

MINING HETEROGENEOUS INFORMATION NETWORKS

BY

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DISSERTATION

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Abstract

Real-world physical objects and abstract data entities are interconnected, forming gigantic networks. By structuring these objects and their interactions into multiple types, such networks become semi-structured heterogeneous information networks. Most real-world applications that handle big data, including interconnected social media and social networks, scientific, engineering, or medical information systems, online e-commerce systems, and most database systems, can be structured into heterogeneous information networks. Therefore, effective analysis of large-scale heterogeneous information networks poses an interesting but critical challenge.

In my thesis, I investigate the principles and methodologies of mining heterogeneous information networks. Departing from many existing network models that view interconnected data as homogeneous graphs or networks, our semi-structured heterogeneous information network model leverages the rich semantics of typed nodes and links in a network and uncovers surprisingly rich knowledge from the network. This semi-structured heterogeneous network modeling leads to a series of new principles and powerful methodologies for mining interconnected data, including (1) ranking-based clustering, (2) meta-path-based similarity search and mining, (3) user-guided relation strength-aware mining, and many other potential developments. This thesis introduces this new research frontier and points out some promising research directions. To my family for all their love.

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Chapter 1

Introduction

We are living in an interconnected world. Most of data or informational objects, individual agents, groups, or components are interconnected or interact with each other, forming numerous, large, interconnected, and sophisticated networks. Without loss of generality, such interconnected networks are called *information networks*. Examples of information networks include social networks, the World Wide Web, research publication networks [41], biological networks [87], highway networks [56], public health systems, electrical power grids, and so on. Clearly, information networks are ubiquitous and form a critical component of modern information infrastructure. The analysis of information networks, or their special kinds, such as social networks and the Web, has gained extremely wide attentions nowadays from researchers in computer science, social science, physics, economics, biology, and so on, with exciting discoveries and successful applications across all the disciplines.

In this thesis, we propose to model real-world systems as semi-structured heterogeneous information networks, by structuring objects and their interactions into different types, and investigate the principles and methodologies for systematically mining such networks. Departing from many existing network models that view interconnected data as homogeneous graphs or networks, our semi-structured heterogeneous information network model leverages the rich semantics of typed nodes and links in a network and uncovers surprisingly rich knowledge from the network.

In this chapter, we first introduce the motivation and overview of this study in Section 1.1, and then introduce the organization of this thesis in Section 1.2.

1.1 Motivation and Overview

In most of the current research on network science, social and information networks are usually assumed to be homogeneous, where nodes are objects of the same entity type (e.g., person) and links are relationships from the same relation type (e.g., friendship). Interesting results have been generated from such studies with numerous influential applications, such as the well-known PageRank algorithm [20] and community detection methods. However, most real world networks are heterogeneous, where nodes and relations are of different types. For example, in a healthcare network, nodes can be patients, doctors, medical tests, diseases, medicines, hospitals, treatments, and so on. On one hand, treating all the nodes as of the same type (e.g., homogeneous information networks) may miss important semantic information. On the other hand, treating every node as of a distinct type (e.g., labeled graph) may also lose valuable schema-level information. It is important to know that patients are of the same kind, comparing with some other kinds, such as doctors or diseases. Thus, a typed, semi-structured heterogeneous network modeling may capture essential semantics of the real world.

Typed, semi-structured heterogeneous information networks are ubiquitous. For example, the network of Facebook consists of persons as well as objects of other types, such as photos, posts, companies, movies, and so on; in addition to friendship between persons, there are relationships of other types, such as person-photo tagging relationships, person-movie liking relationships, person-post publishing relationships, post-post replying relationships, and so on. A university network may consist of several types of objects like students, professors, courses, and departments, as well as their interactions, such as teaching, course registration or departmental association relationships between objects. Similar kinds of examples are everywhere, from social media to scientific, engineering or medical systems, and to online e-commerce systems. Therefore, heterogeneous information networks are powerful and expressive representations of general real-world interactions between different kinds of network entities in diverse domains.

In this thesis, I investigate the principles and methodologies for mining heterogeneous information networks, by leveraging the semantic meaning of the types of nodes and links in a network, and propose models and algorithms that can exploit such rich semantics and solve real-world problems. Heterogeneous information networks often imply rather different semantic structures from that in homogeneous networks. Links in heterogeneous networks indicate the interactions between various types of objects in a network, which can be difficult to be expressed by traditional features. Information is propagated across various kinds of objects in a network, via various kinds of relationships (i.e., heterogeneous links), carrying different semantics and having different strengths in determining the "influence" across linked objects. These principles have laid the foundation for methodologies of handling various mining tasks in heterogeneous information networks, including ranking, clustering, classification, similarity search, relationship prediction and relation strength learning. We will introduce these mining tasks and their associated new principles and methodologies chapter by chapter.

1.1.1 What Are Heterogeneous Information Networks?

An information network represents an abstraction of the real world, focusing on the *objects* and the *interactions* between the objects. It turns out that this level of abstraction has great power in not only representing and storing the essential information about the real-world, but also providing a useful tool to mine knowledge from it, by exploring the power of links. Formally, we define an information network as follows.

Definition 1.1. (Information network) An information network is defined as a directed graph $G = (\mathcal{V}, \mathcal{E})$ with an object type mapping function $\tau : \mathcal{V} \to \mathcal{A}$ and a link type mapping function $\phi : \mathcal{E} \to \mathcal{R}$, where each object $v \in \mathcal{V}$ belongs to one particular object type $\tau(v) \in \mathcal{A}$, each link $e \in \mathcal{E}$ belongs to a particular relation $\phi(e) \in \mathcal{R}$, and if two links belong to the same relation type, the two links share the same starting object type as well as the ending object type.

Different from the traditional network definition, we explicitly distinguish object types and relationship types in the network. Note that, if a relation exists from type A to type B, denoted as ARB, the inverse relation R^{-1} holds naturally for $BR^{-1}A$. R and its inverse R^{-1} are usually not equal, unless the two types are the same and R is symmetric. When the types of objects $|\mathcal{A}| > 1$ or the types of relations $|\mathcal{R}| > 1$, the network is called **heterogeneous information network**; otherwise, it is a **homogeneous information network**.

Given a complex heterogeneous information network, it is necessary to provide its meta level (i.e., schema-level) description for better understanding the object types and link types in the

network. Therefore, we propose the concept of network schema to describe the meta structure of a network.

Definition 1.2. (Network schema) The network schema, denoted as $T_G = (\mathcal{A}, \mathcal{R})$, is a metal template for a heterogeneous network $G = (\mathcal{V}, \mathcal{E})$ with the object type mapping $\tau : \mathcal{V} \to \mathcal{A}$ and the link mapping $\phi : \mathcal{E} \to \mathcal{R}$, which is a directed graph defined over object types \mathcal{A} , with edges as relations from \mathcal{R} .

The network schema of a heterogeneous information network specifies type constraints on the sets of objects and relationships between the objects. These constraints make a heterogeneous information network semi-structured, guiding the exploration of the semantics of the network. An information network following a network schema is then called a *network instance* of the network schema.

Heterogeneous information networks can be constructed from many interconnected, large-scale datasets, ranging from social, scientific, engineering to business applications. Here are a few examples of such networks.

Bibliographic information network: A bibliographic information network, such as the computer science bibliographic information network derived from DBLP, is a typical heterogeneous network, containing objects in four types of entities: paper (P), venue (i.e., conference/journal) (V), author (A), and term (T). For each paper p ∈ P, it has links to a set of authors, a venue, and a set of terms, belonging to a set of link types. It may also contain citation information for some papers, that is, these papers have links to a set of papers cited by the paper and links from a set of papers citing the paper.

The network schema for a bibliographic network and an instance of such a network are shown in Fig. 1.1.

2. Twitter information network: Twitter as a social media can also be considered as an information network, containing objects types such as user, tweet, hashtag and term, and relation (or link) types such as follow between users, post between users and tweets, reply between tweets, use between tweets and terms, and contain between tweets and hashtags.

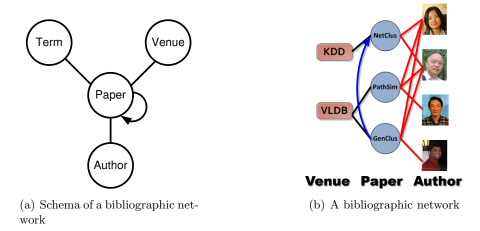


Figure 1.1: A bibliographic network schema and a bibliographic network instance following the schema (only papers, venues and authors are shown).

- 3. Flickr information network: The photo sharing website Flickr can be viewed as an information network, containing a set of object types: *image*, *user*, *tag*, *group*, and *comment*, and a set of relation types, such as *upload* between users and images, *contain* between images and tags, *belong to* between images and groups, *post* between users and comments and *comment* between comments and images.
- 4. **Healthcare information network:** A healthcare system can be modeled as a healthcare information network, containing a set of object types, such as *doctor*, *patient*, *disease*, *treatment*, and *device*, and a set of relation types, such as *used-for* between treatments and diseases, *have* between patients and diseases, and *visit* between patients and doctors.

Heterogeneous information networks can be constructed almost in any domain, such as social networks (e.g., Facebook), e-commerce (e.g., Amazon and eBay), online movie databases (e.g., IMDB), and numerous database applications. Heterogeneous information networks can also be constructed from text data, such as news collections, by entity and relationship extraction using natural language processing and other advanced techniques.

Diverse information can be associated with information networks. Attributes can be attached to the nodes or links in an information network. For example, location attributes, either categorical or numerical, are often associated with some users and tweets in a Twitter information network. Also, temporal information is often associated with nodes and links to reflect the dynamics of

an information network. For example, in a bibliographic information network, new papers and authors emerge every year, as well as their associated links. Beside the structure information of information networks, such information is also helpful or even critical in some tasks on mining information networks.

1.1.2 Why Is Mining Heterogeneous Networks a New Game?

Numerous methods have been developed for the analysis of homogeneous information networks, especially on social networks [2], such as ranking, community detection, link prediction, and influence analysis. However, most of these methods cannot be directly applied to mining heterogeneous information networks. This is not only because heterogeneous links across entities of different types may carry rather different semantic meanings but also because a heterogeneous information network in general captures much richer information than its homogeneous network counterpart. A homogeneous information network is usually obtained by projection from a heterogeneous information network, but with significant information loss. For example, a co-author network can be obtained by projection on co-author information from a more complete heterogeneous bibliographic network. However, such projection will lose valuable information on what subjects and which papers the authors were collaborating on. Moreover, with rich heterogeneous information preserved in an original heterogeneous information network, many powerful and novel data mining functions need to be developed to explore the rich information hidden in the heterogeneous links across entities.

Why is mining heterogeneous networks a new game? Clearly, information propagation across heterogeneous node and links can be very different from that across homogeneous nodes and links. Based on our research into mining heterogeneous information networks, especially our studies on ranking-based clustering [100, 103], ranking-based classification [55, 54], meta-path-based similarity search [99], relationship prediction [96, 97], relation strength learning [95, 101], and network evolution [102], we believe there are a set of new principles that may guide systematic analysis of heterogeneous information networks. We summarize these principles as follows.

1. Information propagation across heterogeneous types of nodes and links. Similar to most of the network analytic studies, links should be used for information propagation in mining

tasks. However, the new game is how to propagate information across heterogeneous types of nodes and links, in particular, how to compute ranking scores, similarity scores, and clusters, and how to make good use of class labels, across heterogeneous nodes and links. No matter how we work out new, delicate measures, definitions, and methodologies, a golden principle is that objects in the networks are interdependent, and knowledge can only be mined using the holistic information in a network.

- 2. Search and mining by exploring network meta structures. Different from homogeneous information networks where objects and links are being treated either as of the same type or as of un-typed nodes or links, heterogeneous information networks in our model are semi-structured and typed, that is, associated with nodes and links structured by a set of types, forming a network schema. The network schema provides a meta structure of the information network. It provides guidance of search and mining of the network and help analyze and understand the semantic meaning of the objects and relations in the network. Meta-path-based similarity search and mining introduced in this thesis has demonstrated the usefulness and the power of exploring network meta structures.
- 3. User-guided exploration of information networks. In a heterogeneous information network, there often exist numerous semantic relationships across multiple types of objects, carrying subtly different semantic meanings. A certain weighted combination of relations or meta-paths may best fit a specific application for a particular user. Therefore, it is often desirable to automatically select the right relation (or meta-path) combinations with appropriate weights for a particular search or mining task based on user's guidance or feedback. User-guided or feedback-based network exploration is a useful strategy.

1.2 Thesis Organization

The first chapter introduces the overview of mining heterogeneous information networks. After that, this thesis is organized into three parts largely following the three principles, each containing two chapters that present methodologies and algorithms for mining heterogeneous information networks, organized by different mining tasks. Finally, Chapter 8 concludes this thesis and outlines

a few open research themes in this domain. The major contents of Chapters 2-7 are summarized as follows.

In Part I: Ranking-Based Clustering, we introduce two studies on the clustering problem, which is one of the basic mining tasks, in heterogeneous information networks, by distinguishing the information propagation across different types of links.

- Chapter 2: Ranking-Based Clustering on Bi-Typed Networks. For link-based clustering of heterogeneous information networks, we need to explore links across heterogeneous types of data. We propose a ranking-based clustering approach, RankClus [100], that generates both clustering and ranking results for a target type of objects in a bi-typed network. This approach is based on the observation that ranking and clustering can mutually enhance each other because objects highly ranked in each cluster may contribute more towards unambiguous clustering, and objects more dedicated to a cluster will be more likely to be highly ranked in the same cluster.
- Chapter 3: Ranking-Based Clustering on Star Networks. Later, we extend our ranking-based clustering study to a more general scenario, where more types of objects exist in the network. Specially, we focus on star networks that have a center type of objects, and propose NetClus [103] that can cluster different types of objects simultaneously. A discussion will be given on how to extend ranking-based clustering to general network schemas.

In Part II: Meta-Path-Based Similarity Search and Mining, we introduce a systematic approach for dealing with general heterogeneous information networks with a specified network schema, using a meta-path-based methodology. Under this framework, similarity search and other mining tasks such as relationship prediction can be addressed by systematic exploration of the network meta structure.

• Chapter 4: Meta-Path-Based Similarity Search. Similarity search plays an important role in the analysis of networks. By considering different linkage paths (i.e., meta-path) in a network, one can derive various semantics on similarity in a heterogeneous information network. A meta-path-based similarity measure, PathSim, is introduced in [99], which aims at finding similar peer objects in the network. PathSim turns out to be more meaningful in many scenarios compared with random-walk based similarity measures.

• Chapter 5: Meta-Path-Based Relationship Prediction. Heterogeneous information network brings interactions among multiple types of objects and hence the possibility of predicting relationships across heterogeneous typed objects. By systematically designing meta-path-based topological features and measures in the network, supervised models can be used to learn appropriate weights associated with different topological features in relationship prediction [96, 97].

In Part III: User-Guided Relation Strength-Aware Mining, we address the issue that the heterogeneity of relations between object types often leads to different mining results that can be chosen by users. With user guidance, the strength of each relation can be automatically learned for improved mining.

- Chapter 6: Relation Strength-Aware Clustering with Incomplete Attributes. Links in networks are frequently used to regularize the attribute-based clustering tasks, *i.e.*, linked objects should have similar cluster labels. However, shall we trust links from different types equally? In this chapter, we propose GenClus [95] to address this problem. By specifying a set of (incomplete) attributes, the strengths of different relations in heterogeneous information networks can be automatically learned to help network clustering.
- Chapter 7: Integrating User-Guided Clustering with Meta-Path Selection. Different meta-paths in a heterogeneous information network represent different relations and carry different semantic meanings. User guidance, such as providing a small set of training examples for some object types, can indicate user preference on the clustering results. Then a preferred meta-path or a weighted meta-path combination of multiple paths can be learned to achieve better consistency between mining results and the training examples [101].

Note that, although three parts largely correspond to the three principles mentioned in Section 1.1, to solve a mining task in heterogeneous information networks usually involves multiple principles simultaneously. For example, the first principle that "information propagation across heterogeneous types of nodes and links" is used throughout all the chapters of this thesis. The second principle "search and mining by exploring network meta structures" can further help to guide the information propagation in heterogeneous information networks, and serves as meta-features for tasks involving learning process. The third principle "user-guided exploration of information

networks" can help us select the desirable ranking function in Part I for the clustering tasks, as well as select the best meta-paths for similarity search in Chapter 4.

Part I

Ranking-Based Clustering

Chapter 2

Ranking-Based Clustering on Bi-Typed Networks

For link-based clustering of heterogeneous information networks, we need to explore links across heterogeneous types of data. In this part, we study how ranking can be computed for different types of objects using different types of links, and show how ranking and clustering mutually enhance each other and finally achieve reasonable ranking and clustering results. In this chapter, we study the problem in a special case of heterogeneous information networks, *i.e.*, the bi-typed networks. In next chapter (Chapter 3), we will study another special but more general case, *i.e.*, the star networks.

2.1 Overview

A great many analytical techniques have been proposed toward a better understanding of information networks, though largely on homogeneous information networks, among which are two prominent ones: ranking and clustering. On one hand, ranking evaluates objects of information networks based on some ranking function that mathematically demonstrates characteristics of objects. With such functions, two objects can be compared, either qualitatively or quantitatively, in a partial order. PageRank [20] and HITS [59], among others, are perhaps the most well-known ranking algorithms over information networks. On the other hand, clustering groups objects based on a certain proximity measure so that similar objects are in the same cluster, whereas dissimilar ones are in different clusters. After all, as two fundamental analytical tools, ranking and clustering can be used to show the overall views of an information network, and hence have been widely used in various applications.

Clustering and ranking are often regarded as *orthogonal* techniques, each applied independently to information network analysis. However, applying only one of them over an information network

often leads to incomplete, or sometimes rather biased, analytical results. For instance, ranking objects over a whole information network without considering which clusters they belong to often leads to dumb results, e.g., ranking database and computer architecture venues or authors together may not make much sense; alternatively, clustering a large number of objects (e.g., thousands of authors) into one cluster without distinction is dull as well. However, integrating two functions together may lead to more comprehensible results, as shown in Example 2.1.

Example 2.1. (Ranking without/with clustering) Consider a set of venues from two areas of (1) DB/DM (i.e., Database and Data Mining) and HW/CA (i.e., Hardware and Computer Architecture), each having 10 venues, as shown in Table 2.1. We choose top 100 authors in each area from DBLP, according to their number of publications in the selected venues. With the authority ranking function specified in Section 2.3.2, our ranking-only algorithm gives top-10 ranked results in Table 2.2. Clearly, the results are rather dumb (because of the mixture of the areas) and are biased towards (i.e., ranked higher for) the HW/CA area. Moreover, such biased ranking result is caused not by the specific ranking function we chose, but by the inherent incomparability between the two areas.

Still consider the same dataset. If we cluster the venues in the DB/DM area and rank both venues and the authors relative to this cluster, the ranking results are shown in Table 2.3.

Table 2.1: A set of venues from two research areas.				
DB/DM	{SIGMOD, VLDB, PODS, ICDE, ICDT, KDD, ICDM, CIK-			
	M, PAKDD, PKDD}			
HW/CA	{ASPLOS, ISCA, DAC, MICRO, ICCAD, HPCA, ISLPED,			
	CODES, DATE, VTS }			

This example shows that good clustering indeed enhances ranking results. Furthermore, assigning ranks to objects often leads to better understanding of each cluster. By integrating both clustering and ranking, one can get more comprehensible results on networks. In this chapter, we introduce RankClus that integrates clustering and ranking for a special case of heterogeneous information networks, *i.e.*, *bi-typed networks*.

Table 2.2: Top-10 ranked venues and authors without clustering.

Rank	Venue	Rank	Authors		
1	DAC	1	Alberto L. Sangiovanni-Vincentelli		
2	ICCAD	2	Robert K. Brayton		
3	DATE	3	Massoud Pedram		
4	ISLPED	4	Miodrag Potkonjak		
5	VTS	5	Andrew B. Kahng		
6	CODES	6	Kwang-Ting Cheng		
7	ISCA	7	Lawrence T. Pileggi		
8	VLDB	8	David Blaauw		
9	SIGMOD	9	Jason Cong		
10	ICDE	10	D. F. Wong		

Table 2.3: Top-10 ranked venues and authors in DB/DM cluster.

Rank	Venue	Rank Authors	
1	VLDB	1	H. V. Jagadish
2	SIGMOD	2 Surajit Chaudhuri	
3	ICDE	3 Divesh Srivastava	
4	PODS	4	Michael Stonebraker
5	KDD	5	Hector Garcia-Molina
6	CIKM	6	Jeffrey F. Naughton
7	ICDM	7	David J. DeWitt
8	PAKDD	8	Jiawei Han
9	ICDT	9 Rakesh Agrawal	
10	PKDD	10 Raghu Ramakrishnan	

2.2 Preliminaries

For the ranking-based clustering problem on a bi-typed network, we are interested in the task of clustering one type of objects (target objects) using the other type of objects (attribute objects) and the links in the network, as well as ranking the objects in each cluster at the same time. For example, given a bi-typed bibliographic network containing venues and authors, where links exist between venues and authors, and between authors and authors, we are interested in clustering venues into different clusters representing different research areas, using the authors and links in the network. At the same time, we could answer who are the top venues and researchers in a research area.

Figure 2.1 illustrates a bi-typed bibliographic network, which contains two types of objects, venues (X) and authors (Y). Two types of links exist in this network: the author-venue publication links, with the weight indicating the number of papers an author has published in a venue, and the author-author co-authorship links, with the weight indicating the number of times two authors have collaborated. The bi-typed network can be represented by a block-wise adjacency matrix:

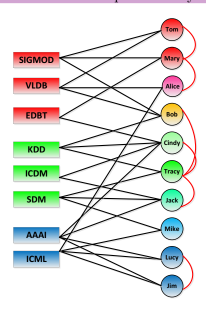


Figure 2.1: A bi-typed bibliographic network.

$$W = \left(\begin{array}{cc} W_{XX} & W_{XY} \\ W_{YX} & W_{YY} \end{array}\right)$$

where W_{XX} , W_{XY} , W_{YX} and W_{YY} each denotes a type of relation between types of the subscripts. Formally, a bi-typed information network can be defined as follows.

Definition 2.1. (Bi-typed information network) Given two types of object sets X and Y, where $X = \{x_1, x_2, ..., x_m\}$, and $Y = \{y_1, y_2, ..., y_n\}$, the graph $G = (\mathcal{V}, \mathcal{E})$ is called a bi-typed information network on types X and Y, if $\mathcal{V} = X \cup Y$ and $\mathcal{E} \subseteq \mathcal{V} \times \mathcal{V}$.

The biggest issue in clustering target objects in a network is that unlike in traditional attributebased dataset, the features for those objects are not explicitly given here. A straightforward way to generate clusters for target objects in a heterogeneous network is to first evaluate the similarity between target objects using a link-based approach, such as SimRank [52], and then apply graph clustering methods [90, 72] to generate clusters. However, to evaluate pair-wise similarity between objects in an information network is a space and time consuming task. Instead, RANKCLUS explores rank distribution for each cluster to generate new measures for target objects, which are low-dimensional. The clusters are improved under the new measure space. More importantly, this measure can be further enhanced during the iterations of the algorithm, because better clustering leads to better ranking, and better ranking gives better ranking-based features thus better clustering results. That is, different from combining ranking and clustering in a two-stage procedure like facet ranking [31, 124], the quality of clustering and ranking can be mutually enhanced in RANKCLUS.

2.3 Ranking Functions

Ranking function is critical in our ranking-based clustering algorithms, which not only provides rank scores for objects to distinguish their importance in a cluster, but also serves as a new feature extraction tool to improve the clustering quality. Current ranking functions are mostly defined on homogeneous networks, such as PageRank [20] and HITS [59]. In this section, we introduce two ranking functions based on the bi-typed bibliographic network: Simple Ranking and Authority Ranking. Ranking functions on more complex heterogeneous networks are discussed at the end of this section.

2.3.1 Simple Ranking

The simplest ranking of venues and authors is based on the number of publications, which is proportional to the numbers of papers accepted by a venue or published by an author.

Formally, given the bi-typed information network with types X and Y, and the adjacency matrix W, simple ranking generates the rank score of type X and type Y as follows:

$$\begin{cases}
\vec{r}_X(x) = \frac{\sum_{j=1}^n W_{XY}(x,j)}{\sum_{i=1}^m \sum_{j=1}^n W_{XY}(i,j)} \\
\vec{r}_Y(y) = \frac{\sum_{i=1}^n W_{XY}(i,y)}{\sum_{i=1}^m \sum_{j=1}^n W_{XY}(i,j)}
\end{cases} (2.1)$$

The time complexity of Simple Ranking is $O(|\mathcal{E}|)$, where $|\mathcal{E}|$ is the number of links. According to simple ranking, authors publishing more papers will have higher rank score, even these papers

are all in junk venues. In fact, simple ranking evaluates importance of each object according to the number of their immediate neighbors.

2.3.2 Authority Ranking

A more useful ranking function we propose here is *authority ranking*, which gives an object higher rank score if it has more authority. Ranking authority merely with publication information seems impossible at the first glance, as citation information could be unavailable or incomplete (such as in the DBLP data, where there is no citation information imported from Citeseer, ACM Digital Library, or Google Scholars). However, two simple empirical rules give us the first clues.

- Rule 1: Highly ranked authors publish many papers in highly ranked venues.
- Rule 2: Highly ranked venues attract many papers from highly ranked authors.

Note that these empirical rules are domain dependent and are usually given by the domain experts who know both the field and the dataset well. From the above heuristics, we define the iterative rank score formulas for authors and venues according to each other as follows.

According to Rule 1, each author's score is determined by the number of papers and their publication forums,

$$\vec{r}_Y(j) = \sum_{i=1}^m W_{YX}(j,i)\vec{r}_X(i)$$
(2.2)

When author j publishes more papers, there are more nonzero and high weighted $W_{YX}(j,i)$, and when the author publishes papers in a higher ranked venue i, which means a higher $\vec{r}_X(i)$, the score of author j will be higher. At the end of each step, $\vec{r}_Y(j)$ is normalized by $\vec{r}_Y(j) \leftarrow \frac{\vec{r}_Y(j)}{\sum_{j'=1}^n \vec{r}_Y(j')}$.

According to Rule 2, the score of each venue is determined by the quantity and quality of papers in the venue, which is measured by their authors' rank scores,

$$\vec{r}_X(i) = \sum_{j=1}^n W_{XY}(i,j)\vec{r}_Y(j)$$
(2.3)

When there are more papers appearing in venue i, there are more non-zero and high weighted $W_{XY}(i,j)$, and if the papers are published by a higher ranked author j, which means a higher $\vec{r}_Y(j)$, the score of venue i will be higher. The score vector is then normalized by $\vec{r}_X(i) \leftarrow \frac{\vec{r}_X(i)}{\sum_{i'=1}^m \vec{r}_X(i')}$.

Note that the normalization will not change the ranking position of an object, but it gives a relative importance score to each object. And as shown in RankClus [100], the two formulas will converge to the primary eigenvector of $W_{XY}W_{YX}$ and $W_{YX}W_{XY}$ respectively.

When considering the co-author information, the scoring function can be further refined by a third rule:

• Rule 3: The rank of an author is enhanced if he or she co-authors with many highly ranked authors.

Adding this new rule, we can calculate rank scores for authors by revising Equation (2.2) as

$$\vec{r}_Y(i) = \alpha \sum_{j=1}^m W_{YX}(i,j)\vec{r}_X(j) + (1-\alpha)\sum_{j=1}^n W_{YY}(i,j)\vec{r}_Y(j)$$
(2.4)

where parameter $\alpha \in [0,1]$ determines how much weight to put on each factor, which can be assigned based on one's belief or learned by some training dataset.

Similarly, we can prove that \vec{r}_Y should be the primary eigenvector of $\alpha W_{YX}W_{XY} + (1-\alpha)W_{YY}$, and \vec{r}_X should be the primary eigenvector of $\alpha W_{XY}(I-(1-\alpha)W_{YY})^{-1}W_{YX}$. Since the iterative process is a power method to calculate primary eigenvectors, the rank score will finally converge.

For authority ranking, the time complexity is $O(t|\mathcal{E}|)$, where t is the iteration number and $|\mathcal{E}|$ is the number of links in the graph. Note that, $|\mathcal{E}| = O(d|\mathcal{V}|) \ll |\mathcal{V}|^2$ in a sparse network, where $|\mathcal{V}|$ is the number of total objects in the network and d is the average link per object.

Different from simple ranking, authority ranking gives an importance measure to each object based on the whole network, rather than its immediate neighborhoods, by the score propagation over the whole network.

2.3.3 Alternative Ranking Functions

Although here we illustrate only two ranking functions, general ranking functions are not confined to them. In practice, a ranking function is not only related to the link property of an information network, but also depends on domain knowledge. For example, in many science fields, journals are given higher weights than conferences when evaluating an author. Moreover, although ranking functions in this section are defined on bi-typed networks, ranking function on heterogeneous networks with more types of objects can be similarly defined. For example, PopRank [82] is a

possible framework for general heterogeneous network, which takes into account the impact both from the same type of objects and from the other types of objects, with different impact factors for different types. When ranking objects in information networks, junk or spam entities are often ranked higher than deserved. For example, authority ranking can be spammed by some bogus venues that accept any submissions due to their large numbers of accepted papers. Techniques that can best use expert knowledge, such as TrustRank [44], can be used to semi-automatically separate reputable, good objects from spam ones. Personalized PageRank [132], that can utilize expert ranking as query and generate rank distributions with respect to such knowledge, can be another choice to integrate expert knowledge.

2.4 From Conditional Rank Distributions to New Clustering Measures

Given a bi-typed bibliographic network, suppose that we have an initial partition on target type X (venue type) and have calculated the conditional rank scores of venues and authors for each clustered network, the next issue becomes how to use the conditional rank scores to further improve the clustering results. Intuitively, for each venue cluster, which could form a research area, the rank scores of authors conditional to this cluster (or research area) should be distinct from that of the authors in other clusters. This implies that these rank scores can be used to derive new features for objects for better clustering. Further, we treat these rank scores as from a discrete rank distribution, as they are non-negative values and summing up to 1, which indicates the subjective belief of how likely one may know an author or a venue according to their authority in each cluster.

Example 2.2. (Conditional rank distribution as cluster feature) Conditional rank distributions in different clusters are distinct from each other, especially when these clusters are reasonably well partitioned. Still using the network of the two-research-area example introduced in Section 2.1, we rank two hundred authors based on two venue clusters, and the two conditional rank distributions are shown in Figure 2.2. From the figure, we can clearly see that DB/DM authors (with IDs from 1 to 100) rank high relative to the DB/DM venues, whereas rank extremely low relative to the HW/CA venues. A similar situation happens for the HW/CA authors (with IDs from 101 to 200).

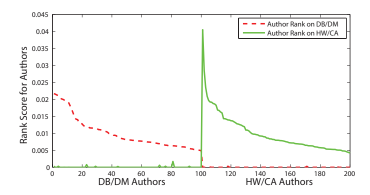


Figure 2.2: Authors' rank distributions over different clusters.

From Example 2.2, one can see that conditional rank distributions for attribute type in each cluster are rather different from each other, and can be used as measures to characterize each cluster. That is, for each cluster X_k , the conditional rank scores of X's and Y's, $\vec{r}_{X|X_k}$ and $\vec{r}_{Y|X_k}$, can be viewed as conditional rank distributions of X and Y, which in fact are the features for cluster X_k .

2.4.1 Cluster Membership for Each Target Object

Suppose we now know the clustering results for type X, which are X_1, X_2, \ldots , and X_K , where K is the number of clusters. Also, according to some given ranking function, we have got conditional rank distribution over Y in each cluster X_k , which is $\vec{r}_{Y|X_k}(k=1,2,\ldots,K)$, and conditional rank distribution over X, which is $\vec{r}_{X|X_k}(k=1,2,\ldots,K)$. In the DBLP scenario, a cluster of venues, e.g., the DB venues, can induce a subnetwork of venues and authors in that area. Conditional rank distributions $\vec{r}_{Y|X_k}$ and $\vec{r}_{X|X_k}$ are calculated on each induced subnetwork of that area. In practice, to avoid the zero rank score for target objects X, we propagate the converged rank scores of Y, $\vec{r}_{Y|X_k}$, for one step further to all target objects, to get a new $\vec{r}_{X|X_k}$. For simplicity, we use $p_k(Y)$ to denote $\vec{r}_{Y|X_k}$ and $p_k(X)$ to denote $\vec{r}_{X|X_k}$ in the following. We use $\pi_{i,k}$ to denote x_i 's cluster membership for cluster k, which in fact is the posterior probability that x_i belongs to cluster k and satisfies $\sum_{k=1}^K \pi_{i,k} = 1$.

According to Bayes' rule, $p(k|x_i) \propto p(x_i|k)p(k)$. Since we already know $p(x_i|k)$, the conditional rank of x_i in cluster k, the goal is thus to estimate p(k), the cluster size of k. In the DBLP scenario, p(k) can be considered as the proportion of papers belonging to the research area that is induced

by the kth venue cluster, where each paper is represented by a link between a venue and an author. According to $p(k|x_i) \propto p(x_i|k)p(k)$, we can see that in general the higher its conditional rank in a cluster $(p(x_i|k))$, the higher possibility an object will belong to that cluster $(p(k|x_i))$. Since the conditional rank scores of X objects are propagated from the conditional rank scores of Y objects, we can also see that highly ranked attribute object has more impact on determining the cluster membership of a target object.

Example 2.3. (Cluster membership as object feature) Following Example 2.2, each venue x_i is represented as a two-dimensional cluster membership vector $(\pi_{i,1}, \pi_{i,2})$. We plot 20 venues according to their cluster membership vectors in Figure 2.3, where different styles of points represent different areas the venues really belong to. From the figure, we can see that the DB/DM venues (denoted as *) and the HW/CA venues (denoted as +) are separated clearly under the new features in terms of cluster membership vectors, which are derived according to the conditional rank distributions of venues and authors with respective to the two research areas.

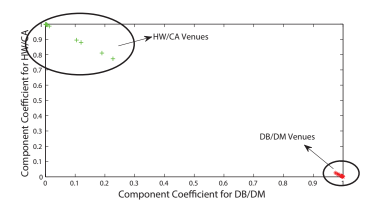


Figure 2.3: Venues' scatter plot based on 2-d cluster membership.

2.4.2 Parameter Estimation Using the EM Algorithm

In order to derive the cluster membership for each target object, we need to estimate the size proportion for each cluster p(k) correctly, which can be viewed as the proportion of the links issued by the target objects belonging to cluster k. In our bi-typed bibliographic information network scenario, this is the proportion of papers belonging to the cluster.

We then build a mixture model for generating links issued by the target objects. Namely,

each link between objects x_i and y_j is generated with the probability $p(x_i, y_j) = \sum_k p_k(x_i, y_j) p(k)$, where $p_k(x_i, y_j)$ denotes the probability of generating such a link in cluster k. We also make an independence assumption that an attribute object y_j issuing a link is independent to a target object x_i accepting this link, which is $p_k(x_i, y_j) = p_k(x_i)p_k(y_j)$. This assumption says once an author writes a paper, he is more likely to submit it to a highly ranked venue to improve his rank; while for venues, they are more likely to accept papers coming from highly ranked authors to improve its rank as well. This idea is similar to preferential attachment [9] of link formation for homogeneous networks, but we are considering more complex rank distributions instead of degrees of objects.

Let Θ be the K-dimensional parameter vector for p(k)'s. The likelihood of observing all the links between types X and Y under the parameter setting is then:

$$L(\Theta|W_{XY}) = p(W_{XY}|\Theta) = \prod_{i=1}^{m} \prod_{j=1}^{n} p(x_i, y_j|\Theta)^{W_{XY}(i,j)}$$
(2.5)

where $p(x_i, y_j | \Theta)$ is the probability to generate link $\langle x_i, y_j \rangle$, given current parameter Θ . The goal is to find the best Θ that maximizes the likelihood. We then apply the EM algorithm [16] to solve the problem. In the E-step, we calculate the conditional distribution $p(z = k | y_j, x_i, \Theta^0)$ based on the current value of Θ^0 :

$$p(z = k|y_j, x_i, \Theta^0) \propto p(x_i, y_j|z = k)p(z = k|\Theta^0) = p_k(x_i)p_k(y_j)p^0(z = k)$$
(2.6)

In the M-Step, we update Θ according to the current Θ^0 :

$$p(z=k) = \frac{\sum_{i=1}^{m} \sum_{j=1}^{n} W_{XY}(i,j) p(z=k|x_i, y_j, \Theta^0)}{\sum_{i=1}^{m} \sum_{j=1}^{n} W_{XY}(i,j)}.$$
 (2.7)

By setting $\Theta^0 = \Theta$, the whole process can be repeated. At each iteration, updating rules from Equations (2.6)-(2.7) are applied, and the likelihood function will converge to a local maximum.

Finally, the cluster membership for each target object x_i in each cluster k, $\pi_{i,k}$, is calculated using Bayes' rule:

$$\pi_{i,k} = p(z = k|x_i) = \frac{p_k(x_i)p(z = k)}{\sum_{l=1}^{K} p_l(x_i)p(z = l)}$$
(2.8)

2.5 Cluster Centers and Distance Measure

After we get the estimations for clustering memberships for each target object x_i by evaluating mixture models, x_i can be represented as a K dimensional vector $\vec{s}_{x_i} = (\pi_{i,1}, \pi_{i,2}, \dots, \pi_{i,K})$. The centers for each cluster can thus be calculated accordingly, which is the mean of \vec{s}_{x_i} for all x_i in each cluster. Next, the distance between an object and cluster $D(x, X_k)$ is defined by 1 minus cosine similarity. The cluster label for each target object can be adjusted accordingly.

2.6 RankClus: Algorithm Summarization

To summarize, the input of RankClus is a bi-typed information network $G = \langle \{X \cup Y\}, W \rangle$, the ranking functions for X and Y, and the cluster number K. The output is K clusters of X with conditional rank scores for each x, and conditional rank scores for each y. The algorithm of RankClus is illustrated in Figure 2.4 and summarized in the following.

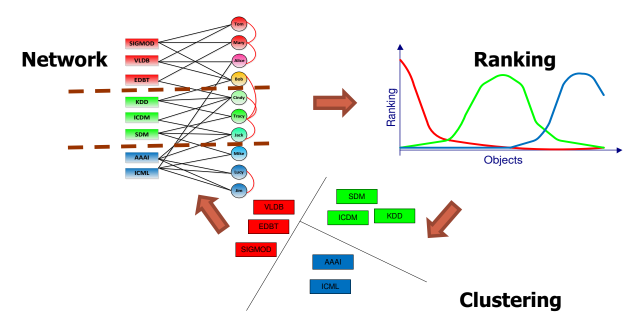


Figure 2.4: The illustration of the RankClus algorithm.

• Step 0: Initialization.

The initial clusters for target objects are generated, by assigning each target object with a cluster label from 1 to K randomly.

- Step 1: Ranking for each cluster.
 - Based on current clusters, K cluster-induced networks are generated accordingly, and the conditional rank distributions for types Y and X are calculated. In this step, we also need to judge whether any cluster is empty, which may be caused by the improper initialization of the algorithm. When some cluster is empty, the algorithm needs to restart in order to generate K clusters.
- Step 2: Estimation of the cluster membership vectors for target objects. In this step, we need to estimate the parameter Θ in the mixture model and get new representations for each target object and centers for each target cluster: \vec{s}_x and \vec{s}_{X_k} . In practice, the number of iterations t for calculating Θ only needs to be set to a small number.
- Step 3: Cluster adjustment.
 In this step, we calculate the distance from each object to each cluster center and assign it to the nearest cluster.
- Repeat Steps 1, 2 and 3 until clusters change only by a very small ratio ε or the number of iterations is bigger than a predefined value iterNum. In practice, we can set $\varepsilon = 0$, and iterNum = 20. In our experiments, the algorithm will converge in less than 5 rounds in most cases for the synthetic dataset and around 10 rounds for the DBLP dataset.

Example 2.4. (Mutual improvement of clustering and ranking) We now apply our algorithm to the two-research-area example. The conditional rank distributions for each cluster and cluster memberships for each venue at each iteration of the running procedure are illustrated in Figure 2.5 (a)-(h). To better explain how our algorithm works, we set an extremely bad initial clustering as the initial state. In Cluster 1, there are 14 venues, half from the DB/DM area and half from the HW/CA area. Cluster 2 contains the remaining 6 venues, which are ICDT, CIKM, PKDD, ASPLOS, ISLPED and CODES. We can see that the partition is quite unbalanced according to the size, and quite mixed according to the area. During the first iteration, the conditional rank distributions for two clusters are very similar to each other (Figure 2.5(a)), and venues are mixed up and biased to Cluster 2 (Figure 2.5(b)). However, we can still adjust their cluster labels according to the cluster centers, and most HW/CA venues go to Cluster 2 and most DB/DM venues

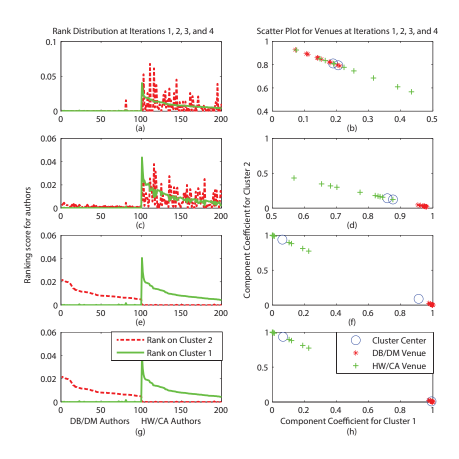


Figure 2.5: Mutual improvement of clusters and ranking through iterations.

go to Cluster 1. At the second iteration, conditional ranking improves somewhat (shown in Figure 2.5(c)) since the clustering (Figure 2.5(b)) is enhanced, and this time clustering results (Figure 2.5(d)) are enhanced dramatically, although they are still biased to one cluster (Cluster 1). At the third iteration, ranking results are improved significantly. Clusters and ranks are further adjusted afterwards, both of which are minor refinements.

At each iteration, the time complexity of RANKCLUS is comprised of three parts: ranking part, mixture model estimation part, and clustering adjustment part. For ranking, if we use simple ranking, the time complexity is $O(|\mathcal{E}|)$. If we use authority ranking, the time complexity is $O(t_1|\mathcal{E}|)$, where $|\mathcal{E}|$ is the number of links, and t_1 is the number of iterations. For mixture model estimation, at each round, we need to calculate the conditional probability for each link in each cluster, the time complexity of which is $O(K|\mathcal{E}|)$. For clustering adjustment, we need to compute the distance between each object (m) and each cluster (K), and the dimension of each object is K, so the time

complexity for this part is $O(mK^2)$. So, overall, the time complexity is $O(t(t_1|\mathcal{E}|+t_2(K|\mathcal{E}|)+mK^2))$, where t is the number of iterations of the whole algorithm and t_2 is the number of iterations of the mixture model. If the network is a sparse network, the time is almost linear to the number of objects.

2.7 Experiments

We now show the effectiveness and efficiency of RANKCLUS algorithm compared with other link-based algorithms, using both synthetic and real datasets.

Case Study on the DBLP Dataset We use the DBLP dataset to generate a bi-typed information network for all the 2676 venues (conferences only) and 20,000 authors with most publications, from the time period of year 1998 to year 2007. Both venue-author relationships and co-author relationships are used. We set the number of clusters K = 15, and apply RANKCLUS with the authority ranking function, with $\alpha = 0.95$. We then pick 5 clusters, and show top-10 venues from each cluster according to the conditional rank scores. The results are shown in Table 2.4, where the research area labels are manually added to each cluster.

Table 2.4: Top-10 venues in 5 clusters generated by RANKCLUS in DBLP.

	DB	Network	AI	Theory	IR
1	VLDB	INFOCOM	AAMAS	SODA	SIGIR
2	ICDE	SIGMETRICS	IJCAI	STOC	ACM Multimedia
3	SIGMOD	ICNP	AAAI	FOCS	CIKM
4	KDD	SIGCOMM	Agents	ICALP	TREC
5	ICDM	MOBICOM	AAAI/IAAI	CCC	JCDL
6	EDBT	ICDCS	ECAI	SPAA	CLEF
7	DASFAA	NETWORKING	RoboCup	PODC	WWW
8	PODS	MobiHoc	IAT	CRYPTO	ECDL
9	SSDBM	ISCC	ICMAS	APPROX-RANDOM	ECIR
10	SDM	SenSys	CP	EUROCRYPT	CIVR

Please note that the clustering and ranking of venues shown in Tables 2.4 have used neither keyword nor citation information, which is the information popularly used in most bibliographic data analysis systems. It is well recognized that citation information is crucial at judging the

influence and impact of a venue or an author in a field. However, by exploring the publication entries only in the DBLP data, the RANKCLUS algorithm can achieve comparable performance as citation studies for clustering and ranking venues and authors. This implies that the collection of publication entries without referring to the keyword and citation information can still tell a lot about the status of venues and authors in a scientific field.

Accuracy and Efficiency Study on Synthetic Data In order to compare accuracy among different clustering algorithms, we generate synthetic bi-typed information networks, which follow the properties of real information networks similar to DBLP. In our experiments, we first fixed the scale of the network and the distribution of links, but change configurations to adjust the density within each cluster and the separateness between different clusters, to obtain 5 different networks (Dataset1 to Dataset5). We set number of clusters K = 3, number of target objects in each cluster as $N_x = [10, 20, 15]$, and number of attribute objects in each cluster as $N_y = [500, 800, 700]$, which are the same for all the 5 datasets. Then we vary the number of links in each cluster (P) and the transition matrix of the proportion of links between different clusters (T), to get the following 5 datasets.

• Dataset1: medium separated and medium density.

```
P = [1000, 1500, 2000],

T = [0.8, 0.05, 0.15; 0.1, 0.8, 0.1; 0.1, 0.05, 0.85]
```

• Dataset2: medium separated and low density.

```
P = [800, 1300, 1200],

T = [0.8, 0.05, 0.15; 0.1, 0.8, 0.1; 0.1, 0.05, 0.85]
```

• Dataset3: medium separated and high density.

```
P = [2000, 3000, 4000],

T = [0.8, 0.05, 0.15; 0.1, 0.8, 0.1; 0.1, 0.05, 0.85]
```

• Dataset4: highly separated and medium density.

```
P = [1000, 1500, 2000],
T = [0.9, 0.05, 0.05, 0.05, 0.9, 0.05; 0.1, 0.05, 0.85]
```

• Dataset5: poorly separated and medium density.

P = [1000, 1500, 2000],

T = [0.7, 0.15, 0.15, 0.15, 0.7, 0.15, 0.15, 0.15, 0.7]

We use the Normalized Mutual Information (NMI) [94] measure to compare the clustering accuracy among different algorithms. For N objects, K clusters, and two clustering results, let $n(i,j), i,j=1,2,\ldots,K$, the number of objects that has the cluster label i in the first clustering result (say generated by the algorithm) and cluster label j in the second clustering result (say the ground truth). We can then define joint distribution $p(i,j) = \frac{n(i,j)}{N}$, row distribution $p_1(j) = \sum_{i=1}^{K} p(i,j)$ and column distribution $p_2(i) = \sum_{j=1}^{K} p(i,j)$, and NMI is defined as:

$$\frac{\sum_{i=1}^{K} \sum_{j=1}^{K} p(i,j) \log(\frac{p(i,j)}{p_1(j)p_2(i)})}{\sqrt{\sum_{j=1}^{K} p_1(j) \log p_1(j) \sum_{i=1}^{K} p_2(i) \log p_2(i)}}$$
(2.9)

We compare RankClus implemented with two ranking functions: Simple Ranking and Authority Ranking, with a state-of-the-art spectral clustering algorithm, the k-way N-cut algorithm [90], implemented with two link-based similarity functions, Jaccard Coefficient and SimRank [52].

Results for accuracy is summarized in Figure 2.6. From the results, we can see that, two versions of RankClus outperform in the first 4 datasets. RankClus with authority ranking is even better, since authority ranking gives a better rank distribution by utilizing the information of the whole network. Through the experiments, we observe that performance of two versions of RankClus and the N-Cut algorithm based on Jaccard coefficient are highly dependent on the data quality, in terms of cluster separateness and link density, while SimRank has a more stable performance, especially on the network that is sparse (Dataset5).

Figure 2.7 summarizes the average execution time of different algorithms over 4 networks with different sizes. We can see that compared with the time-consuming SimRank algorithm, RANKCLUS is very efficient and scalable.

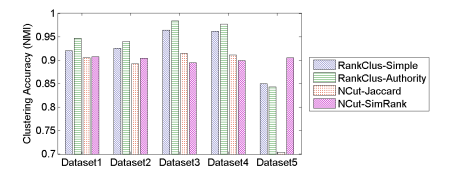


Figure 2.6: Accuracy comparison with baselines in terms of NMI. Dataset1: medium separated and medium density; Dataset2: medium separated and low density; Dataset3: medium separated and high density; Dataset4: highly separated and medium density; and Dataset5: poorly separated and medium density.

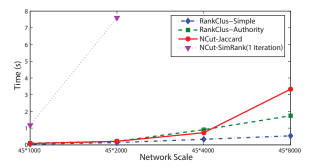


Figure 2.7: Efficiency comparison with baselines in terms of execution time.

2.8 Related Work

In information network analysis, two most important ranking algorithms are PageRank [20] and HITS [59], both of which are successfully applied to the Internet search. PageRank is a link analysis algorithm that assigns a numerical weight to each object of the information network, with the purpose of "measuring" its relative importance within the object set. On the other hand, HITS ranks objects based on two scores: authority and hub. Authority estimates the value of the content of the object, whereas hub measures the value of its links to other objects. Both PageRank and HITS are evaluating the static quality of objects in information network, which is similar to the intrinsic meaning of our ranking methods. However, both PageRank and HITS are designed on the network of web pages, which is a directed homogeneous network, and the weight of the edge is binary. PopRank [82] aims at ranking popularity of web objects. They have considered the role difference of different web pages, and thus turn web pages into a heterogeneous network. They

trained the propagation factor between different types of objects according to partial ranks given by experts. Different from their setting, we will calculate the rank for each type of objects separately (i.e., we do not compare ranks of two objects belonging to different types), rather than consider them in a unified framework. J. E. Hirsch [46] proposed h index originally in the area of physics for characterizing the scientific output of a researcher, which is defined as the number of papers with citation number higher or equal to h. Extensions work [92] shows that it also can work well in computer science area. However, h-index will assign an integer value h to papers, authors, and publication forums, while our work requires that rank sores can be viewed as a rank distribution and thus can serve as a good measure for clustering. What is more, since there are only very limited citation information in DBLP, ranking methods demanding citation cannot work in such kind of data. Instead of proposing a totally new strategy for ranking, we aim at finding empirical rules in the specific area of DBLP data set, and providing ranking function based on these rules, which works well for the specific case. The real novelty lies in our framework is that it tightly integrates ranking and clustering and thus offers informative summary for heterogeneous network such as the DBLP data.

Clustering is another way to summarize information network and discover the underlying structures, which partitions the objects of an information network into subsets (clusters) so that objects in each subset share some common trait. In clustering, proximity between objects is often defined for the purpose of grouping "similar" objects into one cluster, while partitioning dissimilar ones far apart. Spectral graph clustering [90, 72, 28] is state-of-the-art method to do clustering on the homogeneous network. However for heterogeneous network, adjacency matrix of the same type objects does not explicitly exist. Therefore, similarity extraction methods such as SimRank [52] should be applied first, which is an iterative PageRank-like method for computing structural similarity between objects. However, the time cost for SimRank is very high, and other methods such as LinkClus [119] have addressed this issue. Without calculating the pairwise similarity between two objects of the same type, RankClus uses conditional ranking as the measure of clusters, and only needs to calculate the distances between each object and the cluster center.

In web search, there exists an idea of facet ranking [124, 31], which clusters the returned results for each query into different categories, to help users to better retrieve the relevant documents. A

commercial website that illustrates the idea is "vivisimo.com¹." It may seem that facet ranking also integrates ranking with clustering, however, our work is of totally different idea. First, the goal of facet ranking is to help user to better organize the results. The meaning of ranking here is the relevance to the query. RankClus aims at finding higher quality and more informative clusters for target objects with rank information integrated in an information network. Second, facet ranking is a two-stage methodology. In the first stage, relevant results are collected according to the relevance to the query, and then clustering is applied on the collection of returned documents. RankClus integrates ranking and clustering tightly, which are mutually improved during the iterations.

2.9 Discussion

RANKCLUS is the first piece of work that utilizes ranking as cluster feature to improve clustering results and tightly integrates ranking and clustering. However, there are many other issues need to be considered in the future.

First, currently we have only performed experiments on the bi-typed information network. It is interesting to consider utilizing additional information and constraints in the RANKCLUS process. For example, we may interested in adding citation information and text information to the bibliographic data and utilizing the additional information to make refined clustering and ranking. We will extend the framework to a more general type of heterogeneous information networks in next chapter, which focus on star networks.

Second, the empirical rules and its associated weight computation formulas proposed in this study may not be directly transferable to other problem domains. When applying the RANKCLUS methodology to other bibliographic data, such as PubMed, we need to re-consider the empirical rules for ranking functions. When applying the methodology to non-bibliographic data sets, both new ranking functions and the semantics of links need to be reconsidered. Recently, there are some follow-up studies in the domains of PubMed (MedRank [24]) and web photos (RankCompete [22]), which have provided insights to further extend RANKCLUS to other domains.

Third, the quality of ranking function is important to the accuracy of clustering, as it can capture the distinct feature for clusters. However, as we can see, ranking function is highly related

¹http://vivisimo.com

to different domains, how we can automatically extract rules based on a small partial ranking results given by experts could be another interesting problem. A possible solution is to use *meta-path* (Chapter 4) to guide the ranking score propagation in the network, and use user guidance to learn the ranking functions defined on different meta-paths.

2.10 Conclusion

In this chapter, we propose a novel clustering framework called RANKCLUS to integrate clustering with ranking, which generates conditional ranking relative to clusters to improve ranking quality, and uses conditional ranking to generate new measure attributes to improve clustering. As a result, the quality of clustering and ranking are mutually enhanced, which means the clusters are getting more accurate and the ranking is getting more meaningful. Moreover, the clustering results with ranking can provide more informative views of data. Our experiment results show that RANKCLUS can generate more accurate clusters and in a more efficient way than the state-of-the-art link-based clustering method. There are still many research issues to be explored in the RANKCLUS framework. We have identified a few of them in Section 2.9. Clearly, more research is needed to further consolidate this interesting framework and explore its broad applications.

Chapter 3

Ranking-Based Clustering on Star Networks

In this chapter, we study the clustering problem in a more general type of heterogeneous information networks, *i.e.*, star networks, which contains more types of objects. Different from RankClus we are interested in clustering objects from different types simultaneously. A new algorithm NetClus is proposed, which also follows a ranking-based clustering framework.

3.1 Overview

The clustering task we are solving in this chapter is to soft cluster all types of objects for a more general type of heterogeneous information networks that involve more types of objects and more types of links. Among heterogeneous networks, networks with star network schema (called star networks), such as bibliographic networks centered with papers (see Example 3.1) and tagging networks (e.g., http://delicious.com) centered with a tagging event, are popular and important. In fact, any n-nary relation set such as tables in a relational database can be mapped into a star network, with each tuple in the relation as the center object and all attribute entities linking to the center object.

Example 3.1. (A star bibliographic information network) A bibliographic network contains rich information about research publications. It consists of nodes belonging to four types: paper (D), author (A), term (T), and venue (V). Semantically, each paper is written by a group of authors, using a set of terms, and published in a venue (a conference or a journal). Links exist between papers and authors by the relation of "write" and "written by," between papers and terms by the relation of "contain" and "contained in," between papers and venues by the relation of "publish" and "published by." The topological structure of a bibliographic network is shown in the left part of Figure 3.1, which forms a star network schema, where paper is a center type and all other

types (called attribute types) of objects are linked via papers. The network can be represented as $G = (\mathcal{V}, \mathcal{E}, W)$, where $\mathcal{V} = A \cup V \cup T \cup D$, and the weight of the link $\langle x_i, x_j \rangle$, $w_{x_i x_j}$, is defined as:

$$w_{x_ix_j} = \begin{cases} 1, & \text{if } x_i(x_j) \in A \cup V, \ x_j(x_i) \in D, \text{ and } x_i \text{ has link to } x_j \\ c, & \text{if } x_i(x_j) \in T, \ x_j(x_i) \in D, \text{ and } x_i(x_j) \text{ appears c times in } x_j(x_i), \\ 0, & \text{otherwise.} \end{cases}$$

Formally, a general star network with one center type and T attribute types can be defined as follows, where links only exist between the center type and attribute types.

Definition 3.1. (Star network) An information network, $G = (\mathcal{V}, \mathcal{E}, W)$, with T + 1 types of objects (i.e., $\mathcal{V} = \{X_t\}_{t=0}^T$), is called with star network schema, if $\forall e = \langle x_i, x_j \rangle \in \mathcal{E}, x_i \in X_0 \wedge x_j \in X_t \in \mathcal{E}$, or vice versa. G is then called a star network. Type X_0 is the center type (called the target type) and $X_t \in \mathcal{E}$ are attribute types.

In contrast to traditional cluster definition, we propose NetClus to detect net-clusters that contain multiple types of objects and follow the schema of the original network, where each object can softly belong to multiple clusters. A net-cluster example is shown in Example 3.2.

Example 3.2. (The net-cluster of database area) A net-cluster of the database area consists of a set of database venues, authors, terms, and papers, and these objects belong to the database area with a (nontrivial) probability. Accordingly, we can present rank scores for attribute objects such as venues, authors and terms in its own type. With rank distribution, a user can easily grab the important objects in the area. Table 3.1 shows the top ranked venues, authors and terms in the area "database", generated from a 20-venue subnetwork from a "four-area" DBLP dataset (i.e., database, data mining, information retrieval and artificial intelligence) (see Section 3.6), using NETCLUS.

NETCLUS is designed for a heterogeneous network with the star network schema. It is a ranking-based iterative method following the idea of RANKCLUS, that is, ranking is a good feature to help clustering. Different from RANKCLUS, NETCLUS is able to deal with an arbitrary number of types of objects as long as the network is a star network, also the clusters generated are not groups of single typed objects but a set of subnetworks with the same topology as the input network. For a

Table 3.1: Rank scores for venues, authors and terms for the net-cluster of the database research area.

Venue	Rank score	Author	Rank score	Term	Rank score
SIGMOD	0.315	Michael Stonebraker 0.0063		database	0.0529
VLDB	0.306	Surajit Chaudhuri 0.0057		system	0.0322
ICDE	0.194	C. Mohan	0.0053	query	0.0313
PODS	0.109	Michael J. Carey	0.0052	data	0.0251
EDBT	0.046	David J. DeWitt	0.0051	object	0.0138
CIKM	0.019	H. V. Jagadish	0.0043	management	0.0113

given star network and a specified number of clusters K, NETCLUS outputs K net-clusters (Figure 3.1). Each net-cluster is a sub-layer representing a concept of community of the network, which is an induced network from the clustered target objects, and attached with statistic information for each object in the network.

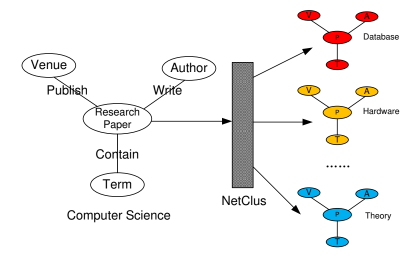


Figure 3.1: Illustration of clustering on a star bibliographic network into net-clusters.

Instead of generating pairwise similarities between objects, which is time consuming and difficult to define under a heterogeneous network, NetClus maps each target object, i.e., that from the center type, into a K-dimensional vector measure, where K is the number of clusters specified by the user. The probabilistic generative model for the target objects in each net-cluster is ranking-based, which factorizes a net-cluster into T independent components, where T is the number of attribute types. In this chapter, we use the star bibliographic network introduced in Example 3.1

to illustrate the NetClus algorithm.

3.2 Ranking Functions

We have introduced ranking functions in Section 2.3, and now we re-examine the two ranking functions for the bibliographic network with a star network schema and illustrate some properties of the two ranking functions for a simple 3-(attribute-)typed star network.

3.2.1 Simple Ranking

Simple ranking is namely the simple occurrence counting for each object normalized in its own type. Given a network G, rank distribution for each attribute type of objects is defined as follows:

$$p(x|T_x, G) = \frac{\sum_{y \in N_G(x)} W_{xy}}{\sum_{x' \in T_x} \sum_{y \in N_G(x')} W_{x'y}}$$
(3.1)

where x is an object from type T_x , and $N_G(x)$ is the set of neighbors of x in G. For example, in the bibliographic network, the rank score for a venue using simple ranking will be proportional to the number of its published papers.

3.2.2 Authority Ranking

Authority ranking for each object is a ranking function that considers the authority propagation of objects in the whole network. Different from the bi-typed information network, we need to consider the rank score propagation over a path in a general heterogeneous information network. For a general star network G, the propagation of authority score from Type X to Type Y through the center type Z is defined as:

$$P(Y|T_Y,G) = W_{YZ}W_{ZX}P(X|T_X,G)$$
(3.2)

where W_{YZ} and W_{ZX} are the weight matrices between the two corresponding types of objects, and can be normalized when necessary. Generally, authority score of one type of objects could be a combination of scores from different types of objects, e.g., that proposed in PopRank [82]. It turns

out that the iteration method of calculating rank distribution is the power method to calculate the primary eigenvector of a square matrix denoting the strength between pairs of objects in that particular type, which can be achieved by selecting a walking path (or a combination of multiple paths) in the network. For more systematic definition of such paths, please refer to Chapter 4 for meta-path-based concepts.

In the DBLP dataset, according to the rules that (1) highly ranked venues accept many good papers published by highly ranked authors, and (2) highly ranked authors publish many good papers in highly ranked venues, we determine the iteration equation as:

$$P(V|T_{V},G) = W_{VD}D_{DA}^{-1}W_{DA}P(A|T_{A},G)$$

$$P(A|T_{A},G) = W_{AD}D_{DV}^{-1}W_{DV}P(V|T_{V},G)$$
(3.3)

where D_{DA} and D_{DV} are the diagonal matrices with the diagonal value equaling to row sum of W_{DA} and W_{DV} , for the normalization purpose. The normalization simply means if a paper was written by multiple authors, we should consider the average rank score of these authors when calculating the rank score of a venue. Since all these matrices are sparse, in practice, the rank scores of objects need only be calculated iteratively according to their limited number of neighbors.

3.2.3 Integrating Ranking Functions with Prior Knowledge

In both ranking functions, prior distributions in different clusters for a certain type of objects can be integrated. For example, a user may give a few representative objects to serve as priors, like terms and venues in each research area. Priors for a given type X are represented in the form $P_P(X|T_X,k), k=1,2,\ldots,K$. The prior is first propagated in the network in a Personalized PageRank [132] way, which propagates scores to objects that are not given in the priors. Then, the propagated prior is linearly combined with the rank distribution calculated by the given ranking function with parameter $\lambda_P \in [0,1]$: the bigger the value, the more dependent on the prior is the final conditional rank distribution.

3.3 Framework of Net Clus Algorithm

Here, we first introduce the general framework of NetClus, and each part of the algorithm will be explained in detail in the following sections. The general idea of the NetClus algorithm given the number of clusters K is composed of the following steps:

- Step 0: Generate initial partitions for target objects and induce initial net-clusters from the original network according to these partitions, i.e., $\{C_k^0\}_{k=1}^K$.
- Step 1: Build ranking-based probabilistic generative model for each net-cluster, i.e., $\{P(x|C_k^t)\}_{k=1}^K$.
- Step 2: Calculate the posterior probabilities for each target object $(p(C_k^t|x))$ and then adjust their cluster assignment according to the new measure defined by the posterior probabilities to each cluster.
- Step 3: Repeat Steps 1 and 2 until the clustering result does not change significantly, i.e., $\{C_k^*\}_{k=1}^K = \{C_k^t\}_{k=1}^K = \{C_k^{t-1}\}_{k=1}^K$.
- Step 4: Calculate the posterior probabilities for each attribute object $(p(C_k^*|x))$ in each netcluster.

In all, the time complexity for NetClus is about linear to $|\mathcal{E}|$, the number of links in the network. When the network is very sparse, which is the real situation in most applications, the time complexity is almost linear to the number of objects in the network.

3.4 Generative Model for Target Objects in a Net-Cluster

According to many studies [37, 9, 79], preferential attachment and assortative mixing exist in many real-world networks, which means an object with a higher degree (i.e., high occurrences) has a higher probability to be attached with a link, and objects with higher occurrences tend to link more to each other. As in the DBLP dataset, 7.64% of the most productive authors publishes 74.2% of all the papers, among which 56.72% of papers are published in merely 8.62% of the biggest venues, which means large size venues and productive authors intend to co-appear via papers. We extend the heuristic by using rank score instead of degree of objects, which denotes the overall importance

of an object in a network. Examples following this intuition include: webpage spammed by many low rank webpages linking to it (high-degree but low rank) will not have too much chance to get a link from a really important webpage, and authors publishing many papers in junk venues will not increase his/her chance to publish a paper in highly ranked venues.

Under this observation, we simplify the network structure by proposing a probabilistic generative model for target objects, where a set of highly ranked attribute objects are more likely to co-appear to generate a center object. To explain this idea, we take the star bibliographic information network as a concrete example and show how the model works, where we assume the number of distinct objects in each type are |A|, |V|, |T|, and |D| respectively, objects in each type are denoted as $A = \{a_1, a_2, \ldots, a_{|A|}\}, V = \{v_1, v_2, \ldots, v_{|V|}\}, T = \{t_1, t_2, \ldots, t_{|T|}\}$ and $D = \{d_1, d_2, \ldots, d_{|D|}\}$.

In order to simplify the complex network with multiple types of objects, we try to factorize the impact of different types of attribute objects and then model the generative behavior of target objects. The idea of factorizing a network is: we assume that given a network G, the probability to visit objects from different attribute types are independent to each other. Also, we make another independence assumption that within the same type of objects, the probability to visit two different objects is independent to each other:

$$p(x_i, x_i | T_x, G) = p(x_i | T_x, G) \times p(x_i | T_x, G)$$

where $x_i, x_j \in T_x$ and T_x is some attribute type.

Now, we build the generative model for target objects given the rank distributions of attribute objects in the network G. Still using bibliographic network as an example, each paper d_i is written by several authors, published in one venue, and comprised of a bag of terms in the title. Therefore, a paper d_i is determined by several attribute objects, say $x_{i1}, x_{i2}, \ldots, x_{in_i}$, where n_i is the number of links d_i has. The probability to generate a paper d_i is equivalent to generating these attribute objects with the occurrence number indicated by the weight of the edge. Under the independency assumptions that we have made, the probability to generate a paper d_i in the network G is defined as follows:

$$p(d_i|G) = \prod_{x \in N_G(d_i)} p(x|T_x, G)^{W_{d_i, x}}$$

where $N_G(d_i)$ is the neighborhood of object d_i in network G, and T_x is used to denote the type of object x. Intuitively, a paper is generated in a cluster with high probability, if the venue it is published in, authors writing this paper and terms appeared in the title all have high probability in that cluster.

3.5 Posterior Probability for Target Objects and Attribute Objects

Once we get the generative model for each net-cluster, we can calculate posterior probabilities for each target object. Now the problem becomes that suppose we know the generative probabilities for each target object generated from each cluster k, k = 1, 2, ..., K, what is the posterior probability that it is generated from cluster k? Here, K is the number of clusters given by the user. As some target objects may not belong to any of K net-cluster, we will calculate K+1 posterior probabilities for each target object instead of K, where the first K posterior probabilities are calculated for each real existing net-clusters $C_1, C_2, ..., C_K$, and the last one in fact is calculated for the original network G. Now, the generative model for target objects in G plays a role as a background model, and target objects that are not very related to any clusters will have high posterior probability in the background model. In this section, we will introduce the method to calculate posterior probabilities for both target objects and attribute objects.

According to the generative model for target objects, the generative probability for a target object d in the target type D in a sub-network G_k is calculated according to the *conditional rank distributions* of attribute types in that sub-network:

$$p(d|G_k) = \prod_{x \in N_{G_k}(d)} p(x|T_x, G_k)^{W_{d,x}}$$
(3.4)

where $N_{G_k}(d)$ denotes the neighborhood of object d in sub-network G_k . In Equation (3.4), in order to avoid zero probabilities in conditional rank scores, each conditional rank score should be first smoothed using global ranking:

$$P_S(X|T_X, G_k) = (1 - \lambda_S)P(X|T_X, G_k) + \lambda_S P(X|T_X, G)$$
(3.5)

where λ_S is a parameter that denotes how much we should utilize the rank distribution from the global ranking.

Smoothing [128] is a well-known technology in information retrieval. One of the reasons that smoothing is required in the language model is to deal with the zero probability problem for missing terms in a document. When calculating generative probabilities of target objects using our ranking-based generative model, we meet a similar problem. For example, for a paper in a given net-cluster, it may link to several objects whose rank score is zero in that cluster. If we simply assign the probability of the target object as zero in that cluster, we will miss the information provided by other objects. In fact, in initial rounds of clustering, objects may be assigned to wrong clusters, if we do not use smoothing technique, they may not have the chance to go back to the correct clusters.

Once a clustering is given on the input network G, say C_1, C_2, \ldots, C_K , we can calculate the posterior probability for each target object (say paper d_i) simply by Bayes' rule: $\pi_{i,k} \propto p(d_i|k) \times p(k)$, where $\pi_{i,k}$ is the probability that paper d_i is generated from cluster k given current generative model, and p(k) denotes the relative size of cluster k, i.e., the probability that a paper belongs to cluster k overall, where $k = 1, 2, \ldots, K, K + 1$.

In order to get the potential cluster size p(k) for each cluster k, we choose cluster size p(k) that maximizes log-likelihood to generate the whole collection of papers and then use the EM algorithm to get the local maximum for p(k).

$$logL = \sum_{i=1}^{|D|} \log(p(d_i)) = \sum_{i=1}^{|D|} \log\left(\sum_{k=1}^{K+1} p(d_i|k)p(k)\right)$$
(3.6)

We use the EM algorithm to get p(k) by simply using the following two iterative formulas, by initially setting $p^{(0)}(k) = \frac{1}{K+1}$:

$$\pi_{i,k}^{(t)} \propto p(d_i|k)p^{(t)}(k); \ p^{(t+1)}(k) = \sum_{i=1}^{|D|} \pi_{i,k}^{(t)}/|D|.$$

When posterior probability is calculated for each target object in each cluster C_k , each target object d can be represented as a K dimensional vector: $\vec{v}(d_i) = (\pi_{i,1}, \pi_{i,2}, \dots, \pi_{i,K})$. The center for each cluster C_k can be represented by the mean vector of all the target objects belonging to

the cluster under the new measure. Next, we calculate cosine similarity between each target object and each center of cluster, and assign the target object into the cluster with the nearest center. A target object is now only belonging to one cluster, and we denote $p(k|d_i)$ as 1 if d_i is assigned to cluster k, 0 otherwise. A new subnetwork G_k can be induced by current target objects belonging to cluster k. The adjustment is an iterative process, until target objects do not change their cluster label significantly under the current measure. Note that, when measuring target objects, we do not use the posterior probability for background model. We make such choices with two reasons: first, the absolute value of posterior probability for background model should not affect the similarity between target objects; second, the sum of the first K posterior probabilities reflects the importance of an object in determining the cluster center.

The posterior probabilities for attribute objects $x \in A \cup V \cup T$ can be calculated as follows:

$$p(k|x) = \sum_{d \in N_G(x)} p(k, d|x) = \sum_{d \in N_G(x)} p(k|d)p(d|x) = \sum_{d \in N_G(x)} p(k|d) \frac{1}{|N_G(x)|}$$

This simply implies, the probability of a venue belonging to cluster C_k equals to the average posterior probability of papers published in the venue; similarly for authors and other attribute objects.

3.6 Experiments

We now study the effectiveness of NetClus and compare it with several state-of-the-art baselines.

Dataset We build star bibliographic networks from DBLP according to Example 3.1. Two networks of different scales are studied. One is a big dataset ("all-area" dataset) which covers all the venues, authors, papers and terms from DBLP. The other is a smaller dataset extracted from DBLP, containing 20 venues from four areas (hence called "four-area" dataset): database, data mining, information retrieval, and artificial intelligence. All authors have ever published papers on any of the 20 venues, and all these papers and terms appeared in the titles of these papers are included in the network. Using the "four-area" dataset, we are able to compare the clustering accuracy with several other methods.

Case Studies We first show the rank distributions in net-clusters we discovered using the "all-area" dataset, which is generated according to authority ranking for venues and authors, by setting venue type as priors and the cluster number as 8. We show four net-clusters in Table 3.2. Also, we can recursively apply NetClus to subnetworks derived from clusters and discover finer level net-clusters. Top-5 authors in a finer level net-cluster about XML area, which is derived from database subnetwork, are shown in Table 3.3.

Table 3.2: Top-5 venues in 4 net-clusters.

Rank	DB and IS	Theory	AI	Software Engineering
1	SIGMOD	STOC	AAAI	ITC
2	VLDB	FOCS	UAI	VTS
3	ICDE	SIAM J. Comput.	IJCAI	POPL
4	SIGIR	SODA	Artif. Intell.	IEEE Trans. Computers
5	KDD	J. Comput. Syst. Sci.	NIPS	IEEE Design & Test of Compu.

Table 3.3: Top-5 authors in "XML" net-cluster.

Rank	Author	
1	Serge Abiteboul	
2	Victor Vianu	
3	Jerome Simeon	
4	Michael J. Carey	
5	Sophie Cluet	

Study on Ranking Functions In Section 3.2, we proposed two ranking functions, namely simple ranking and authority ranking. Here, we study how low dimensional measure derived from rank distributions improve clustering and how clustering can improve this new measure in turn (Figure 3.2). In this study, term is always ranked by simple ranking, and venue and author are ranked by either authority ranking or simple ranking as two different settings.

First, we calculate average KL divergence between each conditional rank distribution and the global rank distribution for each attribute type X to measure the dissimilarity among different

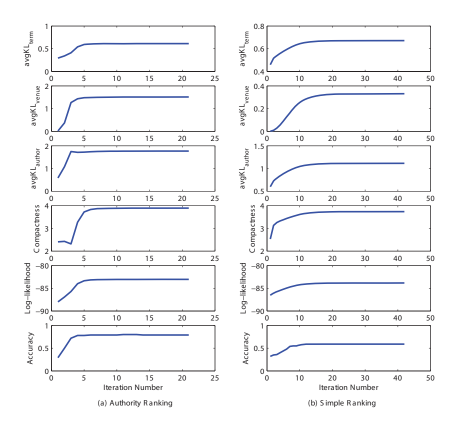


Figure 3.2: The change of ranking and clustering quality in terms of different measurements along with the iterations.

conditional rank distributions, which is denoted as $avg_{KL}(X)$ for type X:

$$avg_{KL}(X) = \frac{1}{K} \sum_{k=1}^{K} D_{KL}(P(X|T_X, G_k)||P(X|T_X, G))$$

Second, in order to evaluate how good the new measure generated in each round for clustering under the ranking function f, we use the compactness, denoted as C_f , which is defined as the average ratio between within-cluster similarity and between-cluster similarity using the new measure:

$$C_f = \frac{1}{|D|} \sum_{k=1}^{K} \sum_{i=1}^{|D_k|} \frac{s(d_{ki}, c_k)}{\sum_{k' \neq k} s(d_{ki}, c_{k'}) / (K - 1)}$$

Third, we trace the accuracy of clustering results for target objects in each round of iteration, which is defined as:

$$accuracy = \frac{1}{|D|} \sum_{i=1}^{D} P_{true}(\cdot|d_i) \cdot P(\cdot|d_i)$$

In other words, we calculate the percentage of papers that are assigned to the correct clusters. However, since |D| is very large even in "four-area" dataset, we manually randomly labeled 100 papers into four clusters and use this paper set to calculate the accuracy.

Fourth, we trace the log-likelihood of the generative model along with the clustering iterations, which is defined in Equation (3.6). From Figure 3.2, we can see authority ranking is better than simple ranking in every measurement.

Study on Parameters In our algorithm, there are two parameters: prior parameter (λ_P) and smoothing parameter setting (λ_S). We use clustering accuracy for sampled papers to test the impact of different settings of parameters to the algorithm. By fixing one of them, we vary the other one. From Figure 3.3(a) and 3.3(b), we find that the larger the prior parameter λ_P , the better the results, while when $\lambda_P > 0.4$, the impact becomes more stable¹; also, the impact of smoothing parameter is very stable, unless it is not too small (less than 0.1) or too big (bigger than 0.8). The results are based on 20 runnings. Priors given for each of the four areas are around 2 or 3 terms. For example, "database" and "system" are priors for database area, with equal prior probabilities.

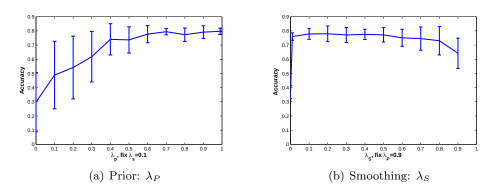


Figure 3.3: Parameter study of λ_P and λ_S .

Accuracy Study In this section, we compare our algorithm with two other algorithms: the topic modeling algorithm PLSA [47] that merely uses term information for documents and RANKCLUS that can only be applied to bi-typed networks. Since both of them cannot directly applied to heterogeneous networks with star network schema, we simplify the network when necessary. For

¹Actually, the extremely poor quality when λ_P is very small is partially caused by the improper accuracy measure at those occasions. When the prior is not big enough to attract the papers from the correct cluster, the clusters generated not necessary have the same cluster label with the priors.

PLSA, only the term type and paper type in the network are used, and we use the same term priors as in NetClus. The accuracy results for papers are in Table 3.4.

Table 3.4: Accuracy of paper clustering results using PLSA and NetClus.

	NetClus (A+V+T+D)	PLSA (T+D)
Accuracy	0.7705	0.608

Since RankClus can only cluster venues, we choose to measure the accuracy of venue cluster. For NetClus, cluster label is obtained according to the largest posterior probability, and Normalized Mutual Information (NMI) is used to measure the accuracy. Since the majority of the authors publish only a few papers, which may contain noise for correctly identifying the clustering of venues, we run RankClus algorithm by setting different thresholds to select subsets of authors. The results are shown in Table 3.5, where d(a) > n means we select authors that have more than n publications to build the bi-typed bibliographic network. All the results are based on 20 runs.

Table 3.5: Accuracy of venue clustering results using RANKCLUS and NETCLUS.

	RANKCLUS	RANKCLUS	RANKCLUS	NetClus
	d(a) > 0	d(a) > 5	d(a) > 10	d(a) > 0
NMI	0.5232	0.8390	0.7573	0.9753

We can see that by using more types of objects in the network, NetClus performs much better than the two baselines that can only utilize partial information in the network.

3.7 Related Work

Clustering on networks and graphs has been widely studied in recent years. Clustering on graphs, often called graph partition, aims at partitioning a given graph into a set of subgraphs based on different criteria, such as minimum cut, min-max cut [35] and normalized cut [90]. Spectral clustering [72] provides an efficient method to get graph partitions which is in fact an NP-hard problem. Rather than investigate the global structure like spectral clustering, several density-based methods [113, 117] are proposed to find clusters in networks which utilizes some neighborhood

information for each object. These methods are all based on the assumption that the network is homogeneous and the adjacent matrix of the network is already defined.

SimRank [52] is able to calculate pairwise similarity between objects by links of a given network, which could deal with heterogenous network, such as bipartite network. However, when the structure of network becomes more complex such as network with star network schema, SimRank cannot give reasonable similarity measures between objects any more. Also, high time complexity is another issue of SimRank, which prevents it from being applied to large scale networks.

An algorithm called RankClus [100] is proposed in Chapter 2, which uses a ranking-clustering mutually enhancement methodology to cluster one type of objects in the heterogeneous network. Although the algorithm is efficient comparing to other algorithms that need to calculate pairwise similarity, there are some weaknesses for RankClus: (1) it has not demonstrated the ability to clustering on networks with arbitrary number of types; and (2) the clusters generated by RankClus only contain one type of objects. In contrast, NetClus can generate net-clusters comprised of objects from multiple types, given any star network.

Other related studies include topic model, such as PLSA [47], which purely uses text information and does not consider link information. Some work such as author-topic model [93] utilizes additional information other than text by designing complex generative models that include additional types of objects. Other work such as [76] intends to optimize a combined objective function with both text and graph constraints. All of these studies are extensions to existing topic model framework, and treat text especially important. In our algorithm, we treat text information just as one common type of objects.

Recently, a different view of clustering on heterogeneous networks [69, 7, 13] appears, which aims at clustering objects from different types simultaneously. Given different cluster number needed for each type of objects, clusters for each type are generated by maximizing some objective function. In this paper, net-cluster follows the original network topology and resembles a community that is comprised of multiple types of objects.

3.8 Discussions

In this section, we present several discussions on further extending the ranking-based clustering framework.

From Star Networks to General Networks First, we give some discussion on how to model more general heterogeneous information networks.

In many bibliographic networks, there could be other link types, for example, the citation links between papers. In this case, the papers can be treated as both target objects and attribute objects. That is, in addition to the the ranking distributions for venues, authors, and terms, we also need to model the ranking distribution for papers in terms of the probabilities of being cited in a certain research area. We then can define the generative probability of a paper by considering citation links in addition. In other cases, people may want to directly model the co-authorship between two authors. A straightforward way is to model the probability of the co-occurrences of author pairs in addition to the probability of the occurrence of single authors. Then the probability of a paper in a cluster is determined also by the probability of all pairs of authors for this paper in that cluster. By directly modeling co-authorship and other relationships, the clustering model can be further enriched due to the introduction of more parameters, but this is likely to cause overfitting as well.

For information networks with arbitrary network structure, the major difficulty of directly applying current model lies in the difficulty of identifying target objects. A possible way to handle this issue is to treat each link as a virtual target object, and model the generation of links from different types separately. The ranking distributions of objects from different types are also modeled separately, but share the same value for the same type of objects, even they could be in different relations.

From Static Networks to Dynamic Networks In real life, networks are rather dynamic, it is interesting to study how objects in the networks form different clusters and how clusters evolve over time, where the clusters represent groups of objects that are closely linked to each other, either due to hidden common interests or due to some social events.

This problem is non-trivial, and it poses several challenges: (1) how to take different types of

objects collectively to detect clusters? (2) how to discover the evolutionary structure (split, merge, and evolve) among clusters of different time windows by modeling the co-evolution of objects in each type? and (3) how to develop an efficient algorithm to solve the problem, as real-world information networks can be very large?

Our recent research EvoNetClus [102] addresses this problem, which studies both the evolution of multiple types of objects in each cluster and the evolutionary structure among the clusters. For example, database and software engineering first formed a huge cluster, but later, database, data mining and machine learning merged into a big cluster, and software engineering itself became an independent cluster. Each cluster is composed of objects from different types, and the clusters evolution is determined by the co-evolution of objects of different types.

From Clustering to Classification In many real-world applications, label information is available for some objects in a heterogeneous information network. Learning from such labeled and unlabeled data via transductive classification can lead to good knowledge extraction of the hidden network structure. Although classification on homogeneous networks [73, 74, 135, 131, 122, 71, 88, 77, 104, 14] has been studied for decades, classification on heterogeneous networks has not been explored until recently. Moreover, both classification and ranking of the nodes (or data objects) in such networks are essential for network analysis. But so far these approaches have generally been performed separately.

In GNetMine [55], we have considered the transductive classification problem on heterogeneous networked data objects which share a common hidden space. Only some objects in the given network are labeled, and the aim is to predict labels for all types of the remaining objects. It has been shown that by distinguishing the relation type in heterogeneous information networks, the classification accuracy can be significantly enhanced.

Further, the authors combine ranking and classification [54] in order to perform more accurate analysis of a heterogeneous information network, following the idea of integrating ranking and clustering together. The intuition is that highly ranked objects within a class should play more important roles in classification. On the other hand, class membership information is important for determining a high quality ranking over a dataset. It is therefore beneficial to integrate classification and ranking in a simultaneous, mutually enhancing process.

3.9 Conclusion

In this chapter, we address a new clustering problem to detect net-clusters on a special heterogeneous network with star network schema, which aims at splitting the original network into K layers and differs the concept from current clustering methods on heterogeneous networks. A novel ranking-based algorithm called NetClus is proposed to find these clusters. The algorithm makes the assumption that within each net-cluster, target objects (i.e., objects from the center type) are generated by a ranking-based probabilistic generative model. Each target object is then mapped into a new low dimensional measure by calculating their posterior probabilities belonging to each net-cluster through their generative models. Our experiments on DBLP data show that NetClus generates more accurate clustering results than the baseline algorithms extended from the topic model and a previous ranking-based algorithm RankClus. Further, NetClus generates more informative clusters, presenting good ranking information and cluster membership for each attribute object in each net-cluster.

Part II

Meta-Path-Based Similarity Search and Mining

Chapter 4

Meta-Path-Based Similarity Search

We now introduce a systematic approach for dealing with general heterogeneous information networks with a specified but arbitrary network schema, using a meta-path-based methodology. Under this framework, similarity search (Chapter 4) and other mining tasks such as relationship prediction (Chapter 5) can be addressed by systematic exploration of the network meta structure.

4.1 Overview

Similarity search, which aims at locating the most relevant information for a query in a large collection of datasets, has been widely studied in many applications. For example, in spatial database, people are interested in finding the k nearest neighbors for a given spatial object [60]; in information retrieval, it is useful to find similar documents for a given document or a given list of keywords. Object similarity is also one of the most primitive concepts for object clustering, recommender systems, and many other data mining functions.

In a similar context, it is critical to provide effective similarity search functions in information networks, to find similar entities for a given entity. In a bibliographic network, a user may be interested in the top-k most similar authors for a given author, or the most similar venues for a given venue. In a network of tagged images such as Flickr, a user may be interested in search for the most similar pictures for a given picture. In an e-commerce system, a user would be interested in search for the most similar products for a given product. Different from the attribute-based similarity search, links play an essential role for similarity search in information networks, especially when the full information about attributes for objects is difficult to obtain.

There are a few studies leveraging link information in networks for similarity search, but most of these studies are focused on homogeneous networks or bipartite networks, such as personalized PageRank (P-PageRank) [53] and SimRank [52]. These similarity measures disregard the subtlety of different types among objects and links. Adoption of such measures to heterogeneous networks has significant drawbacks: even if we just want to compare objects of the same type, going through link paths of different types leads to rather different semantic meanings, and it makes little sense to mix them up and measure the similarity without distinguishing their semantics. For example, Table 4.1 shows the top-4 most similar venues for a given venue, DASFAA, based on (a) the common authors shared by two venues, or (b) the common topics (i.e., terms) shared by two venues. These two scenarios are represented by two distinct meta-paths: (a) VPAPV, denoting that the similarity is defined by the connection path "venue-paper-author-paper-venue," whereas (b) VPTPV, by the connection path "venue-paper-topic-paper-venue." A user can choose either (a) or (b) or their combination based on the preferred similarity semantics. According to Path (a), DASFAA is closer to DEXA, WAIM, and APWeb, that is, those that share many common authors, whereas according to Path (b), it is closer to Data Knowl. Eng., ACM Trans. DB Syst., and Inf. Syst., that is, those that address many common topics. Obviously, different connection paths lead to different semantics of similarity definitions, and produce rather different ranking lists even for the same query object.

Table 4.1: Top-4 most similar venues to "DASFAA" with two meta-paths.

Rank	path: VPAPV	path:VPTPV
1	DASFAA	DASFAA
2	DEXA	Data Knowl. Eng.
3	WAIM	ACM Trans. DB Syst.
4	APWeb	Inf. Syst.

To systematically distinguish the semantics among paths connecting two objects, we introduce a meta-path-based similarity framework for objects of the same type in a heterogeneous network. A meta-path is a sequence of relations between object types, which defines a new composite relation between its starting type and ending type. The meta-path framework provides a powerful mechanism for a user to select an appropriate similarity semantics, by choosing a proper meta-path, or learn it from a set of training examples of similar objects.

In this chapter, we introduce the meta-path-based similarity framework, and relate it to two

well-known existing link-based similarity functions for homogeneous information networks. Especially, we define a novel similarity measure, PathSim, that is able to find peer objects that are not only strongly connected with each other but also share similar visibility in the network. Moreover, we propose an efficient algorithm to support online top-k queries for such similarity search.

4.2 PathSim: A Meta-Path-Based Similarity Measure

The similarity between two objects in a link-based similarity function is determined by how the objects are connected in a network, which can be described using paths. For example, in a co-author network, two authors can be connected either directly or via common co-authors, which are length-1 and length-2 paths respectively. In a heterogeneous information network, however, due to the heterogeneity of the types of links, the way to connect two objects can be much more diverse. For example, in Table 4.2, Column I gives several path instances between authors in a bibliographic network, indicating whether the two authors have co-written a paper; whereas Column II gives several path instances between authors following a different connection path, indicating whether the two authors have ever published papers in the same venue. These two types of connections represent different relationships between authors, each having some different semantic meaning.

Table 4.2: Path instance vs. meta-path in heterogeneous information networks.

	Column I: Connection Type I	Column II: Connection Type II
	$\operatorname{Jim-}P_1\operatorname{-Ann}$	$\operatorname{Jim-}P_1\operatorname{-SIGMOD-}P_2\operatorname{-Ann}$
Path instance	$Mike-P_2-Ann$	$Mike-P_3-SIGMOD-P_2-Ann$
	$Mike-P_3-Bob$	Mike- P_4 -KDD- P_5 -Bob
Meta-path	Author-Paper-Author	Author-Paper-Venue-Paper-Author

Now the questions is, given an arbitrary heterogeneous information network, is there any way to systematically identify all the possible connection types (i.e., relations) between two object types? In order to do so, we propose two important concepts in the following.

4.2.1 Network Schema and Meta-Path

First, given a complex heterogeneous information network, it is necessary to provide its meta level (i.e., schema-level) description for better understanding the network. Therefore, we propose the

concept of **network schema** to describe the meta structure of a network. The formal definition of network schema has been given in Definition 1.2 in Chapter 1. We repeat the definition in the following.

Definition 4.1. (Network schema) The network schema, denoted as $T_G = (\mathcal{A}, \mathcal{R})$, is a meta template for a heterogeneous network $G = (\mathcal{V}, \mathcal{E})$ with the object type mapping $\tau : \mathcal{V} \to \mathcal{A}$ and the link mapping $\phi : \mathcal{E} \to \mathcal{R}$, which is a directed graph defined over object types \mathcal{A} , with edges as relations from \mathcal{R} .

The concept of network schema is similar to that of the ER (Entity-Relationship) model in database systems, but only captures the entity type and their binary relations, without considering the attributes for each entity type. Network schema serves as a template for a network, and tells how many types of objects there are in the network and where the possible links exist. Note that although a relational database can often be transformed into an information network, the latter is more general and can handle more unstructured and non-normalized data and links, and is also easier to deal with graph operations such as calculating the number of paths between two objects.

As we illustrated previously, two objects can be connected via different paths in a heterogeneous information network. For example, two authors can be connected via "author-paper-author" path, "author-paper-venue-paper-author" path, and so on. Formally, these paths are called *meta-paths*, defined as follows.

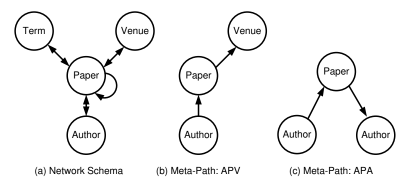


Figure 4.1: Bibliographic network schema and meta-paths.

Definition 4.2. (Meta-path) A meta-path \mathcal{P} is a path defined on the graph of network schema $T_G = (\mathcal{A}, \mathcal{R})$, and is denoted in the form of $A_1 \xrightarrow{R_1} A_2 \xrightarrow{R_2} \dots \xrightarrow{R_l} A_{l+1}$, which defines a composite

relation $R = R_1 \circ R_2 \circ ... \circ R_l$ between types A_1 and A_{l+1} , where \circ denotes the composition operator on relations.

For the bibliographic network schema shown in Figure 4.1 (a), we list two examples of metapaths in Figure 4.1 (b) and (c), where an arrow explicitly shows the direction of a relation. We say a path $p = (a_1 a_2 \dots a_{l+1})$ between a_1 and a_{l+1} in network G follows the meta-path \mathcal{P} , if $\forall i$, $a_i \in A_i$ and each link $e_i = \langle a_i a_{i+1} \rangle$ belongs to each relation R_i in \mathcal{P} . We call these paths as path instances of \mathcal{P} , denoted as $p \in \mathcal{P}$. The examples of path instances have been shown in Table 4.2.

In addition to pointing out the meta-path we are interested in, we also need to consider how to quantify the connection between two objects following a given meta-path. Analogously, a metapath-based measure in an information network corresponds to a feature in a traditional data set, which can be used in many mining tasks.

4.2.2 Meta-Path-Based Similarity Framework

Given a user-specified meta-path, say $\mathcal{P} = (A_1 A_2 \dots A_l)$, several similarity measures can be defined for a pair of objects $x \in A_1$ and $y \in A_l$, according to the path instances between them following the meta-path. We use s(x, y) to denote the similarity between x and y, and list several straightforward measures in the following.

- Path count: the number of path instances p between x and y following \mathcal{P} : $s(x,y) = |\{p : p \in \mathcal{P}\}|$.
- Random walk: s(x,y) is the probability of the random walk that starts form x and ends with y following meta-path \mathcal{P} , which is the sum of the probabilities of all the path instances $p \in \mathcal{P}$ starting with x and ending with y, denoted as Prob(p): $s(x,y) = \sum_{p \in \mathcal{P}} Prob(p)$.
- Pairwise random walk: for a meta-path \mathcal{P} that can be decomposed into two shorter meta-paths with the same length $\mathcal{P} = (\mathcal{P}_1 \mathcal{P}_2)$, s(x, y) is then the pairwise random walk probability starting from objects x and y and reaching the same middle object: $s(x, y) = \sum_{(p_1 p_2) \in (\mathcal{P}_1 \mathcal{P}_2)} Prob(p_1) Prob(p_2^{-1})$, where $Prob(p_1)$ and $Prob(p_2^{-1})$ are random walk probabilities of the two path instances.

In general, we can define a meta-path-based similarity framework for two objects x and y as: $s(x,y) = \sum_{p \in \mathcal{P}} f(p)$, where f(p) is a measure defined on the path instance p between x and y. Note that, P-PageRank and SimRank, two well-known network similarity functions, are weighted

combinations of random walk measure or pairwise random walk measure, respectively, over metapaths with different lengths in homogeneous networks. In order to use P-PageRank and SimRank in heterogeneous information networks, we need to specify the meta-path(s) we are interested in and restrict the random walk on the given meta-path(s).

4.2.3 PathSim: A Novel Similarity Measure

Although there have been several similarity measures as presented above, they are biased to either highly visible objects or highly concentrated objects but cannot capture the semantics of peer similarity. For example, the path count and random walk-based similarity always favor objects with large degrees, and the pairwise random walk-based similarity favors concentrated objects where the majority of the links goes to a small portion of objects. However, in many scenarios, finding similar objects in networks is to *find similar peers*, such as finding similar authors based on their fields and reputation, finding similar actors based on their movie styles and productivity, and finding similar products based on their functions and popularity.

This motivated us to propose a new, meta-path-based similarity measure, called PathSim, that captures the subtlety of peer similarity. The intuition behind it is that two similar peer objects should not only be strongly connected, but also share comparable visibility. As the relation of peer should be symmetric, we confine PathSim to symmetric meta-paths. It is easy to see that, round trip meta-paths in the form of $\mathcal{P} = (\mathcal{P}_l \mathcal{P}_l^{-1})$ are always symmetric.

Definition 4.3. (PathSim: A meta-path-based similarity measure) Given a symmetric meta-path \mathcal{P} , PathSim between two objects x and y of the same type is:

$$s(x,y) = \frac{2 \times |\{p_{x \leadsto y} : p_{x \leadsto y} \in \mathcal{P}\}|}{|\{p_{x \leadsto x} : p_{x \leadsto x} \in \mathcal{P}\}| + |\{p_{y \leadsto y} : p_{y \leadsto y} \in \mathcal{P}\}|}$$

where $p_{x \to y}$ is a path instance between x and y, $p_{x \to x}$ is that between x and x, and $p_{y \to y}$ is that between y and y.

This definition shows that given a meta-path \mathcal{P} , s(x,y) is defined in terms of two parts: (1) their connectivity defined by the number of paths between them following \mathcal{P} ; and (2) the balance of their visibility, where the visibility of an object according \mathcal{P} is defined as the number of path

instances between the object itself following \mathcal{P} . Note that we do count multiple occurrences of a path instance as the weight of the path instance, which is the product of weights of all the links in the path instance.

Table 4.3 presents in three measures the results of finding top-5 similar authors for "Anhai Doan," who is an established young researcher in the database field, under the meta-path APVPA (based on their shared venues), in the database and information system (DBIS) area. P-PageRank returns the most similar authors as those published substantially in the area, that is, highly ranked authors; SimRank returns a set of authors that are concentrated on a small number of venues shared with Doan; whereas PathSim returns Patel, Deshpande, Yang and Miller, who share very similar publication records and are also rising stars in the database field as Doan. Obviously, PathSim captures desired semantic similarity as peers in such networks.

Table 4.3: Top-5 similar authors for "AnHai Doan" in the DBIS area.

Rank	P-PageRank	SimRank	PathSim
1	AnHai Doan	AnHai Doan	AnHai Doan
2	Philip S. Yu	Douglas W. Cornell	Jignesh M. Patel
3	Jiawei Han	Adam Silberstein	Amol Deshpande
4	Hector Garcia-Molina	Samuel DeFazio	Jun Yang
5	Gerhard Weikum	Curt Ellmann	Renée J. Miller

The calculation of PathSim between any two objects of the same type given a certain meta-path involves matrix multiplication. Given a network $G = (\mathcal{V}, \mathcal{E})$ and its network schema T_G , we call the new adjacency matrix for a meta-path $\mathcal{P} = (A_1 A_2 \dots A_l)$ a relation matrix, and is defined as $M = W_{A_1 A_2} W_{A_2 A_3} \dots W_{A_{l-1} A_l}$, where $W_{A_i A_j}$ is the adjacency matrix between type A_i and type A_j . M(i,j) represents the number of paths instances between object $x_i \in A_1$ and object $y_j \in A_l$ under meta-path \mathcal{P} .

For example, relation matrix M for the meta-path $\mathcal{P} = (APA)$ is a co-author matrix, with each element representing the number of co-authored papers for the pair of authors. Given a symmetric meta-path \mathcal{P} , PathSim between two objects x_i and x_j of the same type can be calculated as $s(x_i, x_j) = \frac{2M_{ij}}{M_{ii} + M_{jj}}$, where M is the relation matrix for the meta-path \mathcal{P} , M_{ii} and M_{jj} are the visibility for x_i and x_j in the network given the meta-path.

It is easy to see that the relation matrix for the reverse meta-path of \mathcal{P}_l , which is \mathcal{P}_l^{-1} , is

the transpose of relation matrix for \mathcal{P}_l . In this paper, we only consider the meta-path in the round trip form of $\mathcal{P} = (\mathcal{P}_l \mathcal{P}_l^{-1})$, to guarantee its symmetry and therefore the symmetry of the PathSim measure. By viewing PathSim in the meta-path-based similarity framework, $f(p) = 2\frac{w(a_1,a_2)...w(a_{l-1},a_l)}{M_{ii}+M_{jj}}$, for any path instance p starting from x_i and ending with x_j following the meta-path $(a_1 = x_i \text{ and } a_l = x_j)$, where $w(a_m, a_n)$ is the weight for the link $\langle a_m, a_n \rangle$ defined in the adjacency matrix.

Some good properties of PathSim, such as symmetric, self-maximum and balance of visibility, are shown in Theorem 4.1. For the balance property, we can see that the larger the difference of the visibility of the two objects, the smaller the upper bound for their PathSim similarity.

Theorem 4.1. (Properties of PathSim)

- 1. **Symmetric**: $s(x_i, x_j) = s(x_j, x_i)$.
- 2. **Self-maximum**: $s(x_i, x_j) \in [0, 1]$, and $s(x_i, x_i) = 1$.
- 3. Balance of Visibility: $s(x_i, x_j) \leq \frac{2}{\sqrt{M_{ii}/M_{jj}} + \sqrt{M_{jj}/M_{ii}}}$.

Although using meta-path-based similarity we can define similarity between two objects given any round trip meta-paths, the following theorem tells us a very long meta-path is not very meaningful. Indeed, due to the sparsity of real networks, objects that are similar may share no immediate neighbors, and longer meta-paths will propagate similarities to remote neighborhoods. For example, as in the DBLP example, if we consider the meta-path APA, only two authors that are co-authors have a non-zero similarity score; but if we consider longer meta-paths like APVPA or APTPA, authors will be considered to be similar if they have published papers in a similar set of venues or sharing a similar set of terms no matter whether they have co-authored. But how far should we keep going? The following theorem tells us that a very long meta-path may be misleading. We now use \mathcal{P}^k to denote a meta-path repeating k times of the basic meta-path pattern of \mathcal{P} , e.g., $(AVA)^2 = (AVAVA)$.

Theorem 4.2. (Limiting behavior of PathSim under infinity-length meta-path) Let meta-path $\mathcal{P}^{(k)} = (\mathcal{P}_l \mathcal{P}_l^{-1})^k$, $M_{\mathcal{P}}$ be the relation matrix for meta-path \mathcal{P}_l , and $M^{(k)} = (M_{\mathcal{P}} M_{\mathcal{P}}^T)^k$ be the

relation matrix for $\mathcal{P}^{(k)}$, then by PathSim, the similarity between objects x_i and x_j as $k \to \infty$ is:

$$\lim_{k \to \infty} s^{(k)}(i,j) = \frac{2\mathbf{r}(i)\mathbf{r}(j)}{\mathbf{r}(i)\mathbf{r}(i) + \mathbf{r}(j)\mathbf{r}(j)} = \frac{2}{\frac{\mathbf{r}(i)}{\mathbf{r}(j)} + \frac{\mathbf{r}(j)}{\mathbf{r}(i)}}$$

where \mathbf{r} is the primary eigenvector of M, and $\mathbf{r}(i)$ is the i_{th} item of \mathbf{r} .

As primary eigenvectors can be used as authority ranking of objects [100], the similarity between two objects under an infinite meta-path can be viewed as a measure defined on their rankings ($\mathbf{r}(i)$) is the ranking score for object x_i). Two objects with more similar ranking scores will have higher similarity (e.g., SIGMOD will be similar to AAAI). Later experiments (Table 4.9) will show that this similarity, with the meaning of global ranking, is not that useful. Note that, the convergence of PathSim with respect to path length is usually very fast and the length of 10 for networks of the scale of DBLP can almost achieve the effect of a meta-path with an infinite length. Therefore, in this paper, we only aim at solving the top-k similarity search problem for a relatively short meta-path.

Even for a relatively short length, it may still be inefficient in both time and space to materialize all the meta-paths. Thus we propose in Section 4.3 materializing relation matrices for short length meta-paths, and concatenating them online to get longer ones for a given query.

4.3 Online Query Processing for Single Meta-Path

Compared with P-PageRank and SimRank, the calculation for PathSim is much more efficient, as it is a local graph measure. But it still involves expensive matrix multiplication operations for top-k search functions, as we need to calculate the similarity between a query and every object of the same type in the network. One possible solution is to materialize all the meta-paths within a given length. Unfortunately, it is time and space expensive to materialize all the possible meta-paths. For example, in the DBLP network, the similarity matrix corresponding to a length-4 meta-path, APVPA, for identifying similar authors publishing in common venues is a $710K \times 710K$ matrix, whose non-empty elements reaches 5G, and requires storage size more than 40GB.

In order to support fast online query processing for large-scale networks, we propose a methodology that partially materializes short length meta-paths and then concatenates them online to derive longer meta-path-based similarity. First, a baseline method (PathSim-baseline) is proposed, which computes the similarity between query object x and all the candidate objects y of the same type. Next, a co-clustering based pruning method (PathSim-pruning) is proposed, which prunes candidate objects that are not promising according to their similarity upper bounds. Both algorithms return exact top-k results for the given query. Note that the same methodology can be adopted by other meta-path-based similarity measures, such as random walk and pairwise random walk, by taking a different definition of similarity matrix accordingly.

4.3.1 Single Meta-Path Concatenation

Given a meta-path $\mathcal{P} = (\mathcal{P}_l \mathcal{P}_l^{-1})$, where $\mathcal{P}_l = (A_1 \cdots A_l)$, the relation matrix for path \mathcal{P}_l is $M_{\mathcal{P}} = W_{A_1 A_2} W_{A_2 A_3} \cdots W_{A_{l-1} A_l}$, the relation matrix for path \mathcal{P} is $M = M_{\mathcal{P}} M_{\mathcal{P}}^T$. Let n be the number of objects in A_1 . For a query object $x_i \in A_1$, if we compute the top-k most similar objects $x_j \in A_1$ for x_i on-the-fly, without materializing any intermediate results, computing M from scratch would be very expensive. On the other hand, if we have pre-computed and stored the relation matrix $M = M_{\mathcal{P}} M_{\mathcal{P}}^T$, it would be a trivial problem to get the query results: we only need to locate the corresponding row in the matrix for the query x_i , re-scale it using $(M_{ii} + M_{jj})/2$, and finally sort the new vector and return the top-k objects. However, fully materializing the relation matrices for all possible meta-paths is also impractical, since the space complexity $(O(n^2))$ would prevent us from storing M for every meta-path. Instead of taking the above extreme, we partially materialize relation matrix $M_{\mathcal{P}}^T$ for meta-path \mathcal{P}_l^{-1} , and compute top-k results online by concatenating \mathcal{P}_l and \mathcal{P}_l^{-1} into \mathcal{P} without full matrix multiplication.

We now examine the concatenation problem, that is, when the relation matrix M for the full meta-path \mathcal{P} is not pre-computed and stored, but the relation matrix $M_{\mathcal{P}}^T$ corresponding to the partial meta-path \mathcal{P}_l^{-1} is available. In this case, we assume the main diagonal of M, that is, $D = (M_{11}, \ldots, M_{nn})$, is pre-computed and stored. Since for $M_{ii} = M_{\mathcal{P}}(i,:)M_{\mathcal{P}}(i,:)^T$, the calculation only involves $M_{\mathcal{P}}(i,:)$ itself, and only O(nd) in time and O(n) in space are required, where d is the average number of non-zero elements in each row of $M_{\mathcal{P}}$ for each object.

In this study, we only consider concatenating the partial paths \mathcal{P}_l and \mathcal{P}_l^{-1} into the form $\mathcal{P} = \mathcal{P}_l \mathcal{P}_l^{-1}$ or $\mathcal{P} = \mathcal{P}_l^{-1} \mathcal{P}_l$. For example, given a pre-stored meta-path APV, we are able to

answer queries for meta-paths APVPA and VPAPV. For our DBLP network, to store relation matrix for partial meta-path APV only needs around 25M space, which is less than 0.1% of the space for materializing meta-path APVPA. Other concatenation forms that may lead to different optimization methods are also possible (e.g., concatenating several short meta-paths). In the following discussion, we focus on the algorithms using the concatenation form $\mathcal{P} = \mathcal{P}_l \mathcal{P}_l^{-1}$.

4.3.2 Baseline

Suppose we know the relation matrix $M_{\mathcal{D}}$ for meta-path P_l , and the diagonal vector $D = (M_{ii})_{i=1}^n$, in order to get top-k objects $x_j \in A_1$ with the highest similarity for the query x_i , we need to compute s(i,j) for all x_j . The straightforward baseline is: (1) first apply vector-matrix multiplication to get $M(i,:) = M_{\mathcal{D}}(i,:)M_{\mathcal{D}}^T$; (2) calculate $s(i,j) = \frac{2M(i,j)}{M(i,i)+M(j,j)}$ for all $x_j \in A_1$; and (3) sort s(i,j) to return the top-k list in the final step. When n is very large, the vector-matrix computation will be too time consuming to check every possible object x_j . Therefore, we first select x_j 's that are not orthogonal to x_i in the vector form, by following the links from x_i to find 2-step neighbors in relation matrix $M_{\mathcal{D}}$, that is, $x_j \in CandidateSet = \{\bigcup_{y_k \in M_{\mathcal{D}}.neighbors(x_i)} M_{\mathcal{D}}^T.neighbors(y_k)\}$, where $M_{\mathcal{D}}.neighbors(x_i) = \{y_k | M_{\mathcal{D}}(x_i, y_k) \neq 0\}$, which can be easily obtained in the sparse matrix form of $M_{\mathcal{D}}$ that indexes both rows and columns. This will be much more efficient than pairwise comparison between the query and all the objects of that type. We call this baseline concatenation algorithm as PathSim-baseline.

The PathSim-baseline algorithm, however, is still time consuming if the candidate set is large. Although $M_{\mathcal{P}}$ can be relatively sparse given a short length meta-path, after concatenation, M could be dense, i.e., the CandidateSet could be very large. Still, considering the query object and one candidate object represented by query vector and candidate vector, the dot product between them is proportional to the size of their non-zero elements. The time complexity for computing PathSim for each candidate is O(d) on average and O(m) in the worst case, that is, O(nm) in the worst case for all the candidates, where n is the row size of $M_{\mathcal{P}}$ (i.e., the number of objects in type A_1), m the column size of $M_{\mathcal{P}}$ (i.e., the number of objects in type A_l), and d the average non-zero element for each object in $M_{\mathcal{P}}$. We now propose a co-clustering based top-k concatenation algorithm, by which non-promising target objects are dynamically filtered out to reduce the search

space.

4.3.3 Co-Clustering Based Pruning

In the baseline algorithm, the computational costs involve two factors. First, the more candidates to check, the more time the algorithm will take; second, for each candidate, the dot product of query vector and candidate vector will at most involve m operations, where m is the vector length. The intuition to speed up the search is to prune unpromising candidate objects using simpler calculations. Based on the intuition, we propose a co-clustering-based (i.e., clustering rows and columns of a matrix simultaneously) path concatenation method, which first generates co-clusters of two types of objects for partial relation matrix, then stores necessary statistics for each of the blocks corresponding to different co-cluster pairs, and then uses the block statistics to prune the search space. For better illustration, we call clusters of type A_1 as target clusters, since the objects in A_1 are the targets for the query; and call clusters of type A_l as **feature clusters**, since the objects in A_l serve as features to calculate the similarity between the query and the target objects. By partitioning A_1 into different target clusters, if a whole target cluster is not similar to the query, then all the objects in the target cluster are likely not in the final top-k lists and can be pruned. By partitioning A_l into different feature clusters, cheaper calculations on the dimensionreduced query vector and candidate vectors can be used to derive the similarity upper bounds. This pruning idea is illustrated in Figure 4.2 using a toy example with 9 target objects and 6 feature objects.

Now we introduce the detailed algorithms in the following.

1. Block-wise Commuting Matrix Materialization

The first problem is how to generate these clusters for each commuting matrix $M_{\mathcal{P}}$. Since one commuting matrix can be used for the concatenation into two longer meta-paths, i.e., $M_{\mathcal{P}}M_{\mathcal{P}}^T$ and $M_{\mathcal{P}}^TM_{\mathcal{P}}$, we hope to find co-clusters of feature cluster and target cluster, within which all values are similar to each other. We use a greedy KL-divergence based co-clustering method (summarized in Algorithm 4.1), which is similar to the information-theoretic co-clustering proposed in [34], but simplifies the feature space for each object by merely using the feature cluster information. For example, for $\mathcal{P}_l = (APC)$, we will use the conditional probability of author clusters appearing in

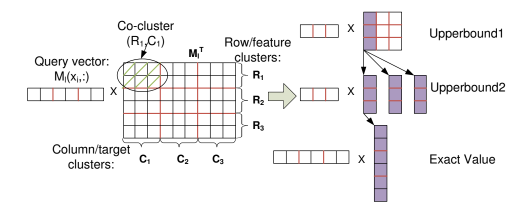


Figure 4.2: Illustration of pruning strategy. Given the partial relation matrix M_l^T and its 3×3 co-clusters, and the query vector $M_l(x_i,:)$ for query object x_i , first the query vector is compressed into the aggregated query vector with the length of 3, and the upper bounds of the similarity between the query and all the 3 target clusters are calculated based on the aggregated query vector and aggregated cluster vectors; second, for each of the target clusters, if they cannot be pruned, calculate the upper bound of the similarity between the query and each of the 3 candidates within the cluster using aggregated vectors; third, if the candidates cannot be pruned, calculate the exact similarity value using the non-aggregated query vector and candidate vectors.

some conference c, say $p(\hat{A}_u|c = \text{``VLDB''})$, as the feature for conference c, use the conditional probability of author clusters in some conference cluster \hat{C}_v , say $p(\hat{A}_u|\hat{C}_v = \text{``DB''})$, as the feature for conference cluster \hat{C}_v , and assign the conference to the conference cluster with the minimum KL-divergence. The adjustment is the same for author type given current conference clusters. The whole process is repeated for conference type and author type alternately, until the clusters do not change any more.

The time complexity of Algorithm 4.1 is O(t(m+n)(UV)), where t is the number of iterations, m and n are the number of objects for feature type and target type, U and V are the numbers of clusters for feature type and target type. Compared with the original O(mn(U+V)) algorithm in [34], it is much more efficient. Sampling-based variation algorithm such as in [83] can be applied for further faster co-clustering. In our experiment setting, we will select objects with higher degrees for the clustering, and assign those with smaller degrees to the existing clusters.

Once the clusters for each type of objects are obtained, the commuting matrix can be decomposed into disjoint blocks. To facilitate further concatenation on two meta-paths for queries, necessary statistical information is stored for each block. For each block b denoted by row cluster R_u and column cluster C_v , we store:

Algorithm 4.1 Greedy Co-Clustering Algorithm

Input: Commuting Matrix $M_{\mathcal{P}}^T$, number of feature clusters (row clusters) U, number of target clusters (column clusters) V

```
Output: row clusters \{R_u\}_{u=1}^U, column clusters \{C_v\}_{v=1}^V
 1: //Initialization.
 2: Randomly assign row objects into \{R_u\}_{u=1}^U;
 3: Randomly assign column objects into \{C_v\}_{v=1}^V;
 5:
         //get center vector of each R_u:
        f(R_u) = \frac{1}{|R_u|} \sum_{v=1}^{V} M_{\mathcal{P}}^T(R_u, C_v);
//Adjust row objects
 6:
 7:
         foreach object x_i in row objects do
 8:
            f(x_i) = \sum_{v=1}^{V} M_{\mathcal{P}}^T(x_i, C_v);
assign x_i into R_u, u = \arg\min_k KL(f(x_i)||f(R_u));
 9:
10:
         //get center vector of each C_v:
11:
         f(C_v) = \frac{1}{|C_v|} \sum_{u=1}^{U} M_{\mathcal{P}}^T(R_u, C_v)
12:
         //Adjust column objects
13:
         foreach object y_j in row objects do
14:
            f(y_j) = \sum_{u=1}^{U} M_{\mathcal{P}}(R_u, y_j);
assign y_j into C_v, v = \arg\min_l KL(f(y_j)||f(C_v));
15:
17: until \{R_u\}, \{C_v\} do not change significantly.
```

1. Element sum of each block $T^{\{U \times V\}}$:

$$t_{uv} = \sum_{i \in R_u} \sum_{j \in C_v} M_{\mathcal{P}}^T(i, j);$$

2. Sum of row vectors (1-norm of each column vector) of each block $T_1^{\{U \times m\}}$:

$$t_{uv,1}(j) = \sum_{i \in R_u} M_{\mathcal{D}}^T(i,j), \text{ for } j \in C_v;$$

3. Square root of sum of square of row vectors (2-norm of each column vector) of each block $TT_1^{\{U\times m\}}$:

$$t_{uv,1}^2(j) = \sqrt{\sum_{i \in R_u} (M_{\mathcal{P}}^T(i,j))^2}$$
, for $j \in C_v$;

4. Sum of column vectors (1-norm of each row vector) of each block $T_2^{\{n \times V\}}$:

$$t_{uv,2}(i) = \sum_{j \in C_v} M_{\mathcal{P}}^T(i,j), \text{ for } i \in R_u;$$

5. Square root of sum of square of column vectors (2-norm of each row vector) of each block $TT_2^{\{n\times V\}}$:

$$t_{uv,2}^{2}(i) = \sqrt{\sum_{j \in C_{v}} (M_{\mathcal{P}}^{T}(i,j))^{2}}, \text{ for } i \in R_{u}.$$

2. Pruning Strategy in Path Concatenation

Now let's focus on how we can get top-k results efficiently for a query given the materialized block-wise commuting matrix. The intuition is that we first check the most promising target cluster,

then if possible, prune the whole target cluster; if not, we first use simple calculations to decide whether we need to further calculate the similarity between the query and the candidate object, then compute the exact similarity value using more complex operations only for those needed.

Theorem 4.3. Bounds for block-based similarity measure approximation. Given a query object x, the query vector is $\mathbf{x} = M_{\mathcal{P}}(x,:)$. Let D be the diagonal vector of M, let $\hat{\mathbf{x}}_1$ be the compressed query vector given feature clusters $\{R_u\}_{u=1}^U$, where $\hat{\mathbf{x}}_1(u) = \max_{j \in R_u} \{\mathbf{x}(j)\}$, and let $\hat{\mathbf{x}}_2$ be the 2-norm query vector given feature clusters R_u , where $\hat{\mathbf{x}}_2(u) = \sqrt{\sum_{j \in R_u} \mathbf{x}(j)^2}$, the similarity between x and target cluster C_v , and the similarity between x and candidate $y \in C_v$ can be estimated using the following upper bounds:

1. upperbound 1:
$$\forall y \in C_v, s(x,y) \leq s(x,C_v) = \sum_{y \in C_v} s(x,y) \leq \frac{2\hat{\mathbf{x}}_1^T T(:,v)}{D(x)+1};$$

2. upperbound 2:
$$\forall y \in C_v, s(x,y) \leq \frac{2\hat{\mathbf{x}}_2^T T T_1(:,y)}{D(x) + D(y)}$$
.

Proof. See Proof in the Appendix A.

In Theorem 4.3, the upper bound for $s(x, C_v)$ can be used to find the most promising target clusters as well as to prune target clusters if it is smaller than the lowest similarity in the current top-k results. The upper bound for s(x, y) can be used to prune target objects that are not promising, which only needs at most U times calculation, whereas the exact calculation needs at most m times calculation. Here, U is the number of feature clusters and m is the number of feature objects, i.e., objects of type A_l .

The search strategy is to first sort the target clusters according to their upper bound of the similarity between the query x and the cluster C_v , i.e., $s(x, C_v)$, in a decreasing order. The higher the similarity the more likely this cluster contains more similar objects to x. It is very critical to use the order to check the most promising target clusters first, by which the most desirable objects are retrieved at an early stage and the upper bounds then have stronger power to prune the remaining candidates. When a new target cluster needs to be checked, the upper bound can be used to prune the whole target cluster and all the remaining target clusters, if it is smaller than the k-th value of the current top-k list. Next, when going to check the candidates within the target cluster, the upper bound between query object x and candidate y can be used to prune non-promising candidates if it is smaller than the current threshold. The algorithm PathSim-pruning

is summarized in Algorithm 4.3. On Line 5, min(S) is the lowest similarity in the current top-k result set S. Similar to PathSim-baseline (Algorithm 4.2), before the pruning steps, we still need to first derive the candidate set. Compared with the baseline algorithm, the pruning-based algorithm at most checks the same number of candidates with the overhead to calculate the upper bounds. In practice, a great number of candidates can be pruned, and therefore the performance can be enhanced.

Algorithm 4.2 (PathSim-Baseline) Vector-Matrix Multiplication Based Path Concatenation

```
Input: Query x_i, Commuting Matrix M_{\mathcal{D}}, Diagonal Vector D, top-k K
Output: Top-k List SortList
 1: CandidateSet = \emptyset;
 2: foreach y_k \in M_{\mathcal{D}}.neighbors(x_i) do
       foreach x_j \in M_{\mathcal{D}}^T.neighbors(y_k) do
          CandidateSet = CandidateSet \cup \{x_i\};
 5: List = \emptyset;
 6: foreach x_i \in CandidateSet do
       value = 2 * M_{\mathcal{D}}(i,:)M_{\mathcal{D}}(j,:)^T/(D(i) + D(j));
       List.update(x_j, value, K);
 9: List.sort();
10: SortList = List.topk(K);
11: return SortList;
```

Algorithm 4.3 (PathSim-Pruning) Cluster-based Top-k Search on Path Concatenation

```
Input: Query x_i, Commuting matrix M_{\mathcal{D}}^T, Feature clusters \{R_u\}_{u=1}^U, Target clusters \{C_v\}_{v=1}^V, Diagonal
    vector D, top-k K.
Output: Top-k list S.
 1: Set CandidateSet = x_i.neighbors.neighbors;
 2: S = \emptyset;
 3: Sort clusters in \{C_v\}_{v=1}^V according to upper bound of s(x_i, C_v);
 4: foreach C_v with decreasing order do
       if the upper bound of s(x_i, C_v) < min(S) then
 6:
          break;
 7:
       else
          foreach x_j \in C_v and x_j \in CandidateSet do
 8:
 9:
             if the upper bound of s(x_i, x_j) < min(S) then
                continue;
10:
11:
                s(x_i, x_j) = \frac{2M \mathcal{P}^{(x_i,:)}(M \mathcal{P}^{(x_j,:)})^T}{D(x_i) + D(x_j)};
12:
                Insert x_i into S;
13:
14: return S:
```

Experiments show that PathSim-Pruning can significantly improve the query processing speed comparing with the baseline algorithm, without affecting the search quality.

4.4 Multiple Meta-Paths Combination

In Section 4.3, we presented algorithms for similarity search using single meta-path. Now, we present a solution to combine multiple meta-paths. Formally, given r round trip meta-paths from Type A back to Type A, $\mathcal{P}_1, \mathcal{P}_2, \ldots, \mathcal{P}_r$, and their corresponding relation matrix M_1, M_2, \ldots, M_r , with weights w_1, w_2, \ldots, w_r specified by users, the combined similarity between objects $x_i, x_j \in A$ are defined as: $s(x_i, x_j) = \sum_{l=1}^r w_l s_l(x_i, x_j)$, where $s_l(x_i, x_j) = \frac{2M_l(i,j)}{M_l(i,i) + M_l(j,j)}$.

Example 4.1. (Multiple meta-paths combination for venue similarity search) Following the motivating example in the introduction section, Table 4.4 shows the results of combining two meta-paths $\mathcal{P}_1 = VPAPV$ and $\mathcal{P}_2 = VPTPV$ with different weights specified by w_1 and w_2 , for query "DASFAA."

Table 4.4: Top-5 similar venues to "DASFAA" using multiple meta-paths.

Rank	$w_1 = 0.2, w_2 = 0.8$	$w_1 = 0.5, w_2 = 0.5$	$w_1 = 0.8, w_2 = 0.2$
1	DASFAA	DASFAA	DASFAA
2	Data Knowl. Eng.	DEXA	DEXA
3	CIKM	CIKM	WAIM
4	EDBT	Data Knowl. Eng.	CIKM
5	Inf. Syst.	EDBT	APWeb

The reason why we need to combine several meta-paths is that, each meta-path provides a unique angle (or a unique feature space) to view the similarity between objects, and the ground truth may be a cause of different factors. Some useful guidance of the weight assignment includes: longer meta-path utilize more remote relationships and thus should be assigned with a smaller weight, such as in P-PageRank and SimRank; and, meta-paths with more important relationships should be assigned with a higher weight. For automatically determining the weights, users could provide training examples of similar objects to learn the weights of different meta-paths using learning algorithms.

We now evaluate the quality of similarity measure generated by combined meta-paths, according to their performance for clustering tasks in the "four-area" dataset. First, two meta-paths for the venue type, namely, VAV and VTV (short for VPAPV and VPTPV), are selected and their linear

combinations with different weights are considered. Second, two meta-paths with the same basic path but different lengths, namely AVA and $(AVA)^2$, are selected and their linear combinations with different weights are considered. The clustering accuracy measured by NMI for conferences and authors is shown in Table 4.5, which shows that the combination of multiple meta-paths can produce better similarity than the single meta-path in terms of clustering accuracy.

Table 4.5: Clustering accuracy for PathSim for meta-path combinations on the "four-area" dataset.

w_1	0	0.2	0.4	0.6	0.8	1
w_2	1	0.8	0.6	0.4	0.2	0
VAV; VTV	0.7917	0.7936	0.8299	0.8587	0.8123	0.8116
$AVA; (AVA)^2$	0.6091	0.6219	0.6506	0.6561	0.6508	0.6501

4.5 Experiments

To show the effectiveness of the PathSim measure and the efficiency of the proposed algorithms, we use the bibliographic networks extracted from DBLP and Flickr in the experiments.

We use the DBLP dataset downloaded in Nov. 2009 as the main test dataset. It contains over 710K authors, 1.2M papers, and 5K venues (conferences/journals). After removing stopwords in paper titles, we get around 70K terms appearing more than once. This dataset is referred as the full-DBLP dataset. Two small subsets of the data (to alleviate the high computational costs of P-PageRank and SimRank) are used for the comparison with other similarity measures in effectiveness: (1) the DBIS dataset, which contains all the 464 venues and top-5000 authors from the database and information system area; and (2) the four-area dataset, which contains 20 venues and top-5000 authors from 4 areas: database, data mining, information retrieval and artificial intelligence [98], and cluster labels are given for all the 20 venues and a subset of 1713 authors.

For additional case studies, we construct a Flickr network from a subset of the Flickr data, which contains four types of objects: images, users, tags, and groups. Links exist between images and users, images and tags, and images and groups. We use 10,000 images from 20 groups as well as their related 664 users and 10284 tags appearing more than once to construct the network.

Table 4.6: Case study of five similarity measures on query "PKDD" on the DBIS dataset.

Rank	P-PageRank	SimRank	RW	PRW	PathSim
1	PKDD	PKDD	PKDD	PKDD	PKDD
2	KDD	Local Pattern Detection	KDD	Local Pattern Detection	ICDM
3	ICDE	KDID	ICDM	DB Support for DM Appl.	SDM
4	VLDB	KDD	PAKDD	Constr. Min. & Induc. DB	PAKDD
5	SIGMOD	Large-Scale Paral. DM	SDM	KDID	KDD
6	ICDM	SDM	TKDE	MCD	DMKD
7	TKDE	ICDM	SIGKDD Expl.	Pattern Detection & Disc.	SIGKDD Expl.
8	PAKDD	SIGKDD Expl.	ICDE	RSKD	Knowl. Inf. Syst.
9	SIGIR	Constr. Min. & Induc. DB	SEBD	WImBI	J IIS
10	CIKM	TKDD	CIKM	Large-Scale Paral. DM	KDID

4.5.1 Effectiveness

Comparing PathSim with other measures When a meta-path $\mathcal{P} = (\mathcal{P}_l \mathcal{P}_l^{-1})$ is given, other measures such as random walk (RW) and pairwise random walk (PRW) can be applied to the same meta-path, and P-PageRank and SimRank can be applied to the sub-network extracted from \mathcal{P} . For example, for the meta-path VPAPV (VAV in short) for finding venues sharing the same set of authors, the bipartite graph M_{CA} , derived from the relation matrix corresponding to VPA can be used in both P-PageRank and SimRank algorithms. In our experiments, the damping factor for P-PageRank is set as 0.9 and that for SimRank is 0.8.

First, a case study is shown in Table 4.6, which is applied to the *DBIS* dataset, under the meta-path *VAV*. One can see that for query "PKDD" (short for "Principles and Practice of Knowledge Discovery in Databases," a European data mining conference), P-PageRank favors the venues with higher visibility, such as KDD and several well-known venues; SimRank prefers more concentrated venues (*i.e.*, a large portion of publications goes to a small set of authors) and returns many not well-known venues such as "Local Pattern Detection" and KDID; RW also favors highly visible objects such as KDD, but brings in fewer irrelevant venues due to that it utilizes merely one short meta-path; PRW performs similar to SimRank, but brings in more not so well-known venues due to the short meta-path it uses; whereas PathSim returns the venues in both the area and the reputation similar to PKDD, such as ICDM and SDM.

We then labeled top-15 results for 15 queries from the venues in the *DBIS* dataset (i.e., SIG-MOD, VLDB, ICDE, PODS, EDBT, DASFAA, KDD, ICDM, PKDD, SDM, PAKDD, WWW, SIGIR, TREC and APWeb), to test the quality of the ranking lists given by 5 measures. We label

Table 4.7: Comparing the accuracy of top-15 query results for five similarity measures on the DBIS dataset measured by nDCG.

	P-PageRank	SimRank	RW	PRW	PathSim
Accuracy	0.5552	0.6289	0.7061	0.5284	0.7446

each result object with a relevance score at one of the three levels: 0 (non-relevant), 1 (somewhat relevant), and 2 (very relevant). Then we use the measure nDCG (i.e., Normalized Discounted Cumulative Gain, with the value between 0 and 1, the higher the better) [51] to evaluate the quality of a ranking algorithm by comparing its output ranking results with the labeled ones (Table 4.7). The results show that PathSim gives the best ranking quality in terms of human intuition, which is consistent with the previous case study.

Table 4.8: Top-10 similar authors to "Christos Faloutsos" under different meta-paths on the *full-DBLP* dataset.

(a)	Path:	APA
(00)	I COLII.	2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

Rank	Author
1	Christos Faloutsos
2	Spiros Papadimitriou
3	Jimeng Sun
4	Jia-Yu Pan
5	Agma J. M. Traina
6	Jure Leskovec
7	Caetano Traina Jr.
8	Hanghang Tong
9	Deepayan Chakrabarti
10	Flip Korn

(b) Path: APVPA

Rank	Author
1	Christos Faloutsos
2	Jiawei Han
3	Rakesh Agrawal
4	Jian Pei
5	Charu C. Aggarwal
6	H. V. Jagadish
7	Raghu Ramakrishnan
8	Nick Koudas
9	Surajit Chaudhuri
10	Divesh Srivastava

Semantic meanings of different meta-paths As we pointed out, different meta-paths give different semantic meanings, which is one of the reasons that similarity definitions in homogeneous networks cannot be applied directly to heterogeneous networks. Besides the motivating example in the introduction section, Table 4.8 shows the author similarity under two scenarios for author Christos Faloutsos: co-authoring papers and publishing papers in the same venues, represented by the meta-paths APA and APVPA respectively. One can see that the first path returns co-authors

who have strongest connections with Faloutsos (i.e., students and close collaborators) in DBLP, whereas APVPA returns those publishing papers in the most similar venues.

Table 4.9: Top-10 similar venues to "SIGMOD" under meta-paths with different lengths on the full-DBLP dataset.

(a) Path: $(VPAPV)^2$ (b) Path: $(VPAPV)^4$ (c) Path: $(VPAPV)^{\infty}$

Rank	Venue	Score	Rank	Venue	Score	Rank	Venue	Score
1	SIGMOD	1	1	SIGMOD	1	1	SIGMOD	1
2	VLDB	0.981	2	VLDB	0.997	2	AAAI	0.9999
3	ICDE	0.949	3	ICDE	0.996	3	ESA	0.9999
4	TKDE	0.650	4	$ ext{TKDE}$	0.787	4	ITC	0.9999
5	SIGMOD Record	0.630	5	SIGMOD Record	0.686	5	STACS	0.9997
6	IEEE Data Eng. Bul.	0.530	6	PODS	0.586	6	PODC	0.9996
7	PODS	0.467	7	KDD	0.553	7	NIPS	0.9993
8	ACM Trans. DB Sys.	0.429	8	CIKM	0.540	8	Comput. Geom.	0.9992
9	EDBT	0.420	9	IEEE Data Eng. Bul.	0.532	9	ICC	0.9991
10	CIKM	0.410	10	J. Comp. Sys. Sci.	0.463	10	ICDE	0.9984

The impact of path length The next interesting question is how the length of meta-path impacts the similarity definition. Table 4.9 shows an example of venues similar to "SIGMOD" with three meta-paths, using exactly the same basic meta-path, but with different repeating times. These meta-paths are $(VPAPV)^2$, $(VPAPV)^4$ and its infinity form (global ranking-based similarity). Note that in $(VPAPV)^2$, two venues are similar if they share many similar authors who publish papers in the same venues; while in $(VPAPV)^4$, the similarity definition of those venues will be further relaxed, namely, two venues are similar if they share many similar authors who publish papers in similar venues. Since venue type only contains 5K venues, we are able to get the full materialization relation matrix for $(VPAPV)^2$. $(VPAPV)^4$ is obtained using meta-path concatenation from $(VPAPV)^2$. The results are summarized in Table 4.9, where longer paths gradually bring in more remote neighbors, with higher similarity scores, and finally, it degenerates into global ranking comparison. Through this study, one can see that a meta-path with relatively short length is good enough to measure similarity, where a long meta-path may even reduce the quality.

4.5.2 Efficiency Comparison

The time complexity for SimRank is $O(KN^2d^2)$, where K is the number of iterations, N is the total number of objects, and d is the average neighbor size; the time complexity for calculating P-PageRank for one query is O(KNd), where K, N, d has the same meaning as in SimRank; whereas the time complexity for PathSim using PathSim-baseline for single query is O(nd), where n < N is the number of objects in the target type, d is the average degree of objects in target type for partial relation matrix $M_{\mathcal{P}_l}$. The time complexity for RW and PRW are the same as PathSim. We can see that similarity measure only using one meta-path is much more efficient than those also using longer meta-paths in the network (e.g., SimRank and P-PageRank).

Now we compare the pruning power of PathSim-pruning vs. PathSim-baseline by considering two factors: the size of the neighbors of a query (Fig. 4.3) and the density of the partial commuting matrix $M_{\mathcal{P}}$ (Fig. 4.4). 500 queries are randomly chosen for two meta-paths (VPAPV and $(VPAPV)^2$, denoted as VAV and VAVAV for short), and the execution time is averaged with 10 runs. The results show that the execution time for PathSim-baseline is almost linear to the size of the candidate set, and the improvement rate for PathSim-pruning is larger for queries with more neighbors, which requires more calculation for exact dot product operation between a query vector and candidate vectors. Also, the denser that the commuting matrix corresponding to the partial meta-path (M_{VPAPV} in comparison with M_{VPA}), the greater the pruning power. The improvement rates are 18.23% and 68.04% for the two meta-paths.

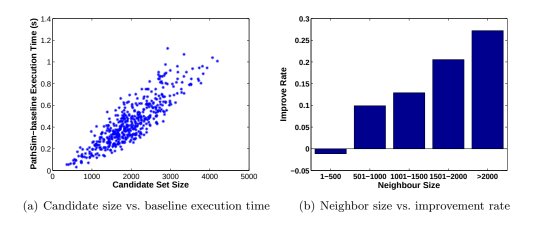


Figure 4.3: Efficiency study for queries with different neighbor size under meta-path VAV on the full-DBLP dataset based on 500 queries.

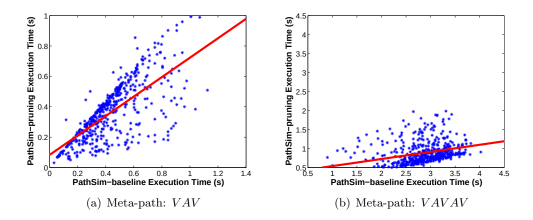


Figure 4.4: Pruning power denoted by the slope of the fitting line under two meta-paths for type conference on the *full-DBLP* dataset. Each dot represents a query under the indicated meta-path.

4.5.3 Case study on Flickr network

In this case study, we show that to retrieve similar images for a query image, one can explore links in the network rather than the content information. Let "I" represent images, "T" tags that associated with each image, and "G" groups that each image belongs to. Two meta-paths are used and compared. One is ITI, which means common tags are used by two images at evaluation of their similarity. The results are shown in Figure 4.5. The other is ITIGITI, which means tags similarities are further measured by their shared groups, and two images can be similar even if they do not share many exact same tags as long as these tags are used by many images of the same groups. One can see that the second meta-path gives better results than the first, as shown in Figure 4.6, where the first image is the input query. This is likely due to that the latter meta-path provides additional information related to image groups, and thus improves the similarity measure between images.

4.6 Related Work

Similarity measure has been widely studied in categorical, numerical, or mix-type data sets, such as cosine similarity defined on two vectors, Jaccard coefficient on two sets, and Euclidean distance on two numerical data points. Based on the traditional similarity measures, a recent study [114]

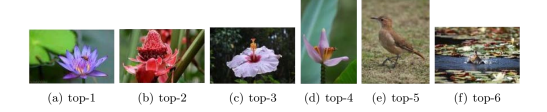


Figure 4.5: Top-6 images in Flickr network under meta-path ITI.

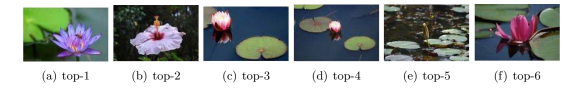


Figure 4.6: Top-6 images in Flickr network under meta-path ITIGITI.

proposes an efficient top-k similarity pair search algorithm, top-k-join, in relational database, which only considers similarity between tuples. Also widely studied are k nearest neighbor search in spatial data [60] and other high dimensional data [15], which aims at finding top-k nearest neighbors according to similarities defined on numerical features. However, these similarity definitions cannot be applied to networks.

Similarity measures defined on homogeneous networks emerged recently. Personalized PageR-ank [53] is an asymmetrical similarity measure that evaluates the probability starting from object x to visit object y by randomly walking on the network with restart. More discussions on how to scale the calculation for online queries are in [38, 108], etc., and how to derive top-k answers efficiently is studied in [42]. SimRank [52] is a symmetric similarity measure defined on homogeneous networks, which can also be directly applied to bipartite networks. The intuition behind SimRank is propagating pairwise similarity to their neighboring pairs. Due to its computational complexity, there are many follow-up studies (e.g., [68]) on speeding up such calculations. SCAN [117] measures similarity of two objects by comparing their immediate neighbor sets.

ObjectRank [5] and PopRank [82] first noticed that heterogeneous relationships could affect the random walk, and assigned different propagation factors to each type of object relationship to either derive a revised version of P-PageRank (ObjectRank) or a global PageRank (PopRank). However, such solutions only give one particular combination of all the possible meta-paths using the fixed

weights determined by the damping factor and propagation factors between different types. In our PathSim definition, users can freely specify the meta-paths they are interested in and assign any weight to them. Random walk style similarity search is not adopted in PathSim, which overcomes the disadvantage of returning highly ranked objects rather than similar peers.

4.7 Discussions

In this study, we assume that users know how to choose meta-path. In practice, there are several ways for a user to select the best meta-path or meta-path combinations. First, a user can make a choice based on her interest and domain knowledge. Second, she can have several experimental trials, such as those done in Section 4.5, and choose the best one according to her intuition. Third, she can label a small portion of data according to specific applications. For example, one can label similar objects or rank them, and then train the best meta-path(s) and their weights by some learning algorithms. By doing so, one can automatically choose appropriate meta-paths as well as the associated weights, and make the similarity search adaptable to different application scenarios. One recent work [115] follows this direction, which can identify different similarity search intentions by learning the weights for different meta-paths. The problem on how to choose and weight different meta-paths is similar to the feature selection process in machine learning. In-depth study for a systematic solution is left as a future research task.

4.8 Conclusion

In this chapter, we have introduced a novel and practical notion of meta-path-based similarity for heterogeneous information networks. We comparatively and systematically examine different semantics of similarity measures in such networks and introduce a new meta-path-based similarity measure to find similar objects of the same type in such networks. Meta-paths give users flexibility to choose different meta-paths and their combinations based on their applications. Moreover, we propose a new similarity measure, PathSim, under this framework, which produces overall better similarity qualities than the existing measures. Since meta-paths can be arbitrarily given, it is unrealistic to fully materialize all the possible similarity results given different meta-paths and their

combinations. However, online calculation requires matrix multiplication, which is time consuming especially when the vector and matrix are not sparse. Therefore, we proposed an efficient solution that partially materializes several short meta-paths and then applies online concatenation and combination among paths to give the top-k results for a query. Experiments on real data sets show the effectiveness of the similarity measure and the efficiency of our method. The framework of meta-path-based similarity search in networks can be enhanced in many ways, e.g., weight learning for different meta-paths, which may help provide accurate similarity measures in real systems and discover interesting relationships among objects.

Chapter 5

Meta-Path-Based Relationship Prediction

In Chapter 4, we introduced a meta-path-based similarity measure, PathSim, for heterogeneous information networks. The concept of meta-path serves not only as a basis for similarity search but also as a key for mining and learning general heterogeneous networks with an arbitrary network schema, because this notion provides a way to guide us to systematically build link-based features. In this chapter, we examine a new mining task, relationship prediction in heterogeneous information networks, by exploring meta-path-based features.

5.1 Overview

Link prediction, that is, predicting the emergence of links in a network based on certain current or historical network information, has been a popular theme of research in recent years, thanks to the popularity of social networks and other online systems. The applications of link prediction range from social networks to biological networks, as it addresses the fundamental question of whether a link will form between two nodes in the future. Most of the existing link prediction methods [66, 45, 111, 67, 63] are designed for homogeneous networks, in which only one type of objects exists in the network. For example, in a friendship network or a co-author network, a user may like to predict possible new friendship between two persons or new co-authorship between two authors, based on the existing links in a network.

In the real world, most networks are heterogeneous, where multiple types of objects and links exist. In such networks, objects are connected by different types of relationships. Objects are connected together not only by immediate links, but also by more sophisticated relationships that follow some meta-path-based relations. Here we extend the *link prediction* problem in homogeneous information networks to the *relationship prediction* problem in heterogeneous information networks,

where a relationship could be an immediate link or a path instance following some meta-path.

Many real-world problems can be considered as relationship prediction tasks, such as citation prediction in a bibliographic network, product recommendation in an e-commerce network, and online advertisement click prediction in an online system-based network.

The heterogeneity of objects and links makes it difficult to use well-known topological features in homogeneous networks for algorithmic design. For example, the number of the common neighbors is frequently used as a feature for link prediction in homogeneous networks. However, the neighbors of an object in a heterogeneous network often are of different types, and a simple measure like the number of shared neighbors cannot reflect this heterogeneity.

We thus propose a meta-path-based relationship prediction framework to overcome this difficulty. Instead of treating objects and links of different types equally or extracting homogeneous subnetworks from the original network, we propose a meta-path-based topological feature framework for heterogeneous networks. The goal is to systematically define the relations between objects encoded in different paths using the meta structure of these paths, that is, the meta-paths.

Two case studies using the meta-path-based relationship prediction framework are presented in this chapter. The first is on co-authorship prediction in the DBLP network, whereas the second proposes a novel prediction model that can predict when a relationship is going to built in a given heterogeneous information network.

5.2 Meta-Path-Based Relationship Prediction Framework

Different from traditional link prediction tasks for homogeneous information networks, in a heterogeneous information network scenario, it is necessary to specify which type of relationships to predict. The relationship to be predicted is called the target relation and can be described using a meta-path. For example, the relation co-authorship can be described as a meta-path A - P - A. Moreover, in order to build an effective prediction model, one need to examine how to construct the meta-path-based topological features between two objects for each potential relationship. In this section, we first examine how to systematically build topological feature space using meta-paths, and then present a supervised prediction framework where the meta-path-based topological measures are used as features.

5.2.1 Meta-Path-based Topological Feature Space

Topological features, also known as structural features, reflect the essential connectivity properties for pairs of objects. Topological feature-based link prediction aims at inferring the future connectivity by leveraging the current connectivity of the network. There are some frequently used topological features defined in homogeneous networks, such as the number of common neighbors, preferential attachment [10, 78], and $katz_{\beta}$ [57]. We first review several commonly used topological features in homogeneous networks, and then propose a systematic meta-path-based methodology to define topological features in heterogeneous networks.

Existing Topological Features

We introduce several well-known and frequently used topological features in homogeneous networks. For more topological features, the readers can refer to [66].

- Common neighbors. Common neighbors is defined as the number of common neighbors shared by two objects a_i and a_j , namely $|\Gamma(a_i) \cap \Gamma(a_j)|$, where $\Gamma(a)$ is the notation for neighbor set of the object a and $|\cdot|$ denotes the size of a set.
- Jaccard's coefficient. Jaccard's coefficient is a measure to evaluate the similarity between two neighbor sets, which can be viewed as the normalized number of common neighbors, namely $\frac{|\Gamma(a_i)\cap\Gamma(a_j)|}{|\Gamma(a_i)\cup\Gamma(a_j)|}.$
- $Katz_{\beta}$. $Katz_{\beta}$ [57] is a weighted summation of counts of paths between two objects with different lengths, namely $\sum_{l=1}^{\infty} \beta^{l} |path_{a_{i},a_{j}}^{\langle l \rangle}|$, where β^{l} is the damping factor for the path with length l.
- *PropFlow*. In a recent study [67], a random walk-based measure PropFlow is proposed to measure the topological feature between two objects. This method assigns the weighs to each path (with fixed length *l*) using the products of proportions of the flows on the edges.

One can see that most of the existing topological features in homogeneous networks are based on neighbor sets or paths between two objects. However, as there are multi-typed objects and multityped relations in heterogeneous networks, the neighbors of an object could belong to multiple types, and the paths between two objects could follow different meta-paths and indicate different relations. Thus, it is necessary to design a more complex strategy to generate topological features in heterogeneous networks.

Meta-path-based Topological Features

To design topological features in heterogeneous networks, we first define the topology between two objects using meta-paths, and then define measures on a specific topology. In other words, a meta-path-based topological feature space is comprised of two parts: the meta-path-based topology and the measure functions that quantify the topology.

Meta-path-based topology As introduced in Chapter 4, a meta-path is a path defined over a network schema and denotes a composition relation over a heterogeneous network. By checking the existing topological features defined in a homogeneous network, we can find that both the neighbor set-based features and path-based features can be generalized in the heterogeneous information network, by considering paths following different meta-paths. For example, if we treat each type of neighbors separately and extend the immediate neighbors to n-hop neighbors (i.e., the distance between one object and its neighbors are n), the common neighbor feature between two objects then becomes the count of paths between the two objects following different meta-paths. For path-based features, such as $Katz_{\beta}$, it can be extended as a combination of paths following different meta-paths, where each meta-path defines a unique topology between objects, representing a special relation.

Meta-paths between two object types can be obtained by traversing the graph of network schema, using standard traversal methods such as the BFS (breadth-first search) algorithm. As the network schema is a much smaller graph compared with the original network, this stage is very fast. We can enumerate all the meta-paths between two object types by setting a length constraint. For example, in order to predict co-authorship in the DBLP network, we extract all the meta-paths within a length constraint, say 4, starting and ending with the author type A. The meta-paths between authors up to length 4 are summarized in Table 5.1, where the semantic meaning of each relation denoted by each meta-path are given in the second column.

Table 5.1: Meta-paths under length 4 between authors in the DBLP network.

Meta-path	Semantic Meaning of the Relation
A-P-A	a_i and a_j are co-authors
$A-P \rightarrow P-A$	a_i cites a_j
$A - P \leftarrow P - A$	a_i is cited by a_j
A-P-V-P-A	a_i and a_j publish in the same venues
A-P-A-P-A	a_i and a_j are co-authors of the same authors
A-P-T-P-A	a_i and a_j write the same topics
$A - P \to P \to P - A$	a_i cites papers that cite a_j
$A - P \leftarrow P \leftarrow P - A$	a_i is cited by papers that are cited by a_j
$A - P \rightarrow P \leftarrow P - A$	a_i and a_j cite the same papers
$A - P \leftarrow P \rightarrow P - A$	a_i and a_j are cited by the same papers

Measure functions on meta-paths Once the topologies given by meta-paths are determined, the next stage is to propose measures to quantify these meta-paths for pairs of objects. Here we list four measures along the lines of topological features in homogeneous networks. They are path count, normalized path count, random walk, and symmetric random walk, defined as follows. Additional measures can be proposed, such as pairwise random walk mentioned in Chapter 4.

- Path count. Path count measures the number of path instances between two objects following a given meta-path R, denoted as PC_R . Path count can be calculated by the products of adjacency matrices associated with each relation in the meta-path.
- Normalized path count. Normalized path count is to discount the number of paths between two objects in the network by their overall connectivity, and is defined as $NPC_R(a_i, a_j) = \frac{PC_R(a_i, a_j) + PC_{R-1}(a_j, a_i)}{Z_R(a_i, a_j)}$, where R^{-1} denotes the inverse relation of R, $Z_R(a_i, a_j)$ is some normalization factor. For example, PathSim [99] is a special case of normalized path count, where $Z_R(a_i, a_j) = PC_R(a_i, a_i) + PC_R(a_j, a_j)$ for symmetric R's.
- Random walk. Random walk measure along a meta-path is defined as $RW_R(a_i, a_j) = \frac{PC_R(a_i, a_j)}{PC_R(a_i, \cdot)}$, where $PC_R(a_i, \cdot)$ denotes the total number of paths following R starting with a_i , which is a natural generalization of PropFlow [67].
- Symmetric random walk. Symmetric random walk considers the random walk from two directions along the meta-path, and defined as $SRW_R(a_i, a_j) = RW_R(a_i, a_j) + RW_{R^{-1}}(a_j, a_i)$.

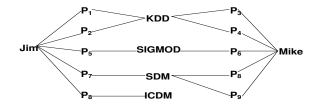


Figure 5.1: An example of path instances between two authors following A-P-V-P-A

Taking the example in Figure 5.1, we show the calculation of these measures. Let R denote the relation represented by meta-path A - P - V - P - A. It is easy to check it is symmetric, i.e., $R = R^{-1}$. Let J denote Jim, and M denote Mike. We can see that $PC_R(J, M) = 7$, $NPC_R(J, M) = \frac{7+7}{7+9} = 7/8$ (under PathSim), $RW_R(J, M) = 1/2$, $RW_R(M, J) = 7/16$, and $SRW_R(J, M) = 15/16$.

For each meta-path, we can apply any measure functions on it and obtain a unique topological feature. So far, we have provided a systematic way to define the topological features in heterogeneous networks, which is a large space defined over $topology \times measure$. These meta-path-based topological features can serve a good feature space for mining and learning tasks, such as relationship prediction.

5.2.2 Supervised Relationship Prediction Framework

The supervised learning framework is summarized in Figure 5.2. Generally, given a past time interval $T_0 = [t_0, t_1)$, we want to use the topological features extracted from the aggregated network in the time period T_0 , to predict the relationship building in a future time interval, say $T_1 = [t_1, t_2)$. In the **training stage**, we first sample a set of object pairs in T_0 , collect their associated topological features represented as \mathbf{x} 's in T_0 , and record relationship building facts between them represented as \mathbf{y} 's in the future interval T_1 . A training model is then built to learn the best coefficients associated with each topological feature by maximizing the likelihood of relationship building. In the **test stage**, we apply the learned coefficients to the topological features for the test pairs, and compare the predicted relationship with the ground truth. Note that, the test stage may have different past interval T'_0 and future interval T'_1 as in the training stage, but we require they have the same lengths as the intervals in the training stage, namely using the same amount of past information to predict the same length of future.

For most of the existing link prediction studies, the tasks are predicting whether a new link will

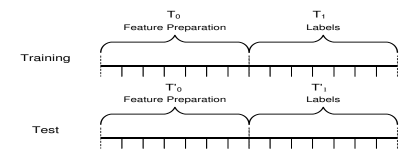


Figure 5.2: Supervised framework for relationship prediction.

appear in the future. In other words, y is a binary variable and is usually modeled as following Bernoulli distribution. While in a more general case, y can be variables related to any reasonable value of the relationship for a pair of objects. For example, in order to predict when a relationship is going to be built, y could be modeled a positive real value following exponential distribution; in order to predict the frequency of a relationship (e.g., how many times two authors are going to collaborate), y could be modeled as a non-negative integer following Poisson distribution. Then statistical models can be built based on the distribution assumptions of y, such as logistic regression model for binary variables and generalized linear model for more sophisticated assumptions.

Two case studies of relationship prediction are shown in the following sections, both of which follow the supervised relationship prediction framework, but with different purposes and thus different assumptions on the response variable y.

5.3 Co-authorship Prediction

For the first case study, we study the problem of co-authorship prediction in the DBLP bibliographic network, that is, whether two authors are going to collaborate in a future interval for the first time. In this case, the target relation for prediction is co-authorship relation, which can be described using meta-path A - P - A. For the topological features, we study all the meta-path listed in Table 5.1 other than A - P - A and all the measures listed in the last section.

We next introduce the relationship prediction model which models the probability of coauthorship between two authors as a function of topological features between them. Given the training pairs of authors, we first extract the topological features for them, and then build the prediction model to learn the weights associated with these features.

5.3.1 The Co-authorship Prediction Model

In order to predict whether two authors are going to collaborate in a future interval, denoted as y, we use the logistic regression model as the prediction model. For each training pair of authors $\langle a_{i_1}, a_{i_2} \rangle$, let \mathbf{x}_i be the (d+1)-dimensional vector including constant 1 and d topological features between them, and y_i be the label of whether they will be co-authors in the future $(y_i = 1)$ if they will be co-authors, and 0 otherwise), which follows Bernoulli distribution with probability p_i $(P(y_i = 1) = p_i)$. The probability p_i is modeled as follows:

$$p_i = \frac{e^{\mathbf{x}_i \boldsymbol{\beta}}}{e^{\mathbf{x}_i \boldsymbol{\beta}} + 1}$$

where β is the d+1 coefficient weights associated with the constant and each topological feature. We then use standard MLE (Maximum Likelihood Estimation) to derive $\hat{\beta}$, that maximizes the likelihood of all the training pairs:

$$L = \prod_{i} p_i^{y_i} (1 - p_i)^{(1 - y_i)}$$

In the test stage, for each candidate author pair, we can predict whether they will collaborate according to $P(y_{test} = 1) = \frac{e^{\mathbf{x}_{test}\hat{\boldsymbol{\beta}}}}{e^{\mathbf{x}_{test}\hat{\boldsymbol{\beta}}}+1}$, where \mathbf{x}_{test} is the (d+1)-dimensional vector including constant 1 and d topological features between the candidate pair.

5.3.2 Experiments

It turns out that the proposed meta-path-based topological features can improve the co-authorship prediction accuracy compared with the baselines that only use homogeneous object and link information.

We consider three time intervals for the DBLP network, according to the publication year associated with each paper: $T_0 = [1989, 1995]$, $T_1 = [1996, 2002]$, and $T_2 = [2003, 2009]$. For the training stage, we use T_0 as the past time interval, and T_1 as the future time interval, which is denoted as $T_0 - T_1$ time framework. For the test stage, we consider the same time framework $T_0 - T_1$ for most of the studies, and consider $T_1 - T_2$ time framework for the query-based case study.

Let an author pair be $\langle a_i, a_j \rangle$, we call a_i the source author, and a_j the target author. Two

sets of source authors are considered. The first set is comprised of highly productive authors, who has published no less than 16 papers in the past time interval; and the second set is comprised of less productive authors, with between 5 and 15 publications. The target authors are selected if they are 2-hop co-authors or 3-hop co-authors of the source author. In all, we have four labeled datasets: (1) the highly productive source authors with 2-hop target authors (denoted as HP2hop); (2) the highly productive source authors with 3-hop target authors (denoted as HP3hop); (3) the less productive source authors with 2-hop target authors (denoted as LP2hop); and (4) the less productive source authors with 3-hop target authors (denoted as LP3hop).

To evaluate the prediction accuracy, two measures are used. The first measure is the classification accuracy rate (accuracy) for binary prediction under the cut-off score as 0.5, and the second one is the area under ROC (receiver operating characteristic) curve [19], which is denoted as AUC.

Overall Accuracy

We first compare the heterogeneous topological features with the homogeneous ones. For the heterogeneous topological features, we use path count measure for 9 meta-paths (denoted as heterogeneous PC) listed in Table 5.1 (not including the target relation itself); for homogeneous topological features, we use (1) the number of common co-authors, (2) the rooted PageRank [66] with restart probability $\alpha=0.2$ for the co-author sub-network, and (3) the number of paths between two authors of length no longer than 4, disregarding their different meta-paths (denoted as homogeneous PC). The rooted PageRank measure is only calculated for the HP3hop dataset, due to its inefficiency in calculation for large number of authors. The comparison results are summarized in Figure 5.3 and Table 5.2. We can see that the heterogeneous topological feature beats the homogeneous ones in all the four datasets, which validates the necessity to consider the different meta-paths separately in heterogeneous networks. We also notice that, in general the co-authorship for highly productive authors is easier to predict than less productive authors, by looking at the overall prediction accuracy on the two groups of source authors. Finally, we can see that the prediction accuracy is higher when the target authors are 3-hop co-authors, which means the collaboration between closer authors in the network is more affected by information that is not available from network topology.

Second, we compare different measures proposed for heterogeneous topological features: (1)

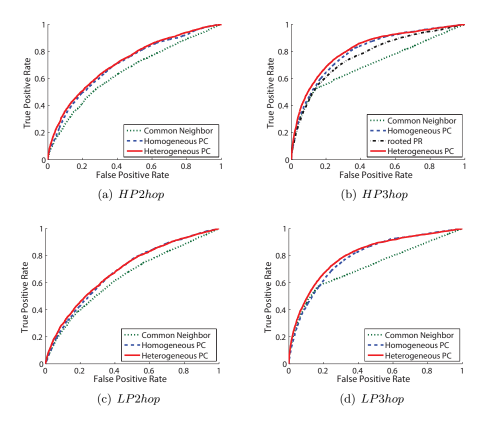


Figure 5.3: Homogeneous features vs. heterogeneous *Path Count* feature. Heterogeneous feature beats homogeneous features in all of the datasets, which is more significant on 3-hop datasets, where topological features play a more important role for co-authorship prediction.

the path count (PC), (2) the normalized path count (NPC, i.e., PathSim in our case), (3) the random walk (RW), (4) the symmetric random walk (SRW), and (5) the hybrid features of (1)-(4) (hybrid). It turns out that in general we have (see Figure 5.4): (1) all the heterogeneous features beat the homogeneous features (common neighbor is denoted as PC1, and homogeneous PC is denoted as PCSum); (2) the normalized path count beats all the other three individual measures; and (3) the hybrid feature produces the best prediction accuracy.

Case Study

For the case study, we first show the learned importance for each topological feature in deciding the relationship building in DBLP, and then show the predicted co-author relationships for some source author in a query mode.

First, we show the learned importance for all the 9 meta-paths with NPC measure, as NPC

Table 5.2: Homogeneous vs. heterogeneous topological features.

Dataset	Topological features	Accuracy	AUC
	common neighbor	0.6053	0.6537
U Doh on	homogeneous PC	0.6433	0.7098
HP2hop	heterogeneous PC	0.6545	0.7230
	common neighbor	0.6589	0.7078
HP3hop	homogeneous PC	0.6990	0.7998
	rooted PageRank	0.6433	0.7098
	heterogeneous PC	0.7173	0.8158
	common neighbor	0.5995	0.6415
LP2hop	homogeneous PC	0.6154	0.6868
	heterogeneous PC	0.6300	0.6935
	common neighbor	0.6804	0.7195
LP3hop	homogeneous PC	0.6901	0.7883
	heterogeneous PC	0.7147	0.8046

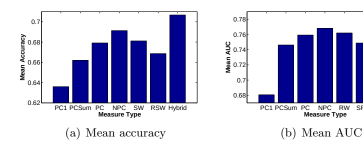


Figure 5.4: Average accuracy over 4 datasets for different features.

is the best measure for co-author relationship prediction overall. We show the p-value for the feature associated with each meta-path under Wald test and their significance level in Table 5.3. From the results, we can see that for the HP3hop dataset, the shared co-authors, shared venues, shared topics and co-cited papers for two authors all play very significant roles in determining their future collaboration(s). For the asymmetric meta-paths that represent the asymmetric relations, such as citing and cited relations between authors, they have different impacts in determining the relationship building. For example, for a highly productive source author, the target authors citing her frequently are more likely to be her future co-authors than the target authors being cited by her frequently.

Second, we study the predicted co-authors for some source author as queries. Note that, pre-

Table 5.3: Significance of meta-paths with Normalized Path Count measure for HP3hop dataset.

Meta-path	<i>p</i> -value	Significance level ¹
$A-P \rightarrow P-A$	0.0378	**
$A - P \leftarrow P - A$	0.0077	***
A-P-V-P-A	1.2974e-174	****
A-P-A-P-A	1.1484e-126	****
A-P-T-P-A	3.4867e-51	****
$A-P \rightarrow P \rightarrow P-A$	0.7459	
$A - P \leftarrow P \leftarrow P - A$	0.0647	*
$A-P o P \leftarrow P-A$	9.7641e-11	****
$A - P \leftarrow P \rightarrow P - A$	0.0966	*

¹ *: p < 0.1; **: p < 0.05; ***: p < 0.01, ****: p < 0.001

dicting co-authors for a given author is an extremely difficult task, as we have too many candidate target authors (3-hop candidates are used), while the number of real new relationships are usually quite small. Table 5.4 shows the top-5 predicted co-authors in time interval T_2 (2003-2009) using the $T_0 - T_1$ training framework, for both the proposed hybrid topological features and the shared co-author feature. We can see that, the results generated by heterogeneous features has a higher accuracy compared with the homogeneous one.

Table 5.4: Top-5 predicted co-authors for Jian Pei in 2003-2009.

Rank	Hybrid heterogeneous features	# of shared authors as features
1	Philip S. Yu	Philip S. Yu
2	Raymond T. Ng	Ming-Syan Chen
3	Osmar R. Zaïane	Divesh Srivastava
4	Ling Feng	Kotagiri Ramamohanarao
5	David Wai-Lok Cheung	Jeffrey Xu Yu

^{*} Bold font indicates true new co-authors of Jian Pei in the period of 2003-2009.

5.4 Relationship Prediction with Time

Traditional link prediction studies have been focused on asking **whether** a link will be built in the future, such as "whether two people will become friends?" However, in many applications, it may be more interesting to predict **when** the link will be built, such as "what is the probability that

two authors will co-write a paper within 5 years," and "by when will a user in Netflix rent the movie *Avatar* with 80% probability?"

In this section, we study the problem of predicting the relationship building time between two objects, such as, when two authors will collaborate for the first in the future, based on the topological structure in a heterogeneous network, by investigating the citation relationship between authors in the DBLP network. First, we introduce the concepts of target relation and topological features for the problem encoded in meta-paths [99]. Then, a generalized linear model (GLM) [36] based supervised framework is proposed to model the relationship building time. In this framework, the building time for relationships are treated as independent random variables conditional on their topological features, and their expectation is modeled as a function of a linear predictor of the extracted topological features. We propose and compare models with different distribution assumptions for relationship building time, where the parameters for each model are learned separately.

5.4.1 Topological Features for Author Citation Relationship Prediction

In the author citation relationship prediction problem, the target relation is $A - P \to P - A$, which is short for $A \xrightarrow{write} P \xrightarrow{cite} P \xrightarrow{write^{-1}} A$, and describes the citation relation between authors. In general, for a target relation $R_T = \langle A, B \rangle$, any meta-paths starting with type A and ending with type B other than the target relation itself can be used as the topological features for predicting new relationships. These meta-paths can be obtained by traversing on the network schema, for example, using BFS (breadth-first search). By reasoning the dynamics of a relationship building, we are in particular considering three forms of relations as topological features:

- 1. $AR_{sim}AR_TB$, where R_{sim} is a similarity relation defined between type A and R_T is the target relation. The intuition is that if a_i in type A is similar to many a_k 's in type A that have relationships with b_j in type B, then a_i is likely to build a relationship with b_j in the future.
- 2. $AR_TBR_{sim}B$, where R_T is the target relation, and R_{sim} is a similarity relation between type B. The intuition is that if a_i in type A has relationships with many b_k 's in type B that are similar to b_j in type B, then a_i is likely to build a relationship with b_j in the future.

Table 5.5: Meta-paths denoting similarity relations between authors.

Meta-path	Semantic Meaning of the Relation
A-P-A	a_i and a_j are co-authors
A-P-V-P-A	a_i and a_j publish in the same venues
A-P-A-P-A	a_i and a_j are co-authors of the same authors
A-P-T-P-A	a_i and a_j write the same topics
$A - P \rightarrow P \leftarrow P - A$	a_i and a_j cite the same papers
$A - P \leftarrow P \rightarrow P - A$	a_i and a_j are cited by the same papers

3. AR_1CR_2B , where R_1 is some relation between A and C and R_2 is some relation between C and B. The intuition is that if a_i in type A has relationships with many c_k 's in type C that have relationships with b_j in type B, then a_i is likely to build a relationship with b_j in the future. Note that the previous two forms are special cases of this one, which can be viewed as triangle connectivity property.

For topological features, we confine similarity relations R_{sim} and other partial relations R_1 and R_2 to those that can be derived from the network using meta-paths. Moreover, we only consider similarity relations that are symmetric.

Taking the author citation relation, which is defined as $A - P \to P - A$, as the target relation, we consider 6 author-author similarity relations defined in Table 5.5. For each similarity relation, we can concatenate the target relation in its left side or in its right side. We then have 12 topology features with the form $AR_{sim}AR_TB$ and $AR_TBR_{sim}B$ in total. Besides, we consider the concatenation of "author-cites-paper" relation $(A - P \to P)$ and "paper-cites-author" relation $(P \to P - A)$ into $(A - P \to P \to P - A)$, as well as all the 6 similarity relations listed in Table 5.5, which can be viewed as the form of AR_1CR_2B themselves. Now we have 19 topological features in total.

For each type of the meta-paths, we illustrate a concrete example to show the possible relationship building in Figure 5.5. In Figure 5.5(a), authors a_1 and a_2 are similar, as they publish papers containing similar terms, and a_2 cites papers published by a_3 . In the future, a_1 is likely to cite papers published by a_3 as well, since she may follow the behavior of her fellows. In Figure 5.5(b), author a_1 cites a_2 , and a_2 and a_3 are cited by common papers together (p_5, p_6, p_7) . Then a_1 is likely to cite a_3 in the future, as she may cite authors similar to a_2 . In Figure 5.5(c), a_1 and

 a_2 publish in the same venue, then a_1 is likely to cite a_2 in the future as they may share similar interests if publishing in the same conference.

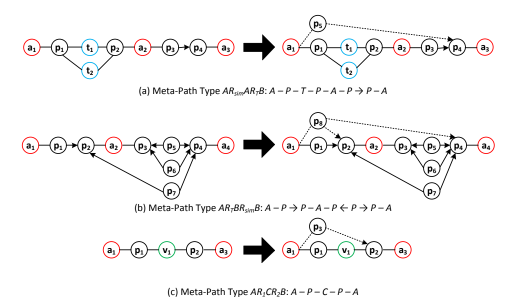


Figure 5.5: Feature meta-path illustration for author citation relationship prediction.

By varying the similarity relations and partial relations, we are able to generate other topological features in arbitrary heterogeneous networks.

Without loss of generality, we use the count of path instances as the default measure. Thus, each meta-path corresponds to a measure matrix. For a single relation $R \in \mathcal{R}$, the measure matrix is just the adjacency matrix of the sub-network extracted by R. Given a composite relation, the measure matrix can be calculated by the matrix multiplication of the partial relations.

In Figure 5.5(a), the count of path instances between a_1 and a_3 following the given meta-path is 2, which are:

(1)
$$a_1 - p_1 - t_1 - p_2 - a_2 - p_3 \rightarrow p_4 - a_3$$
, and

(2)
$$a_1 - p_1 - t_2 - p_2 - a_2 - p_3 \rightarrow p_4 - a_3$$
.

In Figure 5.5(b), the count of path instances between a_1 and a_4 following the given meta-path is 3, which are:

(1)
$$a_1 - p_1 \rightarrow p_2 - a_2 - p_3 \leftarrow p_5 \rightarrow p_4 - a_4$$
,

(2)
$$a_1 - p_1 \rightarrow p_2 - a_2 - p_3 \leftarrow p_6 \rightarrow p_4 - a_4$$
, and

(3)
$$a_1 - p_1 \rightarrow p_2 - a_2 - p_2 \leftarrow p_7 \rightarrow p_4 - a_4$$
.

In Figure 5.5(c), the count of path instances between a_1 and a_3 following the given meta-path is 1, which is:

$$(1) a_1 - p_1 - v_1 - p_2 - a_3.$$

Measures for different meta-paths have different scales. For example, longer meta-paths usually have more path instances due to the adjacency matrix multiplication. We will normalize the measure using Z-score for each meta-path.

5.4.2 The Relationship Building Time Prediction Model

We now propose the generalized linear model-based prediction model, which directly models the relationship building time as a function of topological features, and provides methods to learn the coefficients of each topological feature, under different assumptions for relationship building time distributions. After that, we introduce how to use the learned model to make inferences.

We model the relationship building time prediction problem in a supervised learning framework. In the **training stage**, we first collect the topological features \mathbf{x}_i in the history interval $T_0 = [t_0, t_1)$ for each sampled object pair $\langle a_i, b_i \rangle$, where types of a_i and b_i are $\tau(a_i) = A$ and $\tau(b_i) = B$. Then, we record their relative first relationship building time $y_i = t_i - t_1$, if t_i is in the future training interval $T_1 = [t_1, t_2)$; record the building time $y_i \geq t_2 - t_1$, if no new relationship has been observed in T_1 . Note that in the training stage, we are only given limited time to observe whether and when two objects will build their relationship, it is very possible that two objects build their relationship after t_2 , which needs careful handling in the training model. A generalized linear model (GLM) based relationship building time model is introduced, and the goal is to learn the best coefficients associated with each topological feature that maximize the current observations of the relationship building time. In the **test stage**, we apply the learned coefficients of the topological features to the test pairs, and compare the predicted relationship building time with the ground truth.

Different from the existing link prediction task, in the training stage, we are collecting relationship building time y_i for each training pair, which is a variable ranging from 0 to ∞ , rather than a binary value denoting whether there exists a link in the future interval. Similarly, in the test stage, we are predicting the relationship building time y_i for test pairs that range from 0 to ∞ , rather than predicting whether the link exists or not in the given future interval.

The Generalized Linear Model Framework

The main idea of generalized linear model (GLM) [36] is to model the expectation of random variable Y, E(Y), as some function ("link function") of the linear combination of features, that is, $X\beta$, where X is the observed feature vector, and β is the coefficient vector. Then the goal is to learn β according to the training data set using maximum likelihood estimation. Under different distribution assumptions for Y, usually from the exponential family, E(Y) has different forms of parameter set, and the link functions are with different forms too. Note that the most frequently used Least-Square regression and logistic regression are special cases of GLM, where Y follows Gaussian distribution and Bernoulli distribution respectively.

Suppose we have n training pairs for the target relation $\langle A, B \rangle$. We denote each labeled pair as $r_i = \langle a_i, b_i \rangle$, and y_i as the observed relative relationship building time in the future interval. We denote \mathbf{X}_i as the d dimensional topological feature vector extracted for a_i and b_i in the historical interval plus a constant dimension.

Distributions for Relationship Building Time

The first issue of the prediction model is to select a suitable distribution for the relationship building time. Intuitively, a relationship building between two objects can be treated as an event, and we are interested in when this event will happen.

Let Y be the relationship building time relative to the beginning of the future interval $(y_i = t_i - t_1)$, and let T be the length of future training interval. For training pairs, Y has the observations in $[0,T)\cup\{T^+\}$ in a continuous case, and $\{0,1,2,\ldots,T-1,T^+\}$ in a discrete case, where $y=T^+$ means no event happens within the future training interval. For testing pairs, Y has the observations in $[0,\infty)$ in a continuous case, and nonnegative integers in a discrete case.

We consider three types of distributions for relationship building time, namely exponential, Weibull and geometric distribution. For each of the distribution assumptions over y_i , we set up the models separately.

The first distribution is **exponential distribution**, which is the most frequently used distribution in modeling waiting time for an event. The probability density function of an exponential

distribution is:

$$f_Y(y) = \frac{1}{\theta} \exp\{-\frac{y}{\theta}\}\tag{5.1}$$

where $y \ge 0$, and $\theta > 0$ is the parameter denoting the *mean waiting time* for the event. The cumulative distribution function is:

$$F_Y(y) = Pr(Y \le y) = 1 - \exp\{-\frac{y}{\theta}\}\$$
 (5.2)

The second distribution is **Weibull distribution**, which is a generalized version of exponential distribution and is another standard way to model the waiting time of an event. The probability density function of a Weibull distribution is:

$$f_Y(y) = \frac{\lambda y^{\lambda - 1}}{\theta^{\lambda}} \exp\{-(\frac{y}{\theta})^{\lambda}\}$$
 (5.3)

where $y \geq 0$, and $\theta > 0$ and $\lambda > 0$ are two parameters related to mean waiting time for the event and hazard of happening of the event along with the time. λ is also called the shape parameter, as it affects the shape of probability function. When $\lambda > 1$, it indicates an increasing happening rate along the time (if an event does not happen at an early time, it is getting higher probability to happen at later time); and when $\lambda < 1$, it indicates a decreasing happening rate along the time (if an event does not happen at an early time, it is getting less possible in happening in later time). Note that when $\lambda = 1$, Weibull distribution becomes exponential distribution with mean waiting time as θ , and the happening rate does not change along the time. The cumulative distribution function is:

$$F_Y(y) = Pr(Y \le y) = 1 - \exp\{-\left(\frac{y}{\theta}\right)^{\lambda}\}$$
(5.4)

The third distribution is the **geometric distribution**, which is a distribution that models how many times of failures it needs to take before the first-time success. As in our case, the time of failure is the discrete time that we need to wait before a relationship is built. The probability mass function of a geometric distribution is:

$$Pr(Y = k) = (1 - p)^k p$$
 (5.5)

where k = 0, 1, 2, ..., and p is the probability of the occurrence of the event at each discrete time. The cumulative distribution function is:

$$Pr(Y \le k) = 1 - (1 - p)^{k+1} \tag{5.6}$$

In our case, each relationship building is an independent event, and each relationship building time Y_i is an independent random variable, following the same distribution family, but with different parameters. With the distribution assumptions, we build relationship building time prediction models in the following.

Model under Exponential and Weibull Distribution Note that, as exponential distribution is a special case of Weibull distribution (with $\lambda = 1$), we only discuss prediction model with Weibull distribution.

In this case, we assume relationship building time Y_i for each training pair is independent of each other, following the same Weibull distribution family with the same shape parameter λ , but with different mean waiting time parameters θ_i . Namely, we assume that different relationships for the target relation share the same trend of hazard happening along with the time, but with different expectation in building time. Under this assumption, we can evaluate the expectation for each random variable Y_i as $E(Y_i) = \theta_i \Gamma(1 + \frac{1}{\lambda})$. We then use the link function $E(Y_i) = \exp\{-\mathbf{X}_i\boldsymbol{\beta}\}\Gamma(1+\frac{1}{\lambda})$, that is $\log \theta_i = -\beta_0 - \sum_{j=1}^d X_{i,j}\beta_j = -\mathbf{X}_i\boldsymbol{\beta}$, where β_0 is the constant term. Then we can write the log-likelihood function:

$$\log L = \sum_{i=1}^{n} (f_Y(y_i | \theta_i, \lambda) I_{\{y_i < T\}} + P(y_i \ge T | \theta_i, \lambda) I_{\{y_i \ge T\}})$$

where $I_{\{y_i < T\}}$ and $I_{\{y_i \ge T\}}$ are indicator functions, which equals to 1 if the predicate holds, or 0 otherwise. It is easy to see that the log-likelihood function includes two parts: if y_i is observed in the future interval, we use its real density in the function; otherwise, we are only able to use the probability of $y_i \ge T$ in the function.

By plugging in $\log \theta_i = -\mathbf{X}_i \boldsymbol{\beta}$, we can get the log-likelihood with parameters β and λ :

$$LL_W(\boldsymbol{\beta}, \lambda) = \sum_{i=1}^n I_{\{y_i < T\}} \log \frac{\lambda y_i^{\lambda - 1}}{e^{-\lambda \mathbf{X}_i \beta}} - \sum_{i=1}^n (\frac{y_i}{e^{-\mathbf{X}_i \beta}})^{\lambda}$$
 (5.7)

where LL_W denotes the log-likelihood function under **W**eibull distribution. We refer this model as Weibull model.

Model under Geometric Distribution In this case, we assume relationship building time Y_i for each training pair is independent of each other, following the same geometric distribution family, but with different success probability p_i . Under this assumption, we can evaluate the expectation for each random variable Y_i as $E(Y_i) = \frac{1-p_i}{p_i}$. We then let $E(Y_i) = \exp\{-\mathbf{X}_i\boldsymbol{\beta}\}$, i.e., $\log \frac{1-p_i}{p_i} = -\mathbf{X}_i\boldsymbol{\beta}$. The log-likelihood function is then:

$$LL_G(\boldsymbol{\beta}) = \sum_{i=1}^{n} (Pr(Y_i = y_i)I_{\{y_i < T\}} + P(y_i \ge T)I_{\{y_i \ge T\}})$$

$$= \sum_{i=1}^{n} \left(-I_{\{y_i < T\}}(-\mathbf{X}_i \boldsymbol{\beta}) + (y_i + 1)(-\mathbf{X}_i \boldsymbol{\beta} - \log(e^{-\mathbf{X}_i \boldsymbol{\beta}} + 1)) \right)$$
(5.8)

We refer this model as geometric model.

The learning of the models is becoming an optimization problem, which aims at finding $\hat{\beta}$ and other parameters (e.g., $\hat{\lambda}$ in the Weibull model) that maximize the log-likelihood. As there are no closed form solutions for Eqs. (5.7) and (5.8), we use standard Newton-Raphson method to derive the update formulas, which are based on the first derivative and second derivative (Hessian matrix) of the log-likelihood function.

Model Inference

Once the parameters such as β and λ are learned from the training data set through MLE, we can apply the model to the test pairs of objects, as long as their topological features in the historical network are given. Let the learned parameter values be $\hat{\beta}$ and $\hat{\lambda}$ for β and λ , and let the topological feature vector for the test pairs be \mathbf{X}_{test} (with constant 1 as the first dimension), we now consider three types of questions people may be interested in for the new relationship building time, and provide the solutions in the following.

1. Whether a new relationship between two test objects will be built within t years?

This question is equal to the query for the probability $Pr(y_{test} \leq t)$, which can be evaluated by plugging in the MLE estimators to derive the distribution parameters. Note that for traditional link prediction tasks, t should be the same as the length of training interval. For our task, t can be any nonnegative values. For Weibull model, we have:

$$\hat{\theta}_{test} = \exp\{-\mathbf{X}_{test}\hat{\beta}\}$$

$$Pr(y_{test} \le t) = 1 - \exp\{-(\frac{t}{\hat{\theta}_{test}})^{\hat{\lambda}}\}$$
(5.9)

For geometric model, we have:

$$\hat{p}_{test} = \frac{1}{\exp\{-\mathbf{X}_{test}\hat{\beta}\} + 1}$$

$$Pr(y_{test} \le t) = 1 - (1 - \hat{p}_{test})^{t+1}$$
(5.10)

2. What is the average relationship building time for two test objects?

This is simply the query for $E(Y_{test})$. Using the same estimators for $\hat{\theta}_{test}$ and \hat{p}_{test} as above, we can have the estimator for $E(Y_{test})$ as $E(Y_{test}) = \hat{\theta}_{test}\Gamma(1+\frac{1}{\hat{\lambda}})$ for Weibull model, where $\Gamma(\cdot)$ is the Gamma function, and $E(Y_{test}) = \frac{1-\hat{p}_{test}}{\hat{p}_{test}}$ for geometric model.

3. The quantile: by when a relationship will be built with a probability α ?

This is equal to query for the solution of $F_Y(y_{test}) = \alpha$, and we can get answers as $y_{test} = \hat{\theta}_{test}(-\log(1-\alpha))^{\frac{1}{\lambda}}$ for Weibull model, and $y_{test} = \max\{\frac{\log(1-\alpha)}{\log(1-\hat{p}_{test})} - 1, 0\}$ for geometric model. When $\alpha = 0.5$, the quantile is just the median.

5.4.3 Experiments

We select a subset of authors in the DBLP bibliographic network, who published more than 5 papers in top conferences in the four areas¹ that are related to data mining between years 1996 and 2000 ($T_0 = [1996, 2000]$). The total number of the author set is 2721. Then we sampled 7000

¹Data Mining: KDD, PKDD, ICDM, SDM, PAKDD; Database: SIGMOD Conference, VLDB, ICDE, PODS, EDBT; Information Retrieval: SIGIR, ECIR, ACL, WWW, CIKM; and Artificial Intelligence: NIPS, ICML, ECML, AAAI, IJCAI.

pairs of authors in the form of $\langle a_i, a_j \rangle$ that a_i did not cite a_j in T_0 , but have citation relationship between year 2001 and 2009 ($T_1 = [2001, 2009]$ and T = 9) as positive samples; and we sampled another 7000 pairs of authors that have no citation relationship during either T_0 or T_1 . The citation relationship is defined if a_i cites papers written by a_j published before year 2000. Note that, we have this time constraint for papers as we want to infer citation relationship via the historical network. 19 topological features introduced in Section 5.4.1 are calculated for each training pair. The first (relative) time of the citation relationship is recorded for each pair of authors; and if there is no citation relationship between them in T_1 , the time is recorded as a value bigger than 9.

Experimental Setting

In order to show the power of using time-involved model in relationship prediction, we use logistic regression [84] (denoted as logistic) that is frequently used in binary link prediction tasks as the baseline. Note that, the output of the logistic regression is a probability denoting whether a relationship will be built in T_1 for each test pair. In our models, the output is the parameter set for the distribution of the relationship building time, from which we can infer much more information rather than a simple probability. We denote our models with different distribution assumptions as GLM_geo , GLM_exp , and GLM_weib respectively.

To compare the four models, we use two sets of measures to evaluate the effectiveness of each model. First, we measure the effectiveness according to the predicted probability for each relationship. We define the *accuracy* of the relationship prediction as the ratio between the number of correctly predicted relationship (under the *cut-off* 0.5) and the total number of the test pairs. Also, another frequently used measure AUC (the area under ROC curve) is used to compare the accuracy.

Second, we directly compare the predicted time with the ground truth, among our proposed models. Mean absolute error (MAE) that is the mean of the absolute error between predicted relationship building time and the ground truth is used. Also, we use the ratio of the relationships that occur in some confidence interval derived from the models as another measure to test the accuracy of the predicted time. Note that, relationships yet to happen are not considered in these two measures.

Prediction Power Study

We now compare our time-involved models with the baseline logistic regression, using the first set of measures.

We test the generality power for different models, namely, when the training future interval is not equal to the test future interval $(T^{train} \neq T^{test})$. On one hand, we may want to know the probability of relationship building within each year in the training interval $(T^{test} < T^{train})$; on the other hand, we may want to infer longer term probability given a short term training interval $(T^{test} > T^{train})$. We show the two cases in Tables 5.6 and 5.7. Note that, since logistic regression can only output the probability when $T^{test} = T^{train}$, we use the same predicted probability for different test intervals. In Table 5.6, we fix the training interval with length $T^{train} = 9$, namely, $T_1^{train} = [2001, 2009]$, and vary the test intervals with length from 1 to 4. We can see that when T^{test} is small, time-involved models can give much better prediction accuracy, especially in terms of the measure accuracy. In other words, time-involved models carry more information in telling the probability of relationship building in finer time periods. In Table 5.7, we fix the test interval with length $T^{test} = 9$ and vary the training intervals with length from 2 to 5. We can see that, time-involved models can better utilize the short term training than logistic regression, and output better prediction results for longer term relationship building behavior. It is interesting to note that by using the measure AUC, which does not require users to specify a cut-off value in the predicted probabilities, the performance of logistic regression is still comparable with other models. This is due to AUC only uses the ranking order of the predicted values, while accuracy requires that the absolute values of the predicted probabilities are also correct.

Table 5.6: Prediction generalization power comparison: $T^{test} < T^{train}$ and $T^{train} = 9$.

	$T^{test} = 1$		T^{test}	$T^{test} = 2$		$T^{test} = 3$		$T^{test} = 4$	
	Accuracy	AUC	Accuracy	AUC	Accuracy	AUC	Accuracy	AUC	
logistic	0.7106	0.7619	0.7246	0.7535	0.7669	0.7347	0.7349	0.7731	
GLM-geo	0.9284	0.7626	0.8436	0.7532	0.7829	0.7657	0.7347	0.7696	
GLM-exp	0.9290	0.7553	0.8442	0.7464	0.7821	0.7569	0.7328	0.7603	
GLM-weib	0.9287	0.7273	0.8441	0.7452	0.7826	0.7559	0.7334	0.7597	

In all, for time-involved model, it contains more information and can answer different questions

Table 5.7: Prediction generalization power comparison: $T^{test} > T^{train}$ and $T^{test} = 9$.

	$T^{train} = 2$		$T^{train} = 3$		$T^{train} = 4$		$T^{train} = 5$	
	Accuracy	AUC	Accuracy	AUC	Accuracy	AUC	Accuracy	AUC
logistic	0.5157	0.7810	0.5379	0.7805	0.5599	0.7841	0.5952	0.7896
GLM-geo	0.5942	0.7910	0.6209	0.7926	0.6366	0.7902	0.6522	0.7982
GLM-exp	0.5015	0.7802	0.5214	0.7833	0.6709	0.7841	0.7143	0.7870
GLM-weib	0.7081	0.7816	0.7021	0.7832	0.7002	0.7833	0.7103	0.7862

and with strong generalization power. Logistic regression can only answer the question of whether a relationship will happen or not, given a fixed time interval. However, if we are asking more, it fails in most of the scenarios.

Time Prediction Accuracy Study

We now evaluate the predicted time using different time-involved models. Here, we use the predicted median time as the predicted time. Table 5.8 shows the MAE (mean average error) between the predicted median time and the ground truth under different training and test intervals. It turns out that GLM-exp has the lowest error. Also, both GLM-exp and GLM-weib perform even better using shorter interval as training, whereas GLM-geo has the opposite behavior, that is, longer term of training leads to better performance. Note that, we only calculate the error for the relationships indeed happen in the test interval.

In Table 5.9, we infer different confidence intervals from the predicted relationship building time distribution, and test the ratio of the true relationship in different confidence intervals. A confidence interval (range) rather than a simple value, say the median time, can give users a better view of the relationship building time. It is shown that *GLM-exp* and *GLM-weib* has a higher ratio of giving correct confidence intervals for the true relationship building time, especially when using a small confidence interval. This is very useful in practice as they can give tight bound estimations.

Case Studies

To better understand the output of our model, we now show a case study of predicting when the citation relationship will be build for "Philip S. Yu" with other candidates. The model is

Table 5.8: MAE of predicted time and the ground truth.

	$T^{train} = 5, T^{test} = 9$	$T^{train} = 9, T^{test} = 9$
GLM-geo	4.9883	4.7219
$\overline{GLM\text{-}exp}$	2.7774	3.0685
GLM-weib	3.1025	3.1692

Table 5.9: Ratio of the true relationship occurring in different confidence intervals: $T^{test} = 9$.

	25%-75%		10%-90%		0%-80%	
	$T^{train} = 9$	$T^{train} = 5$	$T^{train} = 9$	$T^{train} = 5$	$T^{train} = 9$	$T^{train} = 5$
GLM-geo	0.5489	0.5336	0.8936	0.8947	0.9650	0.9743
GLM-exp	0.7167	0.7246	0.8619	0.8634	0.9880	0.9889
GLM-weib	0.7278	0.7314	0.8680	0.8686	0.9884	0.9896

trained by GLM-weib using a training interval of 9 years ($T_1^{train} = [2001, 2009]$), with the learned parameter $\lambda = 0.9331$, slightly less than 1, which means the citation relationship has a higher hazard happening at an earlier time. The ground truth of the citation building time, and the predicted median, mean, 25% quantile and 75% quantile for several test pairs are shown in Table 5.10. It can be seen that the predicted median and confidence interval are very suggestive for predicting the true citation relationship building time. For those authors whose predicted being cited time is significantly different from the ground truth, in-depth studies may be needed. For example David Maier is a prolific researcher in database area, and by intuition as well as suggested by the model, Philip should cite him. However, the ground truth says otherwise. Furthermore, this function can be used to recommend authors to any author in DBLP for citation purpose.

Table 5.10: Case studies of relationship building time prediction.

$\overline{a_i}$	a_j	Ground Truth	Median	Mean	25% quant.	75% quant.
Philip S. Yu	Ling Liu	1	2.2386	3.4511	0.8549	4.7370
Philip S. Yu	Christian Jensen	3	2.7840	4.2919	1.0757	5.8911
Philip S. Yu	C. Lee Giles	0	8.3985	12.9474	3.2450	17.7717
Philip S. Yu	Stefano Ceri	0	0.5729	0.8833	0.2214	1.2124
Philip S. Yu	David Maier	9+	2.5675	3.9581	0.9920	5.4329
Philip S. Yu	Tong Zhang	9+	9.5371	14.7028	3.6849	20.1811
Philip S. Yu	Rudi Studer	9+	9.7752	15.0698	3.7769	20.6849

For the above model, the learned *top-4* most important topological features with the highest coefficients are:

- 1. A P T P A, that is, if two authors are very similar in terms of writing similar topics, they tend to cite each other;
- 2. $A-P \leftarrow P \rightarrow P-A$, that is, if two authors are very similar in terms of being frequently co-cited by the common papers, they tend to cite each other;
- 3. $A P A P \rightarrow P A$, that is, an author tends to cite the authors that are frequently cited by her co-authors;
- 4. $A P T P A P \rightarrow P A$, that is, if two authors are similar in terms writing similar topics, they tend to cite the same authors.

These topological features provide insightful knowledge for people in understanding the citation relationship building between authors.

5.5 Related Work

The link prediction problem has been first studied on homogeneous networks. Early work mainly studies unsupervised methods [1, 66], namely they propose different similarity measures according to either topological structures of the networks or proximity of object attributes that are consistent with the link appearance in the future. Later, supervised methods that are able to combine different features with different coefficients via training data sets are proposed by different studies [45, 111, 67]. A recent study [63] has discussed the link prediction problem when the network is not fully observed and thus is modeled in a probabilistic way. A survey in link prediction can be found in [40]. In this paper, we extend the link prediction problem to the more general heterogeneous networks, by extending link prediction to relationship prediction and exploring the topological features in such scenarios.

Recently, some studies [21, 65] propose frequent graph pattern mining-based methodology to detect graph evolution rules, which provides some clues for proposing new topological features

in the network for link prediction. However, the focus on the two papers are still on homogeneous networks, and they have not considered how different frequent evolution patterns affect the link formation speed yet. That is, this methodology cannot answer the "when" problem of link formation.

Another line of study similar to our problem is the link prediction task in relational data [84, 106], as relational data also involves different types of objects and complex relationships between objects. However, these studies have a focus different from our research. As in [84], they study feature selection in a relational environment using relational languages, and feed these features into supervised link prediction models; for [106], their goal is to model the relational data via a probabilistic model. In this chapter, we aim at designing a model for relationship building, either whether or when, by systematically exploring the topological features in heterogeneous networks.

The general setting of link prediction task is set by Liben-Nowell and Kleinberg [66], which is to predict whether a link between two existing objects will be added to the network during the time interval $[t, t + \Delta t]$ given the snapshot of the network at time t. In other words, the task has not considered the issue when a link will appear in this time interval. Recently, several studies have considered the extension on usage of time. In [109], a methodology that assigns weights to events and edges according to their appearing time is proposed, which produces better link prediction accuracy by using more time information in the feature side. In [49], a time series model is proposed to predict the frequency of repeated links in networks. In comparison to these studies, our research focuses on the new relationship prediction and aims at modeling the relationship building in the future.

In all, in this chapter, we extend the traditional link prediction in homogeneous networks into relationship prediction in the more complex heterogeneous networks. we build a framework for general relationship prediction in heterogeneous networks by systematically extracting meta pathbased topological features, and study whether or when the relationship will happen in the future.

5.6 Conclusion

In this chapter, we study the problem of relationship prediction in heterogeneous information networks. In comparison with traditional homogeneous networks, heterogeneous networks contain multiple types of objects and links. Two case studies using the meta-path-based relationship prediction framework are presented in this chapter. The first is on co-authorship prediction in the DBLP network, whereas the second proposes a novel prediction model that can predict when a relationship is going to built in a given heterogeneous information network. Experiments on the DBLP bibliographic network show that by considering heterogeneous topological features, the relationship prediction accuracy can be significantly improved, and the model using hybrid features that have combined different meta-paths and different measures gives the best overall performance. Furthermore, the learned significance for each topological feature can provide better understanding of the relationship building mechanism in such networks.

Part III

User-Guided Relation Strength-Aware Mining

Chapter 6

Relation Strength-Aware Clustering with Incomplete Attributes

A heterogeneous information network contains multiple types of objects as well as multiple types of links, indicating different sorts of interactions among these objects. The heterogeneity of network model brings rich semantic information for mining. It also raises the issue of selecting the right type of information for different mining purposes. For mining different kinds of knowledge, it is desirable to automatically learn the right information encoded in the network, with limited guidance from users. In this chapter, we study a special case of such problems: cluster objects in a network, with user-provided attribute set and relations from the original network schema.

6.1 Overview

The rapid emergence of online social media, e-commerce, and cyber-physical systems brings the necessity to study them with the model of heterogeneous networks in which objects (*i.e.*, nodes) are of different types, and links among objects correspond to different relations, denoting different interaction semantics. In addition, an object is usually associated with some attributes. For example, in a *YouTube* social media network, the object types may include videos, users, and comments; links between objects correspond to different relations, such as publish and like relations between users and videos, post relation between users and comments, and friendship and subscribe relations between users; and attributes may include user's location, the length of video's clips, the number of views, and comments.

Such kinds of heterogeneous information networks are ubiquitous and determining their underlying clustering structures has many interesting applications. For example, clustering objects (e.g., customers, products, and comments) in an online shopping network such as eBay is helpful for customer segmentation in product marketing; and clustering objects (e.g., people, groups, books,

and posts) in an online social network such as Facebook is helpful for voter segmentation in political campaigns.

The clustering task brings two new challenges in such scenarios. First, an object may contain only partial or even no observations for a given attribute set that is critical to determine their cluster labels. That is, a pure attribute-based clustering algorithm cannot correctly detect these clusters. Second, although links have been frequently used in networks to detect clusters [29, 72, 3, 103] in recent research, we consider a more challenging scenario in which the links are of different types and interpretations, each of which may have its own level of semantic importance in the clustering process. That is, a pure link-based clustering without any guidance from attribute specification could fail to meet user demands.

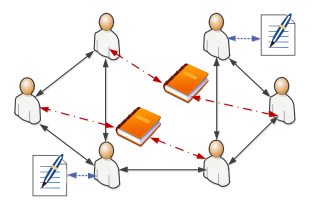


Figure 6.1: A motivating example on clustering political interests in social information networks.

Figure 6.1 shows a toy social information network extracted from a political forum containing users, blogs written by users, books liked by users, and friendship between users. Now suppose we want to cluster users in the network according to their political interests, using the text attributes in user profiles, blogs and books, as well as the link information between objects. On one hand, since not all the users listed their political interests in their profiles, we cannot judge their political interests simply according to the text information contained in their profiles directly. On the other hand, without specifying the purpose of clustering, we cannot decide which types of links to use for the clustering; shall we use the friendship links to detect the social communities, or the user-like-book links to detect the reading groups, or a mix of them? Obviously, to solve such clustering tasks, we need to use both the incomplete attribute information as well as the link information of different types with the awareness of their importance weights. In our example, in order to discover

a user's political interests, we need to learn which link types are more important for our purpose of clustering, among the relationships between her and blogs, books, and her friends.

Recently, some studies [133, 75, 91, 118, 105, 70] show that the combination of attribute and link information in a network can improve the clustering quality. However, none of them has addressed the two challenges simultaneously. Some of them rely on a complete attribute space and the clustering result is considered as a trade-off between attribute-based measures and link-based measures. Moreover, none of the current studies has examined the issue that different types of links have different importance in determining a clustering with a certain purpose.

Here we explore the interplay between different types of links and the specified attribute set in the clustering process, and design a comprehensive and robust probabilistic clustering model for heterogeneous information networks.

6.2 The Relation Strength-Aware Clustering Problem Definition

As defined before, a heterogeneous information network $G = (\mathcal{V}, \mathcal{E}, W)$ is modeled as a directed graph, where each node in the network corresponds to an object (or an event) in real life, and each link corresponds to a relationship between the linked objects. Associated with each link, there is a binary or positive value, denoting its input weight.

Attributes are associated with objects, such as the location of a user, the text description of a book, the text information of a blog, and so on. In this setting, we consider attributes across all different types of objects as a collection of attributes for the network, denoted as $\mathcal{X} = \{X_1, \dots, X_T\}$, in which we are interested only in a subset for a certain clustering purpose. Each object $v \in \mathcal{V}$ contains a subset of the attributes, with observations denoted as $v[X] = \{x_{v,1}, x_{v,2}, \dots, x_{v,N_{X,v}}\}$, where $N_{X,v}$ is the total number of observations of attribute X attached with object v. Note that, some attributes can be shared by different types of objects, such as the text and the location attribute; while some other attributes are unique for a certain type of objects, such as the time length for a video clip. We use \mathcal{V}_X to denote the object set that contains attribute X.

6.2.1 The Clustering Problem

The goal of the clustering problem is to map every object in the network into a unified hidden space, that is, a soft clustering, according to the user-specified subset of attributes in the network, with the help of links from different types.

There are several new challenges for clustering objects in this new scenario. First, the attributes are usually **incomplete** for an object: the attributes specified by a user may be only partially or even not contained in an object type; and the values for these attributes could be missing even if the attribute type is contained in the object type. Moreover, the incompleteness of the data cannot be easily handled by interpolation: the observations for each attribute could be a set or a bag of values, and the neighbors for an object are from different types of objects, which may not be helpful for predicting the missing data. For example, it is impossible to get a user's blog via interpolating techniques. Therefore, none of the existing clustering algorithms that purely based on attribute space can solve the clustering problem in this scenario.

Second, with the awareness that links play a critical role to propagate the cluster information among objects, another challenge is that **different link types** have different semantic meanings and therefore have different strengths in the process of passing cluster information around. In other words, while it is clear that the existence of links between nodes is indicative of clustering similarity, it is also important to understand that different link types may have a different level of importance in the clustering process. In the example of clustering political interests illustrated in Figure 6.1, we expect a higher importance of the relation user-like-book than the relation friendship in deciding the cluster membership of a user. Thus, we need to design a clustering model which can learn the importance of these link types automatically. This will enhance the clustering quality because it marginalizes the impact of low quality types of neighbors of an object during the clustering process.

We present examples of clustering tasks in two concrete heterogeneous information networks in the following.

Example 6.1. (Bibliographic information network) A bibliographic network is a typical heterogeneous network, containing objects from three types of entities, namely papers, publication venues (conferences or journals), and authors. Each paper has different link types to its authors

and publication venue. Each paper is associated with the text attribute as a bag of words. Each author and venue links to a set of papers, but contains no attributes (in our case). The application of a clustering process according to the text attribute in such a scenario can help detect research areas, and decide the research areas for authors, venues and papers.

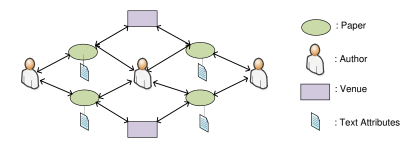


Figure 6.2: Illustration of bibliographic information network.

Note that, we treat text as attributes of papers in this case instead of term entities as in previous chapters. Multiple types of objects and links in this network are illustrated in Figure 6.2. For objects of different types, their cluster memberships may need to be determined by different kinds of information: for authors and venues, the only available information is from the papers linked to them; for papers, both text attributes and links of different types are provided. Note that, even for papers that are associated with text attributes, using link information can further help the clustering quality when the observations of the text data is very limited (e.g., using text merely from titles). Also, we may expect that the neighbors of an author type play a more important role in deciding a paper's cluster compared with the neighbor of a venue type. This needs to be automatically learned in terms of the underlying relation strengths.

Example 6.2. (Weather sensor network) Weather sensor networks typically contain different kinds of sensors for detecting different attributes, such as precipitation or temperature. Some sensors may have incorrect or no readings because of the inaccuracy or malfunctioning of the instruments. The links between sensors are generated according to their k nearest neighbors under geo-distances, in order to incorporate the importance of locality in weather patterns. The clustering of such sensors according to both precipitation and temperature attributes can be useful in determining regional weather patterns.

Figure 6.3 illustrates a weather sensor network containing two types of sensors: temperature

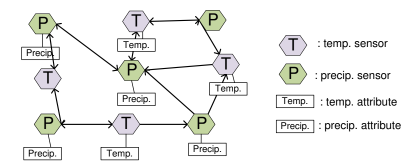


Figure 6.3: Illustration of weather sensor information network.

and precipitation. A sensor may sometimes register none or multiple observations. Although it is desirable to use the complete observations on both temperature and precipitation to determine the weather pattern of a location, in reality a sensor object may contain only partial attribute (e.g., temperature values only for temperature sensors), and both of the attribute and link information are needed for correctly detecting the clusters. Still, which type of links plays a more important role needs to be determined in the clustering process.

Formally, given a network $G = (\mathcal{V}, \mathcal{E}, W)$, a specified subset of its associated attributes $X \in \mathcal{X}$, the attribute observations $\{v[X]\}$ for all objects, and the number of clusters K, our goal is:

- 1. to learn a soft clustering for all the objects $v \in \mathcal{V}$, denoted by a membership probability matrix, $\Theta_{|\mathcal{V}| \times K} = (\boldsymbol{\theta}_v)_{v \in \mathcal{V}}$, where $\Theta(v, k)$ denotes the probability of object v in cluster $k, 0 \leq \Theta(v, k) \leq 1$ and $\sum_{k=1}^K \Theta(v, k) = 1$, and $\boldsymbol{\theta}_v$ is the K dimensional cluster membership vector for object v, and
- 2. to learn the strengths (importance weights) of different link types in determining the cluster memberships of the objects, $\gamma_{|\mathcal{R}|\times 1}$, where $\gamma(r)$ is a real number and stands for the importance weight for the link type $r \in \mathcal{R}$.

6.3 The Clustering Framework

We propose a novel probabilistic clustering model in this section and introduce the algorithm that optimizes the model in Section 6.4.

6.3.1 Model Overview

Given a network G, with the observations of its links and the observations $\{v[X]\}$ for the specified attributes $X \in \mathcal{X}$, a good clustering configuration Θ , which can be viewed as hidden cluster information for objects, should satisfy two properties:

- 1. Given the clustering configuration, the observed attributes should be generated with a high probability. Especially, we model each attribute for each object as a separate mixture model, with each component representing a cluster.
- 2. The clustering configuration should be highly consistent with the network structure. In other words, linked objects should have similar cluster membership probabilities, and larger strength of a link type requires more similarity between the linked objects of this type.

Overall, we can define the likelihood of the observations of all the attributes $X \in \mathcal{X}$ as well as the hidden continuous cluster configuration Θ , given the underneath network G, the relation strength vector γ , and the cluster component parameter β , which can be decomposed into two parts, the generative probability of the observed attributes given Θ and the probability of Θ given the network structure:

$$p(\{\{v[X]\}_{v\in\mathcal{V}_X}\}_{X\in\mathcal{X}},\Theta|G,\boldsymbol{\gamma},\boldsymbol{\beta}) = \prod_{X\in\mathcal{X}} p(\{v[X]\}_{v\in\mathcal{V}_X}|\Theta,\boldsymbol{\beta})p(\Theta|G,\boldsymbol{\gamma})$$
(6.1)

From a generative point of view, this model explains how observations for attributes associated with objects are generated: first, a hidden layer of variables Θ is generated according to the probability $p(\Theta|G, \gamma)$, given the network structure G and the strength vector γ ; second, the observed values of attributes associated with each object are generated according to mixture models, given the cluster membership of the object, as well as the cluster component parameter β , with the probability $\prod_{X \in \mathcal{X}} p(\{v[X]\}_{v \in \mathcal{V}_X} | \Theta, \beta)$.

The goal is then to find the best parameters γ and β , as well as the best clustering configuration Θ that maximize the likelihood. The detailed modeling of the two parts is introduced in the following.

6.3.2 Modeling Attribute Generation

Given a configuration Θ for the network G, namely, the membership probability vector $\boldsymbol{\theta}_v$ for each object v, the attribute observations for each object v are conditionally independent with observations from other objects. Each attribute X associated with each object v is then assumed following the same family of mixture models that share the same cluster components, with the component mixing proportion as the cluster membership vector $\boldsymbol{\theta}_v$. For simplicity, we first assume that only one attribute X is specified for the clustering purpose and then briefly discuss a straightforward extension to the multi-attribute case.

Single Attribute

Let X be the only attribute we are interested in the network, and let v[X] be the observed values for object v, which may contain multiple observations. It is natural to consider that the attribute observation v[X] for each object v is generated from a mixture model, where each component is a probabilistic model that stands for a cluster, with the parameters to be learned, and the component weights denoted by θ_v . Formally, the probability of all the observations $\{v[X]\}_{v \in \mathcal{V}_X}$ given the network configuration Θ is modeled as:

$$p(\lbrace v[X]\rbrace_{v\in\mathcal{V}_X}|\Theta,\boldsymbol{\beta}) = \prod_{v\in\mathcal{V}_X} \prod_{x\in v[X]} \sum_{k=1}^K \theta_{v,k} p(x|\boldsymbol{\beta}_k)$$
(6.2)

where K is the number of clusters, and β_k is the parameter for component k. In this chapter, we consider two types of attributes, one corresponding to text attributes with categorical distributions, and the other numerical attributes with Gaussian distributions.

1. Text attribute with categorical distribution: In this case, objects in the network contain text attributes in the form of a term list, from the vocabulary l=1 to m. Each cluster k has a different term distribution following a categorical distribution, with the parameter $\beta_k = (\beta_{k,1}, \ldots, \beta_{k,m})$, where $\beta_{k,l}$ is the probability of term l appearing in cluster k, that is, $X|k \sim discrete(\beta_{k,1}, \ldots, \beta_{k,m})$. Following the frequently used topic modeling method PLSA [47], each term in the term list for an object v is generated from the mixture model, with each component as a categorical distribution over terms described by β_k , and the component coefficient is θ_v .

Formally, the probability of observing all the current attribute values is:

$$p(\{v[X]\}_{v \in \mathcal{V}_X} | \Theta, \beta) = \prod_{v \in \mathcal{V}_X} \prod_{l=1}^m (\sum_{k=1}^K \theta_{v,k} \beta_{k,l})^{c_{v,l}}$$
(6.3)

where $c_{v,l}$ denotes the count of term l that object v contains.

2. Numerical attribute with Gaussian distribution: In this case, objects in the network contain numerical observations in the form of a value list, from the domain \mathbb{R} . The kth cluster is a Gaussian distribution with parameters $\boldsymbol{\beta}_k = (\mu_k, \sigma_k^2)$, that is, $X|k \sim \mathcal{N}(\mu_k, \sigma_k^2)$, where μ_k and σ_k are mean and standard deviation of normal distribution for component k. Each observation in the observation list for an object v is generated from the Gaussian mixture model, where each component is a Gaussian distribution with parameters μ_k, σ_k^2 , and the component coefficient is $\boldsymbol{\theta}_v$. The probability density for all the observations for all objects is then:

$$p(\{v[X]\}_{v \in \mathcal{V}_X} | \Theta, \beta) = \prod_{v \in \mathcal{V}_X} \prod_{x \in v[X]} \sum_{k=1}^K \theta_{v,k} \frac{1}{\sqrt{2\pi\sigma_k^2}} e^{-\frac{(x-\mu_k)^2}{2\sigma_k^2}}$$
(6.4)

Multiple Attributes

As in the weather sensor network example, we are interested in multiple attributes, namely temperature and precipitation. Generally, if multiple attributes in the network are specified by users, say X_1, \ldots, X_T , the probability density of observed attribute values $\{v[X_1]\}, \ldots, \{v[X_T]\}$ for a given clustering configuration Θ is as follows, by assuming the independence among these attributes:

$$p(\{v[X_1]\}_{v \in \mathcal{V}_{X_1}}, \dots, \{v[X_T]\}_{v \in \mathcal{V}_{X_T}} | \Theta, \boldsymbol{\beta}_1, \dots, \boldsymbol{\beta}_T) = \prod_{t=1}^T p(\{v[X_t]\}_{v \in \mathcal{V}_{X_t}} | \Theta, \boldsymbol{\beta}_t)$$
(6.5)

6.3.3 Modeling Structural Consistency

From the view of links, the more similar the two objects are in terms of cluster memberships, the more likely they are connected by a link. In order to quantitatively measure the consistency of a clustering result Θ with the network structure G, we define a novel probability density function for observing Θ .

We assume that linked objects are more likely to be in the same cluster, if the link type is of

importance in determining the clustering process. That is, for two linked objects v_i and v_j , their membership probability vectors $\boldsymbol{\theta}_i$ and $\boldsymbol{\theta}_j$ should be similar. Within the same type of links, the higher link weight (w(e)), the more similar $\boldsymbol{\theta}_i$ and $\boldsymbol{\theta}_j$ should be. Further, a certain link type may be of greater importance, and will influence the similarity to a greater extent. The consistency of a configuration Θ with the network G, is evaluated with the use of a composite analysis with respect to all the links in the network in the form of a probability density value. A more consistent configuration of Θ will yield a higher probability density value. In the following, we first introduce how the consistency of two cluster membership vectors is defined with respect to a single link, and then show how this analysis can be applied over all links in order to create a probability density value as a function of Θ .

For a link $e = \langle v_i, v_j \rangle \in \mathcal{E}$, with type $r = \phi(e) \in \mathcal{R}$, we denote the *importance of the link type* to the clustering process by a real number $\gamma(r)$. This is different from the weight of the link w(e), which is specified in the network as input, whereas the value of $\gamma(r)$ is defined on link types and needs to be learned. We denote the consistency function of two cluster membership vectors θ_i and θ_j with link e under strength weights for each link type γ by a feature function $f(\theta_i, \theta_j, e, \gamma)$. Higher values of this function imply greater consistency with the clustering results. In the following, we list several desiderata for a good feature function:

- 1. The value of the feature function f should increase with greater similarity of θ_i and θ_j .
- 2. The value of the feature function f should decrease with greater importance of the link e, either in terms of its specified weight w(e), or the learned importance $\gamma(r)$ for its link type. In other words, for the larger strength of a particular link type, two linked nodes are required to be more similar in order to claim the same level of consistency.
- 3. The feature function should not be symmetric between its first two arguments θ_i and θ_j , because the impact from node v_i to node v_j could be different from that of v_j to v_i .

The last criterion may need some further explanation. For example, in a citation network, a paper i may cite paper j, because i feels that j is relevant to itself, while the reverse may not be necessarily true. In the experimental section, we will show that asymmetric feature functions produce higher accuracy in link prediction.

We then propose a cross entropy-based feature function, which satisfies all of the desiderata listed above. For a link $e = \langle v_i, v_j \rangle \in \mathcal{E}$, with relation type $r = \phi(e) \in \mathcal{R}$, the feature function $f(\theta_i, \theta_j, e, \gamma)$ is defined as:

$$f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \boldsymbol{\gamma}) = -\gamma(r)w(e)H(\boldsymbol{\theta}_j, \boldsymbol{\theta}_i) = \gamma(r)w(e)\sum_{k=1}^K \theta_{j,k}\log\theta_{i,k}$$
(6.6)

where $H(\boldsymbol{\theta}_j, \boldsymbol{\theta}_i) = -\sum_{k=1}^K \theta_{j,k} \log \theta_{i,k}$, is the cross entropy from $\boldsymbol{\theta}_j$ to $\boldsymbol{\theta}_i$, which evaluates the deviation of v_j from v_i , in terms of the average coding bits needed if using coding schema based on the distribution of $\boldsymbol{\theta}_i$. For a fixed value of $\gamma(r)$, the value of $H(\boldsymbol{\theta}_j, \boldsymbol{\theta}_i)$ is minimal and (therefore) f is maximal, when the two vectors are identical. It is also evident from Eq. (6.6) that the value of f decreases with increasing learned link type strength $\gamma(r)$ or input link weight w(e). We require $\gamma \geq 0$, in the sense that we do not consider links that connect dissimilar objects. The value of f so defined is a non-positive function, with larger value indicating a higher consistency of the link.

Other distance functions such as KL-divergence could replace the cross entropy in the feature function. However, as cross entropy favors distributions that concentrate on one cluster $(H(\theta_j, \theta_i))$ achieves the lowest distance, when $\theta_j = \theta_i$ and $\theta_{i,k} = 1$ for some cluster k, which agrees with our clustering purpose, we pick it over KL-divergence.

We then propose a log-linear model to model the probability of Θ given the link type weights γ , where the probability of one configuration Θ is defined as the exponential of the summation of feature functions of all the links in G:

$$p(\Theta|G, \gamma) = \frac{1}{Z(\gamma)} \exp\{\sum_{e = \langle v_i, v_j \rangle \in \mathcal{E}} f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \gamma)\}$$
(6.7)

where γ is the strength weight vector for all link types, $f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \gamma)$ is the feature function defined on links of different types, and $Z(\gamma)$ is the partition function that makes the distribution function sum up to 1: $Z(\gamma) = \int_{\Theta} \exp\{\sum_{e=\langle v_i, v_j \rangle \in \mathcal{E}} f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \gamma)\} d\Theta$. The partition function $Z(\gamma)$ is an integral over the space of all the configurations Θ , and it is a function of γ .

6.3.4 The Unified Model

The overall goal of the network clustering problem is to determine the best clustering results Θ , the link type strengths γ and the cluster component parameters β that maximize the generative probability of attribute observations and the consistency with the network structure, described by the likelihood function in Eq. (6.1).

Further, we add a Gaussian prior to γ as a regularization to avoid overfitting, with the mean as 0, and the covariance matrix as $\sigma^2 I$, where σ is the standard deviation of each element in γ , and I is the identity matrix. We set $\sigma = 0.1$ in our experiments, and more complex strategy can be used to select σ according to labeled clustering results, which will not be discussed here. The new objective function is then:

$$g(\Theta, \boldsymbol{\beta}, \boldsymbol{\gamma}) = \log \sum_{X \in \mathcal{X}} p(\{v[X]\}_{v \in \mathcal{V}_X} | \Theta, \boldsymbol{\beta}) + \log p(\Theta|G, \boldsymbol{\gamma}) - \frac{||\boldsymbol{\gamma}||^2}{2\sigma^2}$$
(6.8)

In addition, we have the constraints that $\gamma \geq 0$, and some constraints for β that are dependent on the attribute distribution type. Also, $p(\{v[X]\}_{v \in \mathcal{V}_X} | \Theta, \beta)$ and $p(\Theta|G, \gamma)$ need to be replaced by the specific formulas proposed above for concrete derivations.

6.4 The Clustering Algorithm

This section presents a clustering algorithm that computes the proposed probabilistic clustering model. Intuitively, we begin with the assumption that all the types of links play an equally important role in the clustering process, then update the strength for each type according to the average consistency of links of that type with the current clustering results, and finally achieve a good clustering as well as a reasonable strength vector for link types. It is an iterative algorithm containing two steps in that clustering results and strengths of link types mutually enhance each other, which maximizes the objective function of Eq. (6.8) alternatively.

In the first step, we fix the link type weights γ to the best value γ^* , determined in the last iteration, then the problem becomes that of determining the best clustering results Θ and the attribute parameters β for each cluster component. We refer to this step as the *cluster optimization*

$$step: \ [\Theta^*, \boldsymbol{\beta}^*] = \underset{\Theta, \boldsymbol{\beta}}{\arg\max} \ g(\Theta, \boldsymbol{\beta}, \boldsymbol{\gamma}^*).$$

In the second step, we fix the clustering configuration parameters $\Theta = \Theta^*$ and $\beta = \beta^*$, corresponding to the values determined in the last step, and use it to determine the best value of γ , which is consistent with current clustering results. We refer to this step as the *link type strength* learning step: $\gamma^* = \arg\max_{\alpha > 0} g(\Theta^*, \beta^*, \gamma)$.

The two steps are repeated until convergence is achieved.

6.4.1 Cluster Optimization

In the cluster optimization step, each object has the link information from different types of neighbors, where the strength of each type of link is given, as well as the possible attribute observations. The goal is to utilize both link and attribute information to get the best clustering result for all the objects. Since γ is fixed in this step, the partition function and regularizer term become constants, and can be discarded for optimization purposes. Therefore, we can construct a simplified objective function $g_1(\cdot, \cdot)$, which depends only on Θ and β :

$$g_1(\Theta, \boldsymbol{\beta}) = \sum_{e = \langle v_i, v_j \rangle} f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \boldsymbol{\gamma}) + \sum_{v \in \mathcal{V}_X} \sum_{x \in v[X]} \log \sum_{k=1}^K \theta_{v,k} p(x|\boldsymbol{\beta}_k)$$
(6.9)

We derived an EM-based algorithm [32, 16] to solve Eq. (6.9). In the E-step, the probability of each observation x for each object v and each attribute X belonging to each cluster, usually called the hidden cluster label of the observation, $z_{v,x}$, is derived according to the current parameters Θ and β . In the M-step, the parameters Θ and β are updated according to the new membership for all the observations in the E-step. The iterative formulas for single text attribute and single Gaussian attribute are provided below.

1. Single categorical text attribute: Let $z_{v,l}$ denote the hidden cluster label for the lth term in the vocabulary for object v, Θ^{t-1} be the value of Θ at iteration t-1, and $\boldsymbol{\beta}^{t-1}$ be the value of $\boldsymbol{\beta}$ at iteration t-1. $\mathbf{1}_{\{v \in \mathcal{V}_X\}}$ is the indicator function, which is 1 if v contains this attribute,

and 0 otherwise. Then, we have:

$$p(z_{v,l}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1}) \propto \theta_{v,k}^{t-1} \beta_{k,l}^{t-1}$$

$$\theta_{v,k}^{t} \propto \sum_{e = \langle v, u \rangle} \gamma(\phi(e)) w(e) \theta_{u,k}^{t-1} + \mathbf{1}_{\{v \in \mathcal{V}_X\}} \sum_{l=1}^{m} c_{v,l} p(z_{v,l}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1})$$

$$\beta_{k,l}^{t} \propto \sum_{v \in \mathcal{V}_X} c_{v,l} p(z_{v,l}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1})$$
(6.10)

2. Single Gaussian numerical attribute: Let $z_{v,x}$ denote the hidden cluster label for the observation x for object v, Θ^t be the value of Θ at iteration t, and μ_k^t and σ_k^t be the values of mean and standard deviation for kth cluster at iteration t. $\mathbf{1}_{\{v \in \mathcal{V}_X\}}$ is the indicator function, which is 1 if v contains this attribute, and 0 otherwise. Then, we have:

$$p(z_{v,x}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1}) \propto \theta_{v,k}^{t-1} \frac{1}{\sqrt{2\pi(\sigma_{k}^{t-1})^{2}}} e^{-\frac{(x-\mu_{k}^{t-1})^{2}}{2(\sigma_{k}^{t-1})^{2}}}$$

$$\theta_{v,k}^{t} \propto \sum_{e=\langle v,u \rangle} \gamma(\phi(e)) w(e) \theta_{u,k}^{t-1} + \mathbf{1}_{\{v \in \mathcal{V}_{X}\}} \sum_{x \in v[X]} p(z_{v,x}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1})$$

$$\mu_{k}^{t} = \frac{\sum_{v \in \mathcal{V}_{X}} \sum_{x \in v[X]} x p(z_{v,x}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1})}{\sum_{v \in \mathcal{V}_{X}} \sum_{x \in v[X]} p(z_{v,x}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1})}$$

$$(\sigma_{k}^{2})^{t} = \frac{\sum_{v \in \mathcal{V}_{X}} \sum_{x \in v[X]} (x - \mu_{k}^{t})^{2} p(z_{v,x}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1})}{\sum_{v \in \mathcal{V}_{X}} \sum_{x \in v[X]} p(z_{v,x}^{t} = k | \Theta^{t-1}, \boldsymbol{\beta}^{t-1})}$$

For networks with multiple attributes, the formulae can be derived similarly. The readers can find the formulae for the case of two Gaussian numerical attributes in [95].

From the update rules, we can see that the value of the membership probability for an object is dependent on its neighbors' memberships, the strength of the link types, the weight of the links, and the attribute associated with it (if any). When an object contains no attributes in the specified set, or contains no observations for the specified attributes, the cluster membership is totally determined by its linked objects, which is a weighted average of their cluster memberships and the weight is determined by both the weight of the link and the weight of the link type. When an object contains some observations of the specified attributes, its cluster membership is determined by both its neighbors and these observations for each possible attribute.

6.4.2 Link Type Strength Learning

The link type strength learning step is to find the best strength weight for each type of links that makes the current clustering result to be generated with the highest probability. By doing so, the low quality link types that connect objects not so similar will be punished and assigned with low strength weights; while the high quality link types will be assigned with high strength weights.

Since the values of Θ and β are fixed in this step, the only relevant parts of the objective function (for optimization purposes) are those which depend on γ . These are the structural consistency modeling part and the regularizer over γ . Therefore, we can construct the following simplified objective function $g_2(\cdot)$ as a function of γ :

$$g_2(\gamma) = \sum_{e = \langle v_i, v_j \rangle} f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \gamma) - \log Z(\gamma) - \frac{||\gamma||^2}{2\sigma^2}$$
(6.12)

In addition, we have the linear constraints as $\gamma \geq 0$.

However, g_2 is difficult to be optimized directly, since the partition function $Z(\gamma)$ is an integral over the entire space of valid values of Θ , which is intractable. Instead, we construct an alternate approximate objective function g'_2 , which factorizes $\log p(\Theta|G)$ as the sum of $\log p(\theta_i|\theta_{-i},G)$, namely the pseudo-log-likelihood, where $p(\theta_i|\theta_{-i},G)$ is the conditional probability of θ_i given the remaining objects' clustering configurations, which turns out to be dependent only on its neighbors. The intuition of using pseudo-log-likelihood to approximate the real log-likelihood is that, if the probability of generating the clustering configuration for each object conditional on its neighbors is high, the probability of generating the whole clustering configuration should also be high. In other words, if the local patches of a network are very consistent with the clustering results, the consistency over the whole network should also be high.

In particular, we choose each local patch of the network as an object and all its out-link neighbors. In this case, every link is considered exactly once, and the newly designed objective function $g'_2(\cdot)$ is as follows:

$$g_2'(\gamma) = \sum_{i=1}^{|V|} \left(\sum_{e=\langle v_i, v_j \rangle} f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \boldsymbol{\gamma}) - \log Z_i(\boldsymbol{\gamma}) \right) - \frac{||\boldsymbol{\gamma}||^2}{2\sigma^2}$$
(6.13)

where $\log Z_i(\gamma) = \log \int_{\theta_i} e^{\sum_{e=\langle v_i, v_j \rangle} f(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j, e, \gamma)} d\boldsymbol{\theta}_i$, the local partition function for object v_i , with the linear constraints $\gamma \geq 0$.

As the joint distribution of Θ as well as the conditional distribution of θ_i given its out-link neighbors are both belonging to exponential families, both g_2 and g'_2 are concave functions of γ . Therefore, the maximum value is either achieved at the global maximum point or at the boundary of constraints. The Newton-Raphson method is used to solve the optimization problem. It needs to calculate the first and second derivative of $g'_2(\gamma)$ with respect to γ , which is non-trivial in our case. We discuss the computation of these below.

By re-examining $p(\boldsymbol{\theta}_i | \{\boldsymbol{\theta}_j\}_{\forall e = \langle v_i, v_j \rangle}, G)$, the conditional probability for each object i given its out-link neighbors, we have:

$$p(\boldsymbol{\theta}_i|\{\boldsymbol{\theta}_j\}_{\forall e=\langle v_i, v_j \rangle}, G) \propto \prod_{k=1}^K \theta_{ik}^{\sum_{e=\langle v_i, v_j \rangle} \gamma(\phi(e))w(e)\theta_{j,k}}$$
(6.14)

It is easy to see that $p(\boldsymbol{\theta}_i|\{\boldsymbol{\theta}_j\}_{\forall e=\langle v_i,v_j\rangle},G)$ is a Dirichlet distribution with parameters $\alpha_{ik}=\sum_{e=\langle v_i,v_j\rangle}\gamma(\phi(e))w(e)\theta_{j,k}+1$, for k=1 to K. Therefore, the local partition function for each object $i,Z_i(\boldsymbol{\gamma})$, should be the constant $B(\boldsymbol{\alpha}_i)$ as in Dirichlet distribution, where $\boldsymbol{\alpha}_i=(\alpha_{i1},\ldots,\alpha_{iK})$ and $B(\boldsymbol{\alpha}_i)=\frac{\prod_{k=1}^K\Gamma(\alpha_{ik})}{\Gamma(\sum_{k=1}^K\alpha_{ik})}$. Then the first and second derivatives $(\nabla g_2')$ and $B(\boldsymbol{\alpha}_2')$ can be calculated now as each $B(\boldsymbol{\alpha}_i)$ is a function of Gamma functions. Then, we can use the Newton-Raphson method to determine the value of $\boldsymbol{\gamma}$ that maximizes $B(\boldsymbol{\gamma}_i')$ with the following iterative steps:

1.
$$\boldsymbol{\gamma}^{t+1} = \boldsymbol{\gamma}^t - [Hg_2'(\boldsymbol{\gamma}^t)]^{-1}\nabla g_2'(\boldsymbol{\gamma}^t);$$

2.
$$\forall r \in \mathcal{R}$$
, if $\gamma(r)^{t+1} < 0$, set $\gamma(r)^{t+1} = 0$.

6.4.3 Putting together: The GenClus Algorithm

We integrate the two steps discussed above to construct a <u>Gen</u>eral Heterogeneous Network <u>Clus</u>tering algorithm, GenClus. The algorithm includes an outer iteration that updates Θ and γ alternatively, and two inner iterations that optimize Θ using the EM algorithm and optimize γ using the Newton-Raphson method respectively. For the initialization of γ in the outer iteration, we initialize it as an all-1 vector. This means that all the link types in the network are initially considered equally important. For the initialization of Θ' in the inner iteration for optimizing Θ , we can either (1)

assign Θ'^0 with random assignments, or (2) start with several random seeds, run the EM algorithm for a few steps for each random seed, and choose the one with the highest value of the objective function g_1 as the real starting point. The latter approach will produce more stable results.

The time complexity for the EM algorithm in the first step is $O(t_1(Kd_1|\mathcal{V}|+K|\mathcal{E}|))$, where t_1 is the number of iterations, d_1 is the average number of observations for each object, K is the number of clusters, $|\mathcal{V}|$ is the number of objects, and $|\mathcal{E}|$ is the number of links in the network, which is linear to $|\mathcal{V}|$ for sparse networks. The time complexity of the algorithm in the step of maximizing γ is dependent on the time for calculating the first derivative and Hessian matrix of $g'_2(\gamma)$, and the matrix inversion involved Newton-Raphson algorithm. This is $O(K|\mathcal{E}|+t_2|\mathcal{R}|^{2.376}))$, where K and $|\mathcal{E}|$ are with the same meaning as before, t_2 is the number of iterations, and $|\mathcal{R}|$ is the number of relations in the network. In all, the overall time complexity is $O(t(t_1(Kd_1|\mathcal{V}|+K|\mathcal{E}|)+t_2|\mathcal{R}|^{2.376}))$, where t is the number of outer iterations. In other words, for each outer iteration, the time complexity is approximately linear to the number of objects in the network when the network is sparse. Therefore, the GenClus algorithm is quite scalable.

6.5 Experiments

In this section, we examine the effectiveness of GenClus on several real and synthetic datasets.

6.5.1 Datasets

Two real networks and one synthetic network are used in this study. We extracted two networks from the *DBLP "four-area" dataset* [103, 39], by using different subsets of entities and the links between them to represent the underlying network structures. This dataset was extracted from 20 major conferences from the four areas corresponding to database, data mining, information retrieval, and artificial intelligence. Besides the real networks, we also generated a synthetic weather sensor network. We describe these networks below:

(a) **DBLP four-area A-V network.** This network contains two types of objects, authors (A) and venues (V); and three types of links depending upon publication behavior, namely $publish_in(A, V)$ (short for $\langle A, V \rangle$), $published_by(V, A)$ (short for $\langle V, A \rangle$), and coauthor(A, A) (short for $\langle A, A \rangle$).

The links are associated with a weight corresponding to the number of papers that an author has published in a venue, a venue is contributed by an author, and the two authors have coauthored, respectively. The author nodes and venue nodes contain text corresponding to the text from the titles of all the papers they have ever written or published.

- (b) **DBLP four-area A-V-P network.** This network contains objects corresponding to authors (A), venues (V) and papers (P); and four types of links depending upon the publication behavior, namely write(A, P) (short for $\langle A, P \rangle$), $written_by(P, A)$ (short for $\langle P, A \rangle$), publish(V, P) (short for $\langle V, P \rangle$), and $published_by(P, V)$ (short for $\langle P, V \rangle$). In this case, the links have binary weights, corresponding to presence or absence of the link. Only papers contain text attributes that are from their titles.
- (c) Weather sensor network. This network is synthetically generated, containing two types of objects: temperature (T) and precipitation (P) sensors, and four link types between any two types of sensors denoting the kNN relationship: $\langle T, T \rangle$, $\langle T, P \rangle$, $\langle P, T \rangle$, and $\langle P, P \rangle$. The links are binary weighted according to their k-nearest neighbors. The attributes associated with a sensor correspond to either temperature or precipitation, depend on the type of the sensor. We use the weather network generator to generate two sets of synthetic climate sensor networks, each containing 4 clusters, and each sensor is linked to 5 nearest neighbors for each type (10 in total). In each setting, we vary the number of sensors, by fixing the number of temperature sensors at 1000, and precipitation sensors as 250, 500, and 1000. For each setting, the number of observations for each object may be 1, 5 or 20. In all, for each weather pattern setting, we have 9 networks with different configurations.

6.5.2 Effectiveness Study

We use two measures for our effectiveness study. First, the labels associated with the nodes in the datasets provide a natural guidance in examining the coherence of the clusters. We use *Normalized Mutual Information (NMI)* [94] to compare our clustering result with the ground truth. Second, we use link prediction accuracy to test the clustering accuracy. The similarity between two objects can be calculated by similarity function defined on their two membership vectors, such as using cosine similarity function. Clearly, a better clustering quality will lead to better computation of

similarity (and therefore the accuracy of link prediction). For a certain type of relation $\langle A, B \rangle$, we calculate all the similarity scores between each $v_A \in A$ and all the objects $v_B \in B$, and compare the similarity-based ranked list with the true ranked list determined by the link weights between them. We use the measure *Mean Average Precision (MAP)* [127] to compare the two ranked links.

Clustering Accuracy Test We choose clustering methods that can deal with both the links and attributes as our baselines. None of these baselines is capable of leveraging different link types of different impacts to the clustering process. Therefore, we set each link type strength as 1 for these baselines. Second, we choose different baselines for clustering networks with text attributes and for clustering networks with numerical attributes, since there are no unified clustering methods (other than our presented GenClus) that can address both situations in the same framework.

For the DBLP four-area A-V network and the DBLP four-area A-V-P network that are with text attributes, we use NetPLSA [75] and iTopicModel [98] as baselines, which aim at improving topic qualities by using link information in homogeneous networks. We compare GenClus with the baselines by assuming homogeneity of links for the latter algorithms. The mean and standard deviation of NMI of the 20 running results are shown for the DBLP A-V network and the DBLP A-V-P network in Figures 6.4 and 6.5 respectively. From the results, we can see that the GenClus algorithm is much more effective than both the iTopicModel and the NetPLSA methods in both networks. This is because of the ability of the former algorithms to learn and leverage the strengths of different link types in the clustering process. Furthermore, the standard deviation of NMI over different runs is much lower for GenClus, which suggests that the algorithm is more robust to the initial settings with the learned strength weights for different link types.

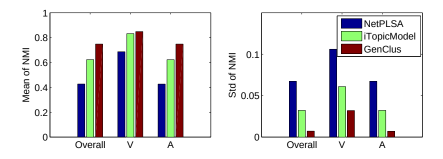


Figure 6.4: Clustering accuracy comparisons for the A-V network.

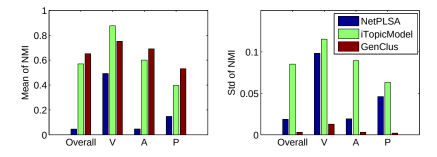


Figure 6.5: Clustering accuracy comparisons for the A-V-P network.

The A-V network is the easiest case among the three networks, since it only contains one type of attribute (the text attribute), and all object types contain this attribute, namely the attribute is complete for every object. The A-V-P network is a more difficult case than the previous one, because not every type of objects contain the text attributes. This requires the clustering algorithm to be more robust to deal with objects with no attributes at all. From the results, we can see that GenClus is more robust than NetPLSA algorithm, which outputs almost random predictions for authors for the A-V-P network. Although the homogenous methodology of the iTopicModel algorithm performs better for objects of type V for A-V network (See Figure 6.5), GenClus still has an overall better performance. This is because our objective function is over all the objects rather than a particular type.

We also examined the actual clusters obtained by the algorithm on *DBLP A-V network*, and list corresponding cluster membership for several venues and authors in Table 6.1, where the research area names are given afterwards according the clustering results. We can see that the clustering results for the *GenClus* algorithm are consistent with human intuition.

Table 6.1: Case Studies of Cluster Membership Results

Object	DB	DM	IR	ML
SIGMOD	0.8577	0.0492	0.0482	0.0449
KDD	0.0786	0.6976	0.1212	0.1026
CIKM	0.2831	0.1370	0.4827	0.0971
Jennifer Widom	0.7396	0.0830	0.1061	0.0713
Jim Gray	0.8359	0.0656	0.0536	0.0449
Christos Faloutsos	0.4268	0.3055	0.1380	0.1296

The synthetic weather sensor network is the most difficult case among the three networks, as

it has two types of attributes corresponding to different types of sensors. Furthermore, all sensor nodes contain incomplete observations of the attributes. Existing algorithms cannot address these issues well. We compare the clustering results of GenClus with two baselines, by comparing the cluster labels with maximum probabilities with the ground truth. In this case, we choose the initial seed for GenClus as one of the tentative running results with the highest objective function, and the number of iterations is set to 5. The first baseline is the k-means algorithm, and the second one is a spectral clustering method that combines the network structure and attribute similarity as a new similarity matrix. We use the framework given in [91], which utilizes modularity objective function in the network part, but we replace the cosine similarity by Euclidean distance in the attribute part as in [125] for better clustering results. As both methods cannot handle the problem of incomplete attributes, we use interpolation to make each sensor have a regular 2-dimensional attribute, by using the mean of all the observations of its neighbors and itself. For the spectral clustering-based framework, we centralize the data by extracting the mean and then normalize them by the standard deviation, in order to make the attribute part comparable with the modularity part in the objective function. Both parts are set to have equal weights.

The results are summarized in Figures 6.6 and 6.7. It is evident that the GenClus algorithm exhibits superior performance to the two baselines in most of the datasets (17 out of 18 cases). Furthermore, GenClus can produce more stable clustering results compared with k-means, which is very sensitive to the number of observations for each object, especially for Setting 2. GenClus is also highly adaptive in that there is no need of any weight specification for combining the network and attribute-contributions to the clustering process. This results in greater stability of GenClus. Another major advantage of GenClus (which is not immediately evident from the presented results) is that we can directly utilize every observation instead of the mean, while the baselines can only use a biased mean value because of the interpolation process.

Link Prediction Accuracy Test Next, the link prediction accuracy measured by MAP is compared between GenClus and the baselines. For the A-V network, we select the link type $\langle A, V \rangle$ for the prediction task, namely we want to predict which venues an author is likely to go. For the A-V-P network, we select the link type $\langle P, V \rangle$ for the prediction task, namely we want to predict which venue a paper is published in. As the prediction is based on the similarity between the two

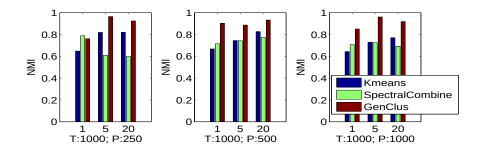


Figure 6.6: Clustering accuracy comparisons for weather sensor network Setting 1.

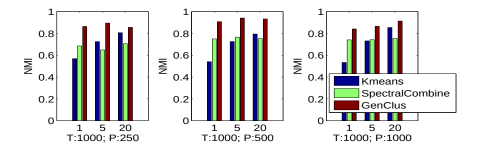


Figure 6.7: Clustering accuracy comparisons for weather sensor network Setting 2.

objects, say query object v_i with clustering membership $\boldsymbol{\theta}_i$ and candidate object v_j with clustering membership $\boldsymbol{\theta}_j$, three similarity functions are used here: (1) cosine similarity denoted as $\cos(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j)$; (2) the negative of Euclidean distance denoted as $-||\boldsymbol{\theta}_i - \boldsymbol{\theta}_j||$; and (3) the negative of cross entropy denoted as $-H(\boldsymbol{\theta}_j, \boldsymbol{\theta}_i)$. The results are summarized in Tables 6.2 and 6.3.

Table 6.2: Prediction accuracy for A-V relation in A-V network.

	NetPLSA	iTopicModel	GenClus
$\cos(oldsymbol{ heta}_i, oldsymbol{ heta}_j)$	0.4351	0.5117	0.7627
$- oldsymbol{ heta}_i - oldsymbol{ heta}_j $	0.4312	0.5010	0.7539
$H(\boldsymbol{\theta}_j, \boldsymbol{\theta}_i)$	0.4323	0.5088	0.7753

Table 6.3: Prediction accuracy for P-V Relation in A-V-P network.

	NetPLSA	iTopicModel	GenClus
$\cos(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j)$	0.2762	0.4609	0.5170
$\overline{- oldsymbol{ heta}_i - oldsymbol{ heta}_j }$	0.2759	0.4600	0.5142
$-H(oldsymbol{ heta}_j,oldsymbol{ heta}_i)$	0.2760	0.4683	0.5183

For the weather sensor network, we select the link type $\langle T, P \rangle$, namely we want to predict the P-

typed neighbors for T-typed sensors. We test the link prediction in the network with configuration as in Setting 1, with #T = 1000 and #P = 250. We only output the link prediction results for *GenClus* algorithm, since the other two baselines can only output hard clusters (exact cluster memberships rather than probabilities). The results are shown in Table 6.4.

Table 6.4: Prediction accuracy for $\langle T, P \rangle$ in weather sensor network.

	$\cos(\boldsymbol{\theta}_i, \boldsymbol{\theta}_j)$	$- oldsymbol{ heta}_i - oldsymbol{ heta}_j $	$-H(\boldsymbol{\theta}_j, \boldsymbol{\theta}_i)$
MAP	0.7285	0.7690	0.8073

From the results, it is evident that the GenClus algorithm has the best link prediction accuracy in terms of different similarity functions. Also, the results show that the asymmetric function $-H(\theta_j, \theta_i)$ provides the best link prediction accuracy, especially for better clustering results such as those obtained by GenClus and in the weather sensor network where the out-link neighbors are different from the in-link neighbors.

Analysis of Link Type Strength Since the process of learning the semantic importance of relations is important in a heterogeneous clustering approach, we present the learned relation strengths in Figure 6.8 for the two DBLP four-area networks. From the figure, it is evident that in the A-V network, the link type $\langle A, V \rangle$ has greater importance to the clustering process than the link type $\langle A, A \rangle$, and thus is more important in deciding an author's membership. This is because the spectrum of co-authors is often broad, whereas authors' publication frequency in each venue can be a more reliable predictor of clustering behavior. For the A-V-P network, we can see that the link type $\langle P, V \rangle$ has the weight 3.13, whereas the link type $\langle P, A \rangle$ has a much higher weight 13.30. This suggests that the latter link type is more reliable in deciding the cluster for papers, since the venues usually have a broader research track than the authors. For example, it is difficult to judge the cluster for a paper if we only know that it is published in the CIKM venue. The ability of our algorithm to learn such important characteristics of different link types is one of the reasons that it is superior to other competing methods.

For the weather sensor network, we summarize the link type strengths for the three networks with different network sizes that contain 5 observations for each sensor and use the configuration

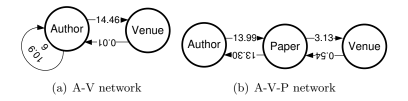


Figure 6.8: Strengths for link types in two DBLP four-area networks.

of Setting 1, in Table 6.5. It is evident that GenClus can correctly detect: (1) the P-typed sensors cannot be trusted as much as the other typed sensors when P-typed sensors are very sparse, due to their farther distance and less similarity to other objects (the strengths of $\langle T, P \rangle$ and $\langle P, P \rangle$ relations decrease as #P decreases); (2) for both types of sensors, T-typed neighbors are more trustable than P-typed ones, due to the higher quality of T-typed data in the network setting.

	$\langle T, T \rangle$	$\langle T, P \rangle$	$\langle P, T \rangle$	$\langle P, P \rangle$
T:1000; P: 250	3.14	2.88	1.60	1.32
T:1000; P: 500	3.16	3.05	2.38	1.98
T:1000; P: 1000	3.14	3.03	3.34	2.78

Table 6.5: Link Type Strength for Weather Sensor Network in Setting 1

6.5.3 A Typical Running Case

One of the core ideas of this algorithm is to enable a mutual learning process between importance of link types for the clustering and the actual clustering results. In this section, we provide some detailed results within the different iterations of the algorithm, which suggests that such a mutual learning process does occur. In particular, a typical running case for the A-V network is illustrated in Figure 6.9. Figure 6.9(a) shows how the clustering accuracy progresses along with the changes in the importance of different link types. Figure 6.9(b) shows how the strength weights change along with the clustering results at different iterations and finally converge to the correct values. Note that, we plotted the initial value γ at iteration 0 in Figure 6.9(b), which is an all-one value.

6.5.4 Efficiency Study

In this part, we study the efficiency of our algorithm. We illustrate the execution time of each inner iteration for the EM algorithm, which is the bottleneck component for the overall time

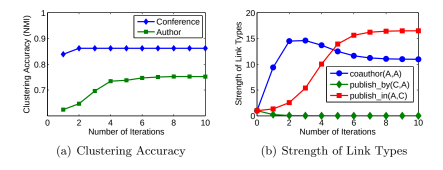


Figure 6.9: A Running Case on A-V network: Iterations 1 to 10

complexity. The results are presented for the weather sensor network with different sizes and number of observations for both the pattern generator settings. The results are illustrated in Figure 6.10, and are consistent with our observations in the complexity section about the scalability with the number of objects.

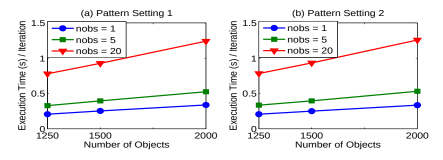


Figure 6.10: Scalability Test over Number of Objects

One observation is that the EM approach is very easy to parallelize, which is the major component for *GenClus* in terms of time complexity. We tested the parallel version of the EM algorithm with the use of 4 parallel threads, and it turned out that the execution time is improved by a factor of 3.19. This suggests that the approach is highly parallelizable.

6.6 Related Work

Clustering is a classical problem in data analysis, and has been studied extensively in the context of multi-dimensional data [50]. Most of these algorithms are attribute based, in which the data corresponds to a multi-dimensional format, and does not contain links. A number of clustering methods [18, 58, 23, 26] have been proposed on the basis of network structure only, mainly in

the context of the community detection problem [4, 64, 29]. A recent piece of work extends the network clustering problem to the heterogeneous scenario [103]. However, this latter method [103] is designed for a specific kind of network structure, referred as the *star network schema*, and is not applicable to networks of general structure. Furthermore, it cannot be easily integrated with attribute information.

Recently, some studies [12, 91, 75, 118] have shown that by considering the link constraints in addition to the attributes, the clustering accuracy can be enhanced. However, most of these algorithms require that the network links, objects and their attributes are all homogeneous. A recent clustering method [133] integrates the network clustering process with categorical attributes by considering the latter as augmented objects, but the same methodology cannot be applied to numerical values. Some other algorithms [91] can cluster objects with numerical attributes by combining the network clustering objective function with a numerical clustering objective function, but it is difficult to decide the weight to combine them, and cannot deal with the incomplete attributes properly. [70] provides a framework for clustering objects in relational networks with attributes. However, they studied a different clustering problem by clustering objects from different types separately, and did not study the interplay of importance of different link types and the clustering results. Probabilistic relational models, such as [105], provide a way to model a rational database containing both attributes and links, but do not consider the scenario studied in this paper that clustering purposes could be different according to the specified attributes. Also, they cannot handle the problem of incomplete attributes due to the discriminative nature of their methods.

There are several different philosophies on using the link information in addition to attributes to help the clustering in networks. First, in [91, 133], links are viewed to provide another angle of similarity measure between objects besides the attribute-based similarity measure, and the final clustering results are generated by combining the two angles. Second, In relational clustering [70] and probabilistic relational models [105], every link is treated as equally important and the probability of a link appearance is modeled explicitly according to the cluster memberships of the two objects of the link, in a way of building mixture of block models [3]. Third, in [75, 98], links are considered to provide additional information about the similarity between objects that are consistent with the attributes, and the final clustering result is a more smoothing version

compared with the one merely using attributes. However, none of these views is able to model the fact that different relations should have different importance in determining the clustering process for a certain purpose. Our philosophy in modeling link consistency is more similar to the third line, that is, two objects linking together indicates a higher chance that they have similar cluster memberships. Moreover, we further associate each type of links with a different importance weight in measuring the consistency under a given clustering purpose, and thus each type of relation carries different strengths in passing the cluster membership between the linked objects.

6.7 Conclusion

We propose GenClus, the first approach to cluster general heterogeneous information networks with different link types and different attribute types, such as numerical or text attributes, with guidance from a specified subset of the attributes. Our algorithm is designed to seamlessly work in the case when some of the nodes may not have the complete attribute information. One key observation of the work is that heterogeneous network clustering provides a tremendous challenge because different types of links may present different levels of semantic importance to the clustering process. The importance of different semantic link types is learned in order to enable an effective clustering algorithm that meets a user's demand. We present experimental results which show the advantages of the approach over competing methods, including a number of interesting case studies and a study of the algorithm efficiency.

Chapter 7

User-Guided Clustering via Meta-Path Selection

In this chapter, we study another relation strength-aware mining problem: user-guided clustering of a certain type of objects, based on their involvement of multiple types of relations, encoded by meta-paths, in a heterogeneous information network. In an application, a user often has the best say on the kinds of clusters she would like to get, and such guidance will lead to the selection of appropriate combination of weighted meta-paths for generation of desired clustering results.

7.1 Overview

With the advent of massive social and information networks, link-based clustering of objects in networks becomes increasingly important since it may help discover hidden knowledge in large networks. Link-based clustering groups objects based on their links instead of attribute values. This is especially useful when attributes of objects cannot be fully obtained. Most existing link-based clustering algorithms are on homogeneous networks, where links carry the same semantic meaning and only differ in their strengths (i.e., weights). However, most real-world networks are heterogeneous, where objects are of multiple types and are linked via different types of relations or sequences of relations, forming a set of meta-paths. These meta-paths indicate different relations among object types and imply diverse semantics, and thus clustering on different meta-paths will generate rather different results, as shown below.

Example 7.1. (Meta-path-based clustering) A toy heterogeneous information network is shown in Figure 7.1, which contains three types of objects: organization (O), author (A) and venue (V), and two types of links: the solid line represents the affiliation relation between author and organization, whereas the dashed one the publication relation between author and venue. Authors are then connected (indirectly) via different meta-paths. For example, A - O - A is a meta-path denoting a

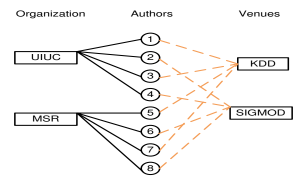


Figure 7.1: A toy heterogeneous information network containing organizations, authors and venues.

relation between authors via organizations (i.e., colleagues), whereas A - V - A denotes a relation between authors via venues (i.e., publishing in the same venues). A question then arises: which type of connections should we use to cluster the authors?

Obviously, there is no unique answer to this question: Different meta-paths lead to different author connection graphs, which may lead to different clustering results. In Figure 7.2(a), au-

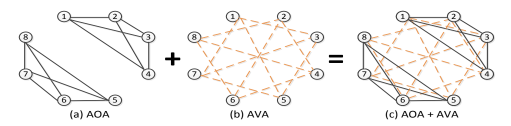


Figure 7.2: Author connection graphs under different meta-paths.

thors are connected via organizations and form two clusters: $\{1,2,3,4\}$ and $\{5,6,7,8\}$; in Figure 7.2(b), authors are connected via venues and form two different clusters: $\{1,3,5,7\}$ and $\{2,4,6,8\}$; whereas in Figure 7.2(c), a connection graph combining both meta-paths generates 4 clusters: $\{1,3\},\{2,4\},\{5,7\}$ and $\{6,8\}$.

This toy example shows that all the three clusterings look reasonable but they carry diverse semantics. It should be a user's responsibility to choose her desired meta-path(s). However, it is often difficult to ask her to explicitly specify one or a weighted combination of meta-paths. Instead, it is easier for her to give some guidance in other forms, such as giving one or a couple of examples for each cluster. For example, it may not be hard to give a few known conferences in each cluster (i.e., field) if one wants to cluster them into K research areas (for a user-desired K), or ask a user

to name a few restaurants if one wants to cluster them into different categories in a business review website (e.g., Yelp).

The new situation is that since we are dealing with heterogeneous networks, the previous work on user-guided clustering or semi-supervised learning approaches on (homogeneous) graphs [61, 134, 135] cannot apply. We need to explore meta-paths that represent heterogeneous connections across objects, leading to rich semantic meanings, hence diverse clustering results. With user guidance, a system will be able to learn the most appropriate meta-paths or their weighted combinations. The learned meta-paths will in turn provide an insightful view to help understand the underlying mechanism in the formation of a specific type of clustering, such as, which meta-path is more important to determine a restaurant's category?—the meta-path connecting them via customers, or the one connecting them via text in reviews, or the one determined by the nearest spatial locations?

We thus integrate meta-path selection with user-guided clustering in order to better cluster a user-specified type of objects (i.e., target objects) in a heterogeneous information network. We assume that user guidance is in the form of a small set of seeds for each cluster. For example, to cluster authors into 2 clusters in Example 7.1, a user may seed $\{1\}$ and $\{5\}$ for two clusters, which implies a selection of meta-path A - O - A; or seed $\{1\}, \{2\}, \{5\},$ and $\{6\}$ for four clusters, which implies a combination of both meta-paths A - O - A and A - V - A with about equal weight. Our goal is to (1) determine the weight of each meta-path for a particular clustering task, which should be consistent with the clustering results implied by the limited user guidance, and (2) output the clustering result according to the user guidance and using the learned weights for each meta-path.

We propose a probabilistic model that models the hidden clusters for target objects, the user guidance, and the quality weights for different meta-paths in a unified framework. An effective and efficient iterative algorithm *PathSelClus* is developed to learn the model, where the clustering quality and the meta-paths quality mutually enhance each other. The experiments with different tasks on two real networks show our algorithm outperforms the baselines.

7.2 The Meta-Path Selection Problem for User-Guided Clustering

Here we illustrate the problem using two heterogeneous information networks: the DBLP network and the Yelp network.

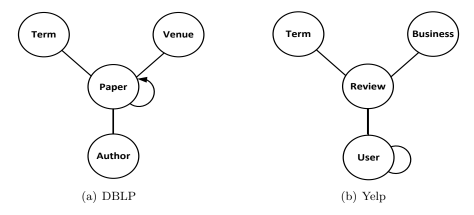


Figure 7.3: Examples of heterogeneous information networks.

Example 7.2. (The DBLP bibliographic network) As introduced before, DBLP is a computer science bibliographic network (see schema in Figure 7.3(a)) containing 4 types of objects: paper(P), author (A), term (T), and venue (V) (i.e., conferences and journals). Links exist between authors and papers by the relation of "write" and "written by," between papers and terms by "mention" and "mentioned by," and between venues and papers by "publish" and "published by." "Citation" relation between papers can be added further using other data source, such as Google scholar.

Example 7.3. (The Yelp network) Yelp is a website (http://www.yelp.com/) where users can write reviews for businesses. The Yelp network (see schema in Figure 7.3(b)) used in this chapter contains 4 types of objects: business (B), user (U), term (T), and review (R). Links exist between users and reviews by the relation of "write" and "written by," between reviews and terms by "mention" and "mentioned by," between businesses and reviews by "commented by" and "comment," and between users by "friendship" (not included in our dataset).

Following our previous discussion, a meta-path is defined by a sequence of relations in the network schema and can be denoted by a sequence of object types when there is no ambiguity. For example, A - P - A is a meta-path denoting the co-authorship between authors, and A - P - V is

a meta-path denoting the publication relation between the author and the venue type. Note that, a single relation defined in the network schema can be viewed as a special case of meta-path, such as the citation relation $P \to P$.

7.2.1 The Meta-Path Selection Problem

Link-based clustering is to cluster objects based on their connections to other objects in the network. In a heterogeneous information network, we need to specify more information for a meaningful clustering. This includes (1) the type of objects to be clustered (called the **target type**), and (2) the type of connections, that is, meta-path(s), to use for the clustering task, and we call the object type that the target type is connecting to via the meta-path as the **feature type**. For example, when clustering authors based on the venues which they have published papers in, the target type is the author type, the meta-path to use is A - P - V, and the feature type is venue.

In a heterogeneous information network, target objects could link to many types of feature objects by multiple meta-paths. For example, authors could connect to other authors via meta-path A-P-A, or connect to terms via meta-path A-P-T. Meta-path selection is to determine which meta-paths or their weighted combination to use for a specific clustering task.

7.2.2 User-Guided Clustering

User guidance is critical for clustering objects in the network. In this study, we consider the guidance as user seeding objects in each cluster. For example, to cluster authors based on their (hidden) research areas, one can first provide several representative authors in each area. These seeds are used as guidance for clustering all the target objects in the network. More importantly, they provide information for selecting the most relevant meta-paths for the specific clustering task. Note that in practice, a user may not be able to provide seeds for *every* cluster, but only for *some* clusters they are most familiar with, which should be handled by the algorithm too.

7.2.3 The Problem Definition

Now we provide the problem definition of user-guided clustering via meta-path selection. Given a heterogeneous information network G, a user needs to specify the following as inputs for a clustering

task:

- 1. The target type for clustering, type T.
- 2. The number of clusters, K, and the object seeds for each cluster, say $\mathcal{L}_1, \ldots, \mathcal{L}_K$, where \mathcal{L}_k denotes the object seeds for cluster k, which could be an empty set. These seeds will be used as hints to learn the purpose/preference of the clustering task.
- 3. A set of M meta-paths starting from type T, denoted as $\mathcal{P}_1, \mathcal{P}_2, \dots, \mathcal{P}_M$, which might be helpful for the clustering task. These meta-paths can be determined either according to users' expert knowledge, or by traversing the network schema starting from type T with a length constraint.

For each meta-path \mathcal{P}_m , we calculate the adjacency matrix W_m , which we call relation matrix, between the target type T and the feature type F_m , by multiplying adjacency matrices for each relation along the meta-path. For example, the relation matrix W for meta-path A-P-V, denoting the number of papers published by an author in a venue, is calculated by $W = W_{AP} \times W_{PV}$, where W_{AP} and W_{PV} are the adjacency matrices for relation A-P and P-V respectively.

The output of the algorithm consists of two parts: (1) the weight $\alpha_m \geq 0$ of each meta-path \mathcal{P}_m for a particular clustering task, which should be consistent with the clustering result implied by the limited user guidance, and (2) the clustering result according to the user guidance and under the learned weights for each meta-path, that is, to associate each target object t_i in T with a K-dimensional soft clustering probability vector, $\boldsymbol{\theta}_i = (\theta_{i1}, \dots, \theta_{iK})$, where θ_{ik} is the probability of t_i belonging to cluster k, i.e., $\theta_{ik} \geq 0$ and $\sum_{k=1}^K \theta_{ik} = 1$.

7.3 The Probabilistic Model

A good clustering result is determined by several factors: First, the clustering result should be consistent with the link structure; second, it should also be consistent with the user guidance; and third, the importance of each meta-path is implied by the user-guided clustering, which should be modeled and learned to further enhance the clustering quality. In the following, we propose a probabilistic approach to model the problem in a unified framework, by considering all the three factors.

7.3.1 Modeling the Relationship Generation

To model the consistency between a clustering result and a relation matrix, we propose a clusteringbased generative model for relationship generation.

For a meta-path \mathcal{P}_m , let its corresponding relation matrix between the target type T and the feature type F_m be W_m . For each target object t_i , we model its relationships as generated from a mixture of multinomial distributions, where the probability of $t_i \in T$ connecting to $f_{j,m} \in F_m$ is conditionally independent on t_i given that the hidden cluster label of the relationship is known. Let $\pi_{ij,m} = P(j|i,m)$ be the generative probability of the relationship starting from t_i and ending at $f_{j,m}$, where $\sum_j \pi_{ij,m} = 1$, then

$$\pi_{ij,m} = P(j|i,m) = \sum_{k} P(k|i)P(j|k,m) = \sum_{k} \theta_{ik}\beta_{kj,m}$$
(7.1)

where $\theta_{ik} = P(k|i)$ denotes the probability of t_i belonging to cluster k and $\beta_{kj,m} = P(j|k,m)$ denotes the probability of $f_{j,m}$ appearing in cluster k. In other words, let $\pi_{i,m} = (\pi_{i1,m}, \dots, \pi_{i|F_m|,m})$ be the generative probability vector for target object t_i , then each $\pi_{i,m}$ can be factorized as a weighted summation of ranking distributions of feature objects in each cluster. The factorization idea is similar to that of PLSA [48], PHITS [30], and RankClus [100], but is built on meta-pathencoded relationships rather than immediate links. This extension will capture more and richer link-based features for clustering target objects in heterogeneous networks.

By assuming each target object t_i is independent with each other and each relationship generated by t_i is independent with each other, the probability of observing all the relationships between all the target objects and feature objects is the production of the probability of all the relationships following meta-path \mathcal{P}_m :

$$P(W_m|\Pi_m, \Theta, B_m) = \prod_i P(\mathbf{w}_{i,m}|\boldsymbol{\pi}_{i,m}, \Theta, B_m) = \prod_i \prod_j (\pi_{ij,m})^{w_{ij,m}}$$
(7.2)

where $\Pi_m = \Theta B_m$ is the probability matrix with cells as $\pi_{ij,m}$'s, Θ is the parameter matrix for θ_{ik} 's, B_m is the parameter matrix for $\beta_{kj,m}$'s, and $w_{ij,m}$ is the weight of the relationship between t_i and $f_{j,m}$. Note that, to model the relationship generation, each meta-path \mathcal{P}_m corresponds

to a different generative probability matrix Π_m . These probability matrices share the same soft clustering probabilities Θ , but they have different ranking distributions B_m in different meta-paths.

7.3.2 Modeling the Guidance from Users

Further, we take the user guidance in the form of object seeds for some clusters as the prior knowledge for the clustering result Θ , by modeling the prior as a Dirichlet distribution rather than treating them as hard labeled ones.

For each target object t_i , its clustering probability vector $\boldsymbol{\theta}_i$ is assumed to be a multinomial distribution, which is generated from some Dirichlet distribution. If t_i is labeled as a seed in cluster k^* , $\boldsymbol{\theta}_i$ is then modeled as being sampled from a Dirichlet distribution with parameter vector $\lambda \mathbf{e}_{k^*} + \mathbf{1}$, where \mathbf{e}_{k^*} is a K-dimensional basis vector, with the k^* th element as 1 and 0 elsewhere. If t_i is not a seed, $\boldsymbol{\theta}_i$ is then assumed as being sampled from a uniform distribution, which can also be viewed as a Dirichlet distribution with parameter vector $\mathbf{1}$. The density of $\boldsymbol{\theta}_i$ given such priors is:

$$P(\boldsymbol{\theta}_i|\lambda) \propto \begin{cases} \prod_k \theta_{ik}^{\mathbf{1}_{\{t_i \in \mathcal{L}_k\}}\lambda} = \theta_{ik^*}^{\lambda}, & \text{if } t_i \text{ is labeled and } t_i \in \mathcal{L}_{k^*}, \\ 1, & \text{if } t_i \text{ is not labeled.} \end{cases}$$
(7.3)

where $\mathbf{1}_{\{t_i \in \mathcal{L}_k\}}$ is an indicator function, which is 1 if $t_i \in \mathcal{L}_k$ holds, and 0 otherwise.

The hyper-parameter λ is a nonnegative value, which controls the strength of users' confidence over the object seeds in each cluster. From Equation (7.3), we can find that:

- when $\lambda = 0$, the prior for θ_i of a labeled target object becomes a uniform distribution, which means no guidance information will be used in the clustering process.
- when $\lambda \to \infty$, the prior for θ_i of a labeled target object converges to a point mass, i.e., $P(\theta_i = \mathbf{e}_{k^*}) \to 1$ or $\theta_i \to \mathbf{e}_{k^*}$, which means we will assign k^* as the hard cluster label for t_i .

In general, a larger λ indicates a higher probability that θ_i is around the point mass \mathbf{e}_{k^*} , and thus a higher confidence for the user guidance.

7.3.3 Modeling the Quality Weights for Meta-Path Selection

Different meta-paths may lead to different clustering results, therefore it is desirable to learn the quality of each meta-path for the specific clustering task. We propose to learn the quality weight for each meta-path by evaluating the consistency between its relation matrix and the user-guided clustering result.

In deciding the clustering result for target objects, a meta-path may be of low quality for the following reasons:

- 1. The relation matrix derived by the meta-path does not contain an inherent cluster structure.

 For example, target objects are connecting to the feature objects randomly.
- 2. The relation matrix derived by the meta-path itself has a good inherent cluster structure, however, it is not consistent with the user guidance. For example, in our motivating example, if the user gives a guidance as: $K = 2, \mathcal{L}_1 = \{1\}, \mathcal{L}_2 = \{2\}$, then the meta-path A - O - A should have a lower impact in the clustering process for authors.

The general idea of measuring the quality of each meta-path is to see whether the relation matrix W_m is consistent with the detected hidden clusters Θ and thus the generative probability matrix Π_m , which is a function of Θ , i.e., $\Pi_m = \Theta B_m$.

In order to quantify the weight for such quality, we model the weight α_m for meta-path \mathcal{P}_m as the relative weight for each relationship between target objects and feature objects following \mathcal{P}_m . In other words, we treat our observations of the relation matrix as $\alpha_m W_m$ rather than original W_m . A larger α_m indicates a higher quality and a higher confidence of the observed relationships, and thus each relationship should count more.

Then, we assume the multinomial distribution $\pi_{i,m}$ has a prior of Dirichlet distribution with parameter vector ϕ_i . In paticular, we consider a discrete uniform prior, which is a special case of Dirichlet distribution with parameters as an all-one vector, i.e., $\phi_{i,m} = 1$. The value of α_m is determined by the consistency between the observed relation matrix W_m and the generative probability matrix Π_m , which can be evaluated as how likely we can get Π_m given the relation matrix W_m and its quality weight α_m . The goal is then to find the α_m^* that maximizes the posterior probability of $\pi_{i,m}$ for all the target objects t_i , given the observation of relationships $\mathbf{w}_{i,m}$ with

relative weight α_m :

$$\alpha_m^* = \underset{\alpha_m}{\operatorname{arg\,max}} \prod_i P(\boldsymbol{\pi}_{i,m} | \alpha_m \mathbf{w}_{i,m}, \boldsymbol{\theta}_i, B_m)$$
 (7.4)

We can show that the posterior of $\pi_{i,m} = \theta_i B_m$ is another Dirichlet distribution with the updated parameter vector as $\alpha_m \mathbf{w}_{i,m} + \mathbf{1}$, according to the multinomial-Dirichlet conjugate:

$$\boldsymbol{\pi}_{i,m} | \alpha_m \mathbf{w}_{i,m}, \boldsymbol{\theta}_i, B_m \sim Dir(\alpha_m w_{ij,m} + 1, \dots, \alpha_m w_{i|F_m|,m} + 1)$$
 (7.5)

which has the following density function:

$$P(\boldsymbol{\pi}_{i,m}|\alpha_m \mathbf{w}_{i,m}, \boldsymbol{\theta}_i, B_m) = \frac{\Gamma(\alpha_m n_{i,m} + |F_m|)}{\prod_j \Gamma(\alpha_m w_{ij,m} + 1)} \prod_j (\pi_{ij,m})^{\alpha_m w_{ij,m}}$$
(7.6)

where $n_{i,m} = \sum_{j} w_{ij,m}$, the total number of path instances from t_i following meta-path \mathcal{P}_m . By modeling α_m in such a way, the meaning of α_m is quite clear:

- $\alpha_m w_{ij,m} + 1$ is the parameter of jth dimension for the new Dirichlet distribution.
- The larger α_m , the more likely it will generate a $\pi_{i,m}$ with a distribution as the observed relationship distribution, i.e., $\pi_{i,m} \to \mathbf{w_{i,m}}/n_{i,m}$ when $\alpha_m \to \infty$, where $n_{i,m}$ is the total number of path instances from t_i following meta-path \mathcal{P}_m .
- The smaller α_m , the more likely it will generate a π_i that is not relevant to the relation matrix W_m , and $\pi_{i,m}$ can be any $|F_m|$ -dimensional multinomial distribution.

Note that, we do not consider negative α_m 's in this model, which means that the relationships with a negative impact in the clustering process are not considered, and the extreme case of $\alpha_m = 0$ means that the relationships in a meta-path are totally irrelevant to the clustering process.

7.3.4 The Unified Model

Putting all the three factors together, we have the joint probability of observing the relation matrices with relative weights α_m 's, and the parameter matrices Π_m 's and Θ :

$$P(\{\alpha_m W_m\}_{m=1}^M, \Pi_{1:M}, \Theta | B_{1:M}, \Phi_{1:M}, \lambda)$$

$$= \prod_i \left(\prod_m P(\alpha_m W_m | \Pi_m, \boldsymbol{\theta}_i, B_m) P(\Pi_m | \Phi_m) \right) P(\boldsymbol{\theta}_i | \lambda)$$
(7.7)

where Φ_m is the Dirichlet prior parameter matrix for Π_m , and an all-one matrix in our case. We want to find the maximum a posteriori probability (MAP) estimate for Π_m 's and Θ , which maximizes the logarithm of posterior probability of $\{\Pi_m\}_{m=1}^M$, given the observations of relation matrices with relative weights $\{\alpha_m W_m\}_{m=1}^M$ and Θ , plus a regularization term over $\boldsymbol{\theta}_i$ for each target object denoting the logarithm of prior density of $\boldsymbol{\theta}_i$:

$$J = \sum_{i} \left(\sum_{m} \log P(\boldsymbol{\pi}_{i,m} | \alpha_{m} \mathbf{w}_{i,m}, \boldsymbol{\theta}_{i}, B_{m}) + \sum_{k} \mathbf{1}_{\{t_{i} \in \mathcal{L}_{k}\}} \lambda \log \theta_{ik} \right)$$
(7.8)

By substituting the posterior probability formula in Equation (7.6) and the factorization form for all $\pi_{i,m}$, we get the final objective function:

$$J = \sum_{i} \left(\sum_{m} \left(\sum_{j} \alpha_{m} w_{ij,m} \log \sum_{k} \theta_{ik} \beta_{kj,m} + \log \Gamma(\alpha_{m} n_{i,m} + |F_{m}|) - \sum_{j} \log \Gamma(\alpha_{m} w_{ij,m} + 1) \right) + \sum_{k} \mathbf{1}_{\{t_{i} \in \mathcal{L}_{k}\}} \lambda \log \theta_{ik} \right)$$

$$(7.9)$$

7.4 The Learning Algorithm

In this section, we introduce the learning algorithm, PathSelClus, for the model (Equation (7.9)) proposed in Section 7.3. It is a two-step iterative algorithm, where the clustering result Θ and the weights for each meta-path α mutually enhance each other. In the first step, we fix the weight vector α , and learn the best clustering results Θ under this weight. In the second step, we fix the clustering matrix Θ and learn the best weight vector α .

7.4.1 Optimize Clustering Result Given Meta-Path Weights

When α is fixed, the terms only involving α can be discarded in the objective function Equation (7.9), which is then reduced to:

$$J_1 = \sum_{m} \alpha_m \sum_{i} \sum_{j} w_{ij,m} \log \sum_{k} \theta_{ik} \beta_{kj,m} + \sum_{i} \sum_{k} \mathbf{1}_{\{t_i \in \mathcal{L}_k\}} \lambda \log \theta_{ik}$$
 (7.10)

The new objective function can be viewed as a weighted summation of the log-likelihood for each relation matrix under each meta-path, where the weight α_m indicates the quality of each meta-path, plus a regularization term over Θ representing the user guidance. Θ and the augmented parameter B_m 's can be learned using the standard EM algorithm, as follows.

• **E-step:** In each relation matrix, we use $z_{ij,m}$ to denote the cluster label for each relationship between a target object t_i and a feature object $f_{j,m}$. According to the generative process described in Section 7.3.1, a cluster k is first picked with probability θ_{ik} , and a feature object $f_{j,m}$ is picked with probability $\beta_{kj,m}$. The conditional probability of the hidden cluster label given the old Θ^{t-1} and B_m^{t-1} values is then:

$$p(z_{ij,m} = k | \Theta^{t-1}, B_m^{t-1}) \propto \theta_{ik}^{t-1} \beta_{kj,m}^{t-1}$$
(7.11)

• M-step: We have the updating formulas for Θ^t and B_m^t as:

$$\theta_{ik}^t \propto \sum_{m} \alpha_m \sum_{j} w_{ij,m} p(z_{ij,m} = k | \Theta^{t-1}, B_m^{t-1}) + \mathbf{1}_{\{t_i \in \mathcal{L}_k\}} \lambda$$
 (7.12)

$$\beta_{kj,m}^t \propto \sum_{i} \sum_{j} w_{ij,m} p(z_{ij,m} = k | \Theta^{t-1}, B_m^{t-1})$$
 (7.13)

From Equation (7.12), we can see that the clustering membership vector $\boldsymbol{\theta}_i$ for t_i is determined by the cluster labels of its relationships to all the feature objects in all the relation matrices. Besides, if t_i is labeled as a seed object in some cluster k^* , $\boldsymbol{\theta}_i$ is also determined by the label. The strength of impacts from these factors is determined by the weight of each meta-path α_m , and the strength of the cluster labels λ , where α_m 's are learned automatically by our algorithm, and λ is given by users.

7.4.2 Optimize Meta-Path Weights Given Clustering Result

Once a clustering result Θ and the augmented parameter B_m 's are given, we can calculate the generative probability matrix Π_m for each meta-path \mathcal{P}_m by: $\Pi_m = \Theta B_m$. By discarding the irrelevant terms, the objective function of Equation (7.9) can be reduced to:

$$J_2 = \sum_{i} \left(\sum_{m} \left(\sum_{j} \alpha_m w_{ij,m} \log \pi_{ij,m} + \log \Gamma(\alpha_m n_{i,m} + |F_m|) - \sum_{j} \log \Gamma(\alpha_m w_{ij,m} + 1) \right) \right)$$
(7.14)

It is easy to check that J_2 is a concave function, which means there is a unique α that maximizes J_2 . We use gradient descent approach to solve the problem, which is an iterative algorithm with the updating formula as: $\alpha_m^t = \alpha_m^{t-1} + \eta_m^t \left. \frac{\partial J_2}{\partial \alpha_m} \right|_{\alpha_m = \alpha_m^{t-1}}$. To guarantee the increase of J_2 , the step size η_m^t is usually set as a small enough number. By setting $\eta_m^t = \frac{\alpha_m^{t-1}}{-\sum_i \sum_j w_{ij,m} \log \pi_{ij,m}}$, following the trick used in non-negative matrix factorization (NMF) [62], we can get updating formula for α_m as:

$$\alpha_m^t = \alpha_m^{t-1} \frac{\sum_i \left(\psi(\alpha_m^{t-1} n_{im} + |F_m|) n_{i,m} - \sum_j \psi(\alpha_m^{t-1} w_{ij,m} + 1) w_{ij,m} \right)}{-\sum_i \sum_j w_{ij,m} \log \pi_{ij,m}}$$
(7.15)

which guarantees α_m^t a non-negative value, where $\psi(x)$ is the digamma function, the first derivative of $\log \Gamma(x)$. Also, by looking at the denominator of the formula, we can see that a larger log-likelihood of observing relationships $w_{ij,m}$ under model probability $\pi_{ij,m}$ (i.e., a smaller denominator as log-likelihood is negative) generally leads to a larger α_m . This is also consistent with the human intuition.

7.4.3 The PathSelClus Algorithm

Overall, the PathSelClus algorithm is an iterative algorithm that optimizes Θ and α alternatively. The optimization of Θ contains an inner loop of EM-algorithm, and the optimization of α contains another inner loop of gradient descent algorithm.

The Weight Setting of Relation Matrices Given a heterogeneous information network G, we calculate the relation matrix W_m for each given meta-path \mathcal{P}_m by multiplying adjacency matrices along the meta-path. It can be shown that, scaling W_m by a factor of $1/c_m$ leads to a scaling of the learned relative weight α_m by a factor of c_m . Therefore, the performance of the clustering result

will not be affected by the scaling of the relation matrix, which is a good property of our algorithm.

Initialization Issues. For the initial value of α , we set it as an all-one vector, which assumes all the meta-paths are equally important. For the initial value of Θ in the clustering step given α , if t_i is not labeled, we assign a random clustering vector to θ_i ; whereas if t_i is labeled as a seed for a cluster k^* , we assign $\theta_i = \mathbf{e}_k^*$.

Time Complexity Analysis. The PathSelClus algorithm is very efficient, as it is proportional to the number of relationships that are used in the clustering process, which is about linear to the number of target objects for short meta-paths in sparse networks. Formally, for the inner EM algorithm that optimizes Θ , the time complexity is $O(t_1(K\sum_m |\mathcal{E}_m| + K|T| + K\sum_m |F_m|)) = O(t_1(K\sum_m |\mathcal{E}_m|))$, where $|\mathcal{E}_m|$ is the number of non-empty relationships in relation matrix W_m , |T| and $|F_m|$ are the numbers of target objects and feature objects in meta-path \mathcal{P}_m , which are typically smaller than $|\mathcal{E}_m|$, and t_1 is the number of iterations. For the inner gradient descent algorithm, the time complexity is $O(t_2(\sum_m |\mathcal{E}_m|))$, where t_2 is the number of iterations. The total time complexity for the whole algorithm is then $O(t(t_1(K\sum_m |\mathcal{E}_m|) + t_2(\sum_m |\mathcal{E}_m|)))$, where t is the number of outer iterations, which usually is a small number. Such a processing efficiency has also be verified by our experiments.

7.5 Experiments

In this section, we compare *PathSelClus* with several baselines and show the effectiveness of our algorithm.

7.5.1 Datasets

We use two real information networks for performance test, the DBLP network and the Yelp network. For each network, we design multiple clustering tasks provided with different user guidance, which are introduced in the following.

1. **The DBLP Network.** For the DBLP network introduced early in the chapter, we design three clustering tasks in the following.

- DBLP-T1: Cluster conferences in the "four-area" dataset [103], which contains 20 major conferences and all the related papers, authors and terms in DM, DB, IR, and AI fields, according to the research areas of the conferences. The candidate meta-paths include: V P A P V and V P T P V.
- DBLP-T2: Cluster top-2000 authors (by their number of publications) in the "four-area" dataset, according to their research areas. The candidate meta-paths include: A P A, A P A P A, A P V P A, and A P T P A.
- DBLP-T3: Cluster 165 authors who have been ever advised by Christos Faloutsos, Michael
 I. Jordan, Jiawei Han, and Dan Roth (including these professors), according to their research
 groups. The candidate meta-paths are the same as in DBLP-T2.
- 2. **The Yelp Network.** For the Yelp network introduced early in the chapter, we are provided by Yelp a sub-network¹, which include 6900 businesses, 152327 reviews, and 65888 users. Hierarchical categories are provided for each business as well, such as "Restaurants," "Shopping" and so on. For Yelp network, we design three clustering tasks in the following.
 - Yelp-T1: We select 4 relatively big categories ("Health and Medical," "Food," "Shopping," and "Beauty and Spas"), and cluster 2224 businesses with more than one reviews according to two meta-paths: B R U R B and B R T R B.
 - Yelp-T2: We select 6 relatively big sub-categories under the first-level category "Restaurant" ("Sandwiches," "Thai," "American (New)," "Mexican," "Italian," and "Chinese"), and cluster 554 businesses with more than one reviews according to the same two meta-paths.
 - Yelp-T3: We select 6 relatively big sub-categories under the first-level category "Shopping" ("Eyewear & Opticians," "Books, Mags, Music and Video," "Sporting Goods," "Fashion," "Drugstores," and "Home & Garden"), and cluster 484 businesses with more than one reviews according to the same two meta-paths.

¹http://www.yelp.com/academic_dataset

7.5.2 Effectiveness Study

First, we study the effectiveness of our algorithm under different tasks, and compare it with several baselines.

Baselines

Three baselines are used for comparison studies. Since none of them has considered the meta-path selection problem, we will use all the meta-paths as features and prepare them to fit the input of each of these algorithms. The first is user-guided, information theoretic-based, k-means clustering (ITC), which is an adaption of seeded k-means algorithm proposed in [11], by replacing Euclidean distance to KL-divergence as used in information theoretic-based clustering algorithms [33, 6]. ITC is a hard clustering algorithm. For the input, we concatenate all the relation matrices side-by-side into one single relation matrix, and thus we get a very high dimensional feature vector for each target object.

The second baseline is the label propagation (LP) algorithm proposed in [135], which utilizes link structure to propagate labels to the rest of the network. For the input, we add all the relation matrices together to get one single relation matrix. As LP is designed for homogeneous networks, we confine our meta-paths to ones that start and end both in the target type. LP is a soft clustering algorithm.

The third baseline is the cluster ensemble algorithm proposed in [85], which can combine soft clustering results into a consensus, which we call ensemble_soft. Different from the previous two baselines that directly combine meta-paths at the input level, cluster ensemble combines the clustering results for different meta-paths at the output level. Besides, we also use majority voting as another baseline (ensemble_voting), which first maps each clustering result for each target object into a hard cluster label and then picks the cluster label that is the majority over different meta-paths. As we can use either ITC or LP as the clustering algorithm for each ensemble method, we get four ensemble baselines in total: ITC_soft, ITC_voting, LP_soft, and LP_voting.

Evaluation Methods

Two evaluation methods are used to test the clustering result compared with the ground truth, where the soft clustering is mapped into hard cluster labels. The first measure is accuracy, which is used when seeds are available for every cluster and is calculated as the percentage of target objects going to the correct cluster. Note that, in order to measure whether the seeds are indeed attracting objects to the right cluster, we do not map the outcome cluster labels to the given class labels. The second measure is normalized mutual information (NMI), which does not require the mapping relation between ground truth labels and the cluster labels obtained by the clustering algorithm. Both measures are in the range of 0 to 1, and a higher value indicates a better clustering result in terms of the ground truth.

Full Cluster Seeds

We first test the clustering accuracy when cluster seeds are given for every cluster. In this case, all the three baselines can be used and compared. Performances under different numbers of seeds in each cluster are tested. Each result is the average of 10 runs.

Table 7.1: Clustering accuracy for DBLP-T1 task.

#S	Measure	PathSelClus	LP	ITC	LP_voting	LP_soft	ITC_voting	ITC_soft
1	Accuracy	0.9950	0.6500	0.6900	0.6500	0.6650	0.6450	0.5100
	NMI	0.9906	0.6181	0.6986	0.6181	0.5801	0.5903	0.5316
2	Accuracy	1	0.7500	0.8450	0.7500	0.8200	0.8950	0.8700
	NMI	1	0.6734	0.7752	0.6734	0.7492	0.8321	0.7942

Table 7.2: Clustering accuracy for DBLP-T2 task.

#S	Measure	PathSelClus	LP	ITC	LP_voting	LP_soft	ITC_voting	ITC_soft
1	Accuracy	0.7951	0.2122	0.3284	0.2109	0.3529	0.2513	0.2548
1	NMI	0.6770	0.0312	0.1277	0.0267	0.0301	0.4317	0.4398
5	Accuracy	0.8815	0.2487	0.3223	0.5117	0.3685	0.3311	0.3495
9	NMI	0.6868	0.0991	0.1102	0.4402	0.0760	0.3092	0.4316
10	Accuracy	0.8863	0.5586	0.3694	0.4297	0.3880	0.4891	0.2969
	NMI	0.6947	0.4025	0.1261	0.1788	0.1148	0.4045	0.4204

Table 7.3: Clustering accuracy for DBLP-T3 task.

#S	Measure	PathSelClus	LP	ITC	LP_voting	LP_soft	ITC_voting	ITC_soft
1	Accuracy	0.8067	0.9273	0.5376	0.7091	0.5424	0.4770	0.2358
1	NMI	0.6050	0.7966	0.5120	0.5870	0.7182	0.3008	0.3416
	Accuracy	0.9036	0.9394	0.5285	0.7333	0.3267	0.5176	0.4085
2	NMI	0.7485	0.8283	0.5056	0.5986	0.8087	0.3898	0.3464
4	Accuracy	0.9248	0.9576	0.7624	0.7636	0.9255	0.6370	0.5485
	NMI	0.7933	0.8841	0.6280	0.6179	0.9057	0.4437	0.4634

The accuracy for all the 6 tasks for two networks are summarized in Table 7.1 through Table 7.3 and Table 7.4 through Table 7.6 respectively. From the results we can see that, *PathSelClus* performs the best in most of the tasks. Even for the task such as DBLP-T3 where other methods give the best clustering result, *PathSelClus* still gives clustering results among the top. This means, *PathSelClus* can give consistently good results across different tasks in different networks.

Also, by looking at the clustering accuracy trend along with the number of seeds used in each cluster, we can see that, more seeds generally leads to better clustering results.

Table 7.4: Clustering accuracy for Yelp-T1 task.

%S	Measure	PathSelClus	LP	ITC	LP_voting	LP_soft	ITC_voting	ITC_soft
1%	Accuracy NMI	0.5384	0.3381	0.2619	0.1632	0.1632	0.2564	0.2769
	NMI	0.5826	0.0393	0.0042	0.0399	0.0399	0.1907	0.2435
2%	$\frac{\text{Accuracy}}{\text{NMI}}$	0.5487	0.3444	0.2798	0.1713	0.1713	0.3581	0.3790
	NMI	0.5800	0.0557	0.0062	0.0567	0.0567	0.2281	0.2734
5%	Accuracy	0.5989	0.3732	0.3136	0.1965	0.1965	0.5215	0.5250
	NMI	0.5796	0.1004	0.0098	0.0962	0.0962	0.2583	0.2878

Table 7.5: Clustering accuracy for Yelp-T2 task.

%S	Measure	PathSelClus	LP	ITC	LP_voting	LP_soft	ITC_voting	ITC_soft
1%	Accuracy	0.7435	0.1137	0.1758	0.2112	0.2112	0.2430	0.2022
1/0	NMI	0.6517	0.0323	0.0178	0.0578	0.0578	0.2308	0.2490
2%	Accuracy	0.8004	0.1264	0.1910	0.2202	0.2202	0.2762	0.2792
470	NMI	0.6803	0.0487	0.0150	0.0801	0.0801	0.2099	0.2907
5%	Accuracy	0.8125	0.2653	0.2200	0.2437	0.2437	0.3049	0.3240
	NMI	0.6894	0.1111	0.0220	0.1212	0.1212	0.2252	0.2692

Table 7.6: Clustering accuracy for Yelp-T3 task.

%S	Measure	PathSelClus	LP	ITC	LP_voting	LP_soft	ITC_voting	ITC_soft
1%	Accuracy	0.4736	0.2789	0.1893	0.0682	0.0682	0.2593	0.1775
1/0	NMI	0.4304	0.0568	0.0155	0.0626	0.0626	0.1738	0.2065
2%	Accuracy	0.4597	0.4008	0.1948	0.0764	0.0764	0.2318	0.2033
2/0	NMI	0.4359	0.0910	0.0172	0.0755	0.0755	0.1835	0.1822
5%	Accuracy	0.4393	0.5351	0.2233	0.1033	0.1033	0.3337	0.3083
	NMI	0.4415	0.1761	0.0194	0.1133	0.1133	0.1793	0.2285

Partial Cluster Seeds

We then test the clustering accuracy when cluster seeds are only available for some of the clusters. We perform this study on DBLP-T3 using *PathSelClus*, which includes 4 clusters, and the results are shown in Figure 7.4. We can see that even if user guidance is only given to some clusters, those seeds can still be used to improve the clustering accuracy. In general, the fewer number of clusters with seeds, the worse the clustering accuracy, which is consistent with the human intuition. Note that, label propagation-based methods like LP cannot deal with partial cluster labels. However, in reality it is quite common that users are only familiar with some of the clusters and are only able to give good seeds in those clusters. That is another advantage of *PathSelClus*.

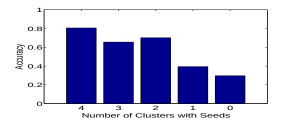


Figure 7.4: Clustering accuracy under partial guidance on DBLP-T3. Number of seeds provided by user for each cluster is 1 (#seeds = 1).

7.5.3 Case Study on Meta-Path Weights

One of the major contributions of *PathSelClus* is that it can select the right meta-paths for a user-guided clustering task. We now show the learned weights of meta-paths for some of the tasks.

In DBLP-T1 task, the total weight α_m for meta-path V-P-A-P-V is 1576, and the

average weight per relationship (a concrete path instance following the meta-path) is 0.0017. The total weight for meta-path V - P - T - P - V is 17001, whereas the average weight per relationship is 0.0003. This means that generally the relationships between two conferences that are connected by an author are more trustable than the ones that are connected by a term, which is consistent with human intuition since many terms can be used in different research areas and authors are typically more focused on confined research topics. However, as there are much more relationships following V - P - T - P - V than following V - P - A - P - V, the former overall provide more information for clustering.

In the Yelp network, similar to DBLP-T1 task, in terms of the average weight for each relationship, meta-path B-R-U-R-B is with higher weight than B-R-T-R-B; whereas in terms of total weight, meta-path B-R-T-R-B is with higher weight. An interesting phenomenon is that, for Yelp-T2 task, which tries to cluster restaurants into different categories, the average weight for relationships following B-R-U-R-B is 0.1716, much lower than the value (0.5864) for Yelp-T3 task, which tries to cluster shopping businesses into finer categories. This simply says that most users actually will try different kinds of food, therefore they will not be served as a good connection between restaurants as they are in other categories.

7.6 Related Work

Recently, there are many clustering algorithms proposed for networks, such as spectral clustering-based methods [90, 72], link-based probabilistic models [30, 3], modularity function-based algorithms [81, 80], and density-based algorithms [117, 113] on homogenous networks; and ranking-based algorithms [100, 103], non-negative matrix factorization [62, 112], spectral clustering-based methods [69], and probabilistic approaches [70] on heterogeneous networks. However, while all these clustering methods use the information given in the networks, none considers that different users may have different purposes for clustering, nor do they ask users to help select different information for link-based clustering. In this chapter, we show that different types of relationships encoded by meta-paths have different semantic meanings in determining the similarity between target objects, and the selection of these meta-paths should be done with user guidance in order to derive user-desired clustering results.

There are several lines of research on how to add user guidance to derive good clustering results, consistent with users' demand in vector space or networked data.

- Clustering with constraints. In [11, 12, 61], clustering algorithms that consider constraints either in the form of seeds in each cluster or pairwise constraints as must-link or cannot-link are proposed. A probabilistic model with an HMRF (hidden Markov random field) as a hidden layer that models the must-link and cannot-link between objects is proposed to solve the problem [12]. This approach can also be extended to graph data with the use of kernels instead of vector-based features [61]. However, these methods assume there is one trustable information source to either define the feature of each object or define the network structure between objects. The goal is to output the clustering result that is consistent with both the similarity defined by the data as well as the user guidance. In this chapter, we dig further and study which type of information source encoded with meta-paths is more trustable in a heterogeneous network.
- Semi-supervised learning on graphs. In [134, 135, 131], algorithms that propagate labels for a small portion of objects into the rest of the network are proposed, which are based on harmonic functions defined between objects using the network structure. Again, this kind of methods totally trust the given network and determine the best labels of the rest of the nodes according to the cost function defined on the network.
- Semi-supervised metric learning. In [17, 8], algorithms that learn the best distance metric functions according to the constrains for the clustering task are proposed. This line of problem is closer to the meta-path selection problem, but still differs significantly. First, they study features of objects in vector space instead of network; second, the metric functions should be given in an explicit format, which is very difficult to determine in a network scenario. In this chapter, we are not finding an explicit metric function that determines the similarity between any two target objects, instead, we model and learn the quality weight for each meta-path in the clustering process, which can be viewed as an implicit way to determine the similarity between two target objects.
- User-guided clustering in relational data. CrossClus [121] deals with another type of guidance from users: the attribute set of the target object type. The algorithm extracts a set of highly

relevant attributes in multiple relations connected via linkages defined in the database schema, and then use the whole attribute set as the feature set to apply traditional vector space-based clustering algorithm. CrossClus works for relational data with complete attributes, but not for purely link-based clustering.

Cluster ensemble [94, 85] is a method that combines clustering results of different methods or different datasets to a single consensus. Most of these cluster ensemble methods try to find a mean partition given different partitions of target objects. However, in reality these clusterings may conflict with each other, representing different purposes of clustering tasks, and a consensus does not necessarily lead to a clustering desired by users. In this study, we do not combine clustering results at the output level, but use intermediate clustering results as feedback to adjust the weight of each meta-path, and thus the clustering results and the quality weight for each meta-path can mutually enhance each other.

Our work also differs from traditional feature selection [43] and recently emerged semi-supervised feature selection [126, 116], which focus on vector space features, and do not have an immediate extension of solutions to our problem. For our meta-path selection problem, each meta-path provides a source of features instead of a concrete feature, and we have shown that simple combinations of features from different sources may lead to no good solution.

7.7 Discussions

The Power of Meta-Path Selection Different meta-paths in heterogeneous networks could be viewed as different sources of information for defining link-based similarity between objects. There are several ways to handle different meta-paths for a mining task such as clustering: (1) to combine them at relation matrix level, such as in baselines ITC and LP; (2) to combine the clustering results at the output level, such as in ensemble baselines; and (3) to learn and improve the quality weights for each meta-path iteratively, such as in *PathSelClus*. Only the third approach is able to select different meta-paths according to different clustering tasks, whereas the other two can only output an "average" clustering result using all the information. It turns out that, in most cases, the third approach is more flexible to combine information from different sources, and its advantage has been

shown in the experiment section.

Meta-Paths vs. Path Instances We now only consider the different semantics encoded by different meta-paths. In practice, different concrete paths (path instances) between two objects may also differ from each other. For example, two objects may be linked via a "bridge" or via a "hub," indicating different meanings. The difference between the two concepts: meta-path and path instance, is similar to the difference between a source of features and a concrete feature in a vector space. In this chapter, we have only discussed the selection of meta-paths. It is possible to select path instance at the object level, and the concrete method is left for future research.

7.8 Conclusion

Link-based clustering for objects in heterogeneous information networks is an important task with many applications. Different from traditional clustering tasks where similarity functions between objects are given and with no ambiguity, objects in heterogeneous networks can be connected via different relationships, encoded by different meta-paths. In this chapter, we integrate the meta-path selection problem with the user-guided clustering problem in heterogeneous networks. An algorithm PathSelClus that can utilize very limited guidance from users in the form of seeds in some of the clusters and automatically learn the best weights for each meta-path in the clustering process, is proposed. The experiments on different tasks on real datasets have demonstrated that our algorithm can output the most stable and accurate clustering results compared with the baselines. Also, the learned weights for each meta-path are very insightful to explain the hidden similarity between target objects under a particular clustering task.

Exploration of other types of user guidance, such as must-link and cannot-link, in meta-path selection for effective link-based clustering is an interesting topic for future study. More generally, meta-path selection problem exists in many other mining tasks, such as classification, ranking, relationship prediction and so on, which requires more future research on integrating meta-path selection with all these different mining tasks.

Chapter 8

Conclusion and Research Frontiers

In this thesis, we have proposed to use heterogeneous information networks to model real-world connected data, and introduced some general principles and methodologies for mining heterogeneous information networks. Although homogeneous networks are interesting subjects to study, real-world objects are usually connected via heterogeneous types of objects in complex ways, carrying critical information and rich data semantics, as shown in the examples like authors linking with papers and venues, and patients linking with diseases and treatments. Clearly, heterogeneous information networks preserve rich semantic information of the real-world data. Mining directly on heterogeneous information networks often leads to in-depth understanding of the relationships among different types of data and their regularities, models, patterns and anomalies, hence a deep insight of the networks, and fruitful mining results.

8.1 Conclusion

In the thesis, we have made the contributions on mining heterogeneous information networks in the following aspects.

- 1. We have proposed **three principles** on systematically mining heterogeneous information networks as follows.
 - Information propagation across heterogeneous types of nodes and links. Heterogeneous information network contains much richer and more complex information compared with projected homogeneous information networks. Objects from different types in the heterogeneous networks are interdependent, and knowledge can only be mined using the holistic information in a network. Thus, we need to study how to compute ranking scores, similarity

scores, cluster and class labels, across heterogeneous nodes and links. This is a principle that goes through all the chapters, which is mainly illustrated in Part I, including Chapter 2 and Chapter 3.

- Search and mining by exploring network meta structures. The network schema provides a meta structure of the information network. It provides guidance of search and mining of the network and help analyze and understand the semantic meaning of the objects and relations in the network. Meta-path-based similarity search and mining introduced in this thesis has demonstrated the usefulness and the power of exploring network meta structures. The second principle is mainly discussed in Part II, including Chapter 4 and Chapter 5.
- User-guided exploration of information networks. In a heterogeneous information network, there often exist numerous semantic relationships across multiple types of objects, carrying subtly different semantic meanings. It is desirable to automatically select the right relation (or meta-path) combinations with appropriate weights for a particular search or mining task based on user's guidance or feedback. User-guided or feedback-based network exploration is a useful strategy. The third principle is mainly discussed in Part III, including Chapter 6 and Chapter 7.
- 2. We have studied **different mining tasks** on heterogeneous information networks, which include ranking (Chapter 2 and Chapter 3), clustering (Chapter 2, Chapter 3, Chapter 6, and Chapter 7), similarity search (Chapter 4), relationship prediction (Chapter 5), user-guided clustering (Chapter 7), and relation strength-aware learning (Chapter 6 and Chapter 7). Some of these mining tasks are unique for heterogeneous information networks, such as relationship prediction and relation strength-aware mining problems. For other mining tasks, although they have also been studied in homogeneous information networks, new solutions are required in the new heterogeneous network scenario.
- 3. We have proposed **models** and **algorithms** to solve the above mining tasks in different applications, which include ranking-based clustering algorithms (RANKCLUS, Chapter 2, and NETCLUS, Chapter 3), meta-path-based similarity search algorithm (PathSim, Chapter 4), meta-path-based

relationship prediction (PathPredict and PathPredict_when, Chapter 5), relation strength-aware clustering with incomplete attributes (GenClus, Chapter 6), and user-guided clustering with meta-path selection (PathSelClus, Chapter 7).

8.2 Research Frontiers

Mining heterogeneous information networks is a young and promising research field. There are many unexplored territories and challenging research issues. Here we illustrate a few of them.

Constructing and Refining Heterogeneous Information Networks. Our study in most of the chapters assumes that a heterogeneous information network to be investigated contains a well-defined network schema and a large set of relatively clean and unambiguous objects and links. However, in the real world, things are more complicated.

A network extracted from a relational database may contain a well-defined schema which can be used to define the schema of its corresponding heterogeneous information network. Nevertheless, objects and links even in such a database-formed information network can still be noisy. For example, in the DBLP network, different authors may share the same name [120], that is, one node in a network may refer to multiple real-world entities; whereas in some other cases, different nodes in a network may refer to the same entity. Entity resolution will need to be integrated with network mining in order to merge and split objects or links and derive high quality results. Moreover, links in a network, roles of a node with respect to some other nodes may not be explicitly given. For example, the advisor-advisee relationship in the DBLP network [110] is not given, but such kind of relationships can be critical for understanding the growth of a research community or for some other data mining tasks. Furthermore, sometimes the connections between different nodes may not be reliable or trustable. For example, the author information for a book provided by an online book store could be erroneous or inaccurate. Multiple Web-sites may provide conflicting or compensating information for the properties of certain objects. Trustworthiness modeling [129] could be critically important for data cleaning, data integration, and quality network construction.

Construction of high-quality heterogeneous information networks becomes increasingly more challenging when we move away from relational databases towards increasingly more complicated,

unstructured data, from text documents, to online web-based systems, multimedia data, and multilingual data. Information extraction, natural language understanding, and many other information processing techniques should be integrated with network construction and analysis techniques to ensure high-quality information networks can be constructed and progressively refined so that quality mining can be performed on better-quality heterogeneous information networks.

Notice that entity extraction, data cleaning, detection of hidden semantic relationships, and trustworthiness analysis should be integrated with the network construction and mining processes to progressively and mutually enhance the quality of construction and mining of information networks.

Diffusion analysis in heterogeneous information networks. Diffusion analysis has been studied on homogeneous networks extensively, from the innovation diffusion analysis in social science [86] to obesity diffusion in health science [27]. However, in the real world, pieces of information or diseases are propagated in more complex ways, where different types of links may play different roles. For example, diseases could propagate among people, different kinds of animals and food, via different channels. Comments on a product may propagate among people, companies, and news agencies, via traditional news feeds, social media, reviews, and so on. It is highly desirable to study the issues on information diffusion in heterogeneous information networks in order to capture the spreading models that better represent the real world patterns.

Discovery and mining of hidden information networks. Although a network can be huge, a user at a time could be only interested in a tiny portion of nodes, links, or sub-networks. Instead of directly mining the entire network, it is more fruitful to mine hidden networks "extracted" dynamically from some existing networks, based on user-specified constraints or expected node/link behaviors. For example, instead of mining an existing social network, it could be more fruitful to mine networks containing suspects and their associated links; or mine subgraphs with nontrivial nodes and high connectivity. How to discover such hidden networks and how to mine knowledge (e.g., clusters, behaviors, and anomalies) from such hidden but non-isolated networks (i.e., still intertwined with the gigantic network in both network linkages and semantics) could be an interesting but challenging problem.

Discovery of application-oriented ontological structures in heterogeneous information networks. As shown in the studies on ranking-based clustering and ranking-based classification, interconnected, multiple typed objects in a heterogeneous information network often provide critical information for generating high quality, fine-level concept hierarchies. For example, it is often difficult to identify researchers just based on their research collaboration networks. However, putting them in a heterogeneous network that links researchers with their publication, conferences, terms and research papers, their roles in the network becomes evidently clear. Moreover, people may have different preferences over ontological structures at handling different kinds of tasks. For example, some people may be interested in the research area hierarchy in the DBLP network, whereas others may be interested in finding the author lineage hierarchy. How to incorporate user's guidance, and generate adaptable ontological structures to meet users's requirement and expectation could be an interesting and useful topic to study.

Online analytical processing of heterogeneous information networks. The power of online analytical processing (OLAP) has been shown in multidimensional analysis of structured, relational data. Similarly, users may like to view a heterogeneous information network from different
angles, in different dimension combinations, and at different levels of granularity. For example, in a
bibliographic network, by specifying the object type as paper and link type as citation relation, and
rolling up papers into research topics, we can immediately see the citation relationships between
different research topics and figure out which research topic could be the driving force for others.
However, the extension of the concept of online analysis processing (OLAP) to multi-dimensional
data analysis of heterogeneous information networks is nontrivial. Not only different applications
may need different ontological structures and concept hierarchies to summarize information networks but also because multiple pieces of semantic information in heterogeneous networks are
intertwined, determined by multiple nodes and links. There are some preliminary studies on this
issue, such as [107, 25, 130], but the large territories of online analytical processing of information
networks are still waiting to be explored.

Intelligent querying and semantic search in heterogeneous information networks. Given real-world data are interconnected, forming gigantic and complex heterogeneous information

networks, it poses new challenges to query and search in such networks intelligently and efficiently. Given the enormous size and complexity of a large network, a user is often only interested in a small portion of the objects and links most relevant to the query. However, objects are connected and inter-dependent on each other, how to search effectively in a large network for a given user's query could be a challenge. Similarity search that returns the most similar objects to a queried object, as studied in this thesis [99] and its follow-up [89], will serve as a basic function for semantic search in heterogeneous networks. Such kind of similarity search may lead to useful applications, such as product search in e-commerce networks and patent search in patent networks. Search functions should be further enhanced and integrated with many other functions. For example, structural search [123], which tries to find semantically similar structures given a structural query, may be useful for finding pattern in an e-commerce network involving buyers, sellers, products, and their interactions. Also, a recommendation system may take advantage of heterogeneous information networks that link among products, customers and their properties to make improved recommendations. Querying and semantic search in heterogeneous information networks opens another interesting frontier on research related to mining heterogeneous information networks.

Appendix A

Proofs of Theorems

Here are the proofs of the theorems introduced in the previous chapters.

Theorem 4.1: Properties of PathSim.

Proof. (1) $s(x_i, x_j) = \frac{2M_{ij}}{M_{ii} + M_{jj}} = \frac{2M_{ji}}{M_{ii} + M_{jj}} = s(x_j, x_i)$, as $M_{ij} = M_l(i, :) \cdot M_l(j, :) = M_l(j, :) \cdot M_l(i, :) = M_{ji}$, where \cdot means the dot product of two vectors.

(2) Let $M_l(i,:) = (a_1, a_2, \ldots, a_p)$, $M_l(j,:) = (b_1, b_2, \ldots, b_p)$, easy to see a_k, b_k are nonnegative for all $1 \le k \le p$, then $M_{ij} = \sum_{k=1}^p a_k b_k \ge 0$, $M_{ii} = \sum_{k=1}^p a_k^2 > 0$ (no dangling object), and $M_{jj} = \sum_{k=1}^p b_k^2 > 0$, therefore $s(x_i, x_j) \ge 0$; also, $\sum_{k=1}^p a_k^2 + \sum_{k=1}^p b_k^2 \ge 2 \sum_{k=1}^p a_k b_k$, with equality holding when $a_k = b_k$ for every k, therefore $s(x_i, x_j) \le 1$, and $s(x_i, x_i) = 1$.

(3)
$$M_{ij} = \sum_k a_k b_k \leq \sqrt{\sum_k a_k^2 \sum_k b_k^2} = \sqrt{M_{ii} M_{jj}}$$
 (by Cauchy-Schwarz inequality), then $s(x_i, x_j) \leq \frac{2}{\sqrt{M_{ii}/M_{jj}} + \sqrt{M_{jj}/M_{ii}}}$.

Theorem 4.2: Limiting behavior of PathSim under infinity length meta-path.

Proof. Since $M=(M_{\mathcal{P}}M_{\mathcal{P}}^T)$ is real symmetric, it can be decomposed as $M=PDP^T$, where D is a diagonal matrix with the values of eigenvalues of M, P is an orthogonal matrix composed of eigenvectors corresponding to eigenvalues in D. Let \mathbf{r} be the first column in P, then $M^k=PD^kP^T$. Let $s_{ij}^{(k)}=\frac{2M^k(i,j)}{M^k(i,i)+M^k(j,j)},\ \lambda_1$ be the largest eigenvalue of M, then $s_{ij}^{(k)}=\frac{2(PD^kP^T/\lambda_1^k)(i,j)}{(PD^kP^T(i,i)+PD^kP^T(j,j))/\lambda_1^k},$ and $\lim_{k\to\infty}s_{ij}^{(k)}=\frac{2\mathbf{r}(i)\mathbf{r}(j)}{\mathbf{r}(i)\mathbf{r}(i)+\mathbf{r}(j)\mathbf{r}(j)}$.

Theorem 4.3: Bounds for block-based similarity measure approximation.

Proof. 1.
$$\sum_{y \in C_v} s(x,y) = \sum_{y \in C_v} \frac{2\mathbf{x}^T \mathbf{y}}{D(x) + D(y)} \leq \frac{2\mathbf{x}^T \sum_{y \in C_v} \mathbf{y}}{D(x) + 1} = \sum_u \frac{2\mathbf{x}(R_u)^T \sum_{y \in C_v} \mathbf{y}(R_u)}{D(x) + 1}$$
$$\leq \sum_u \frac{2\hat{\mathbf{x}}_1(u)T(u,v)}{D(x) + 1} = \frac{2\hat{\mathbf{x}}_1^T T(:,v)}{D(x) + 1}, \text{ since according to Holder's Inequality, } \mathbf{a}^T \mathbf{b} \leq ||\mathbf{a}||_{\infty} ||\mathbf{b}||_1.$$
$$2. \ s(x,y) = \frac{2\mathbf{x}^T \mathbf{y}}{D(x) + D(y)} = \frac{2\sum_u \mathbf{x}(R_u)^T \mathbf{y}(R_u)}{D(x) + D(y)}. \text{ Since } \mathbf{a}^T \mathbf{b} \leq ||\mathbf{a}||_2 ||\mathbf{b}||_2 \text{ according to Cauchy-Schwarz inequality, then the above formula} \leq \frac{2\sum_u \hat{\mathbf{x}}_2(u)TT_1(u,y)}{D(x) + D(y)} = \frac{2\hat{\mathbf{x}}_2^T TT_1(:,y)}{D(x) + D(y)}.$$

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