Constructing Knowledge Graphs and Their Biomedical Applications

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Abstract

Knowledge graphs can support many biomedical applications. These graphs represent biomedical concepts and relationships in the form of nodes and edges. In this review, we discuss how these graphs are constructed and applied with a particular focus on how machine learning approaches are changing these processes. Biomedical knowledge graphs have often been constructed by integrating databases that were populated by experts via manual curation, but we are now seeing a more robust use of automated systems. A number of techniques are used to represent knowledge graphs, but often machine learning methods are used to construct a low-dimensional representation that can support many different applications. This representation is designed to preserve a knowledge graph's local and/or global structure. Additional machine learning methods can be applied to this representation to make predictions within genomic, pharmaceutical, and clinical domains. We frame our discussion first around knowledge graph construction and then around unifying relational learning techniques and unifying applications. Advances in machine learning for biomedicine are creating new opportunities across many domains, and we note potential avenues for future work with knowledge graphs that appear particularly promising.

Introduction

Knowledge graphs are a practical resource for many real world applications. They have been used in social media mining to classify nodes [1] and create recommendation systems [2]. These graphs have also been used in natural language processing to interpret simple questions and use relational information to provide answers [3,4]. In a biomedical setting these graphs have been used to prioritize genes relevant to disease [5,6,7,8], perform drug repurposing [9] and identify drug-target interactions [10].

Despite their utility, precisely defining a knowledge graph is a difficult task because there are multiple conflicting definitions [11]. For this review, we define a knowledge graph as the following: a resource that integrates single or multiple sources of information into the form of a graph. This graph allows for the capacity to make semantic interpretation, continuously incorporate new information and uncover novel hidden knowledge through computational techniques and algorithms. Based on this definition resources like Hetionet [9] would be considered a knowledge graph as Hetionet integrates multiple sources of information into the form of a graph (example shown in Figure 1). Hetionet was used to derive novel information concerning unique drug treatments [9]. We do not consider databases like DISEASES [12] and DrugBank [13] to be knowledge graphs. Although these resources contain essential information, they do not represent their data in the form of a graph.

Knowledge graphs are often constructed from manually curated databases [9,14,15,16]. These databases provide previously established information that can be incorporated into a graph. For example, a graph using DISEASES [12] as a resource would have genes and diseases as nodes, while edges added between nodes would represent an association between a gene and a disease. This example shows a single type of relationship; however, there are graphs that use databases with multiple relationships [17,9]. In addition to manual curation other approaches have used natural language processing techniques to construct knowledge graphs [18,19]. One example used a text mining system to extract sentences that illustrate a protein interacts with another protein [20]. Once identified, these sentences can be incorporated as evidence to establish an edge in a knowledge graph.

In this review we describe various approaches for constructing and applying knowledge graphs in a biomedical setting. We discuss the pros and cons of constructing a knowledge graph via manually curated databases and via text mining systems. We also compare assorted approaches for applying

knowledge graphs to solve biomedical problems. Lastly, we conclude on the practicality of knowledge graphs and point out future applications that have yet to be explored.

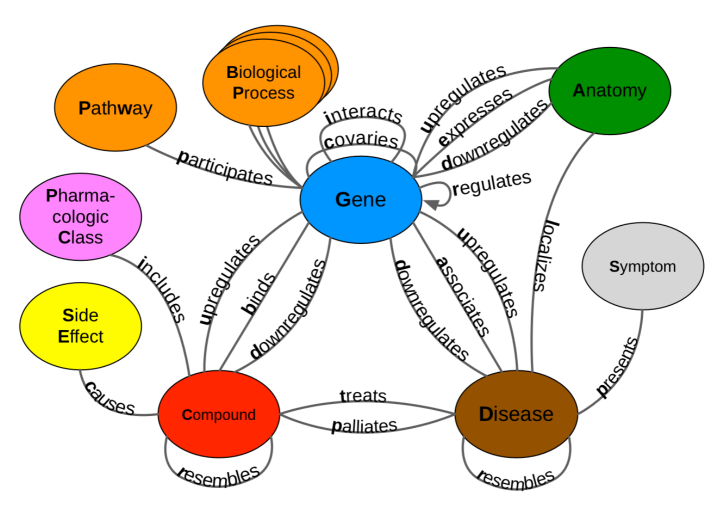


Figure 1: A metagraph (schema) of the heterogeneous network used in the Rephetio project [9]. This undirected network depicts pharmacological and biomedical information. The nodes (circles) represent entities and edges (lines) depict relational information between two entities.

Building Biomedical Knowledge Graphs

Knowledge graphs can be constructed in many ways using resources such as pre-exisitng databases or text. Usually, knowledge graphs are constructed using pre-existing databases. These databases are constructed by domain experts using approaches ranging from manual curation to automated techniques, such as text mining. Manual curation is a time consuming process that requires domain experts to read papers and annotate sentences that assert a relationship. Automated approaches rely on machine learning or natural language processing techniques to rapidly detect sentences of interest. We categorize these automated approaches into the following groups: rule-based extraction, unsupervised machine learning, and supervised machine learning and discuss examples of each type of approach while synthesizing their strengths and weaknesses.

Constructing Databases and Manual Curation

Database construction dates back all the way to 1956 where the first database contained a protein sequence of the insulin molecule [21]. The process of database construction involves gathering relevant text such as journal articles, abstracts, or web-based text and having curators read the gathered text to detect sentences that implicate a relationship (i.e., relationship extraction). Notable databases constructed by this process can be in found in Table 1. An example database, COSMIC [22] was constructed by a group of domain experts scanning the literature for key cancer related genes.

This database contained approximately 35M entries in 2016 [22] and by 2018 had grown to 45M entries [23]. Studies have shown that databases constructed in this fashion contain relatively precise data but the recall is low [24,25,26,27,28,29,30]. This happens because the publication rate is too high for curators to keep up [31]. This bottleneck highlights a critical need for future approaches to scale fast enough to compete with the increasing publication rate.

Semi-automatic methods are a way to accelerate the curation process [28,32,33,34,35,36,37]. The first step of these methods is to use an automated system to initially extract sentences from text. This process removes irrelevant sentences, which dramatically decreases the amount of text that curators must sift through. Following the pre-filtering step, curators then approve or reject the remaining sentences. This approach saved curators an average of 2-2.8 hours compared to manual efforts [32,38]. Despite automated systems excelling in identifying sentences for commonly occurring relationships, they tend to miss lesser-known relationships [32]. These systems also have a hard time parsing ambiguous sentences that naturally occur in text, which makes correcting them a challenging task [32]. Given these caveats, future approaches should look into using techniques that simplify sentences to solve the ambiguity issue [39,40].

Despite the negatives of manual curation, it is still an essential process for extracting relationships from text. This process can be used to generate gold standard datasets that automated systems use for validation [41,42] and can be used during the training process of these systems (i.e., active learning) [43]. It is important to remember that manual curation alone is precise, but results in low recall rates [30]. Future databases should consider initially relying on automated methods to obtain sentences at an acceptable recall level, then incorporate manual curation as a way to fix or remove irrelevant results.

Table 1: A table of databases that used a form of manual curation to populate entries. Reported number of entities and relationships are relative to time of publication.

Database [Reference]	Short Description	Number of Entries	Entity Types	Relationship Types	Method of Population
BioGrid [44]	A database for major model organisms. It contains genetic and proteomic information.	572,084	Genes, Proteins	Protein-Protein interactions	Semi-automatic methods
Comparative Toxicogenomics Database [45]	A database that contains manually curated chemical-genedisease interactions and relationships.	2,429,689	Chemicals (Drugs), Genes, Diseases	Drug-Genes, Drug-Disease, Disease-Gene mappings	Manual curation and Automated systems
Comprehensive Antibiotic Resistance Database [46]	Manually curated database that contains information about the molecular basis of antimicrobial resistance.	174,443	Drugs, Genes, Variants	Drug-Gene, Drug- Variant mappings	Manual curation
COSMIC [<u>22</u>]	A database that contains high resolution human cancer genetic information.	35,946,704	Genes, Variants, Tumor Types	Gene-Variant Mappings	Manual Curation

Database [Reference]	Short Description	Number of Entries	Entity Types	Relationship Types	Method of Population
Entrez-Gene [<u>47</u>]	NCBI's Gene annotation database that contains information pertaining to genes, gene's organism source, phenotypes etc.	7,883,114	Genes, Species and Phenotypes	Gene-Phenotypes and Genes- Species mappings	Semi-automated curation
OMIM [<u>48</u>]	A database that contains phenotype and genotype information	25,153	Genes, Phenotypes	Gene-Phenotype mappings	Manual Curation
PharmGKB [<u>49</u>]	A database that contains genetic, phenotypic, and clinical information related to pharmacogenomi c studies.	43,112	Drugs, Genes, Phenotypes, Variants, Pathways	Gene- Phenotypes, Pathway-Drugs, Gene-Variants, Gene-Pathways	Manual Curation and Automated Methods
UniProt [<u>50</u>]	A protein protein interaction database that contains proteomic information.	560,823	Proteins, Protein sequences	Protein-Protein interactions	Manual and Automated Curation

Text Mining for Relationship Extraction

Rule-Based Relationship Extraction

Rule based-extraction consists of identifying essential keywords and grammatical patterns to detect relationships of interest. Keywords are established via expert knowledge or though the use of pre-existing ontologies, while grammatical patterns are constructed via experts curating parse trees. Parse trees are tree data structures that depict a sentence's grammatical structure and come into two forms: a constituency parse tree (Figure 2) and a dependency parse tree (Figure 3). Both trees use part of speech tags, labels that dictate the grammatical role of a word such as noun, verb, adjective, etc, for construction, but represent the information in two different forms. Constituency parse trees breaks a sentence down into a subphrases (Figure 2) while dependency path trees analyzes the grammatical structure of a sentence (Figure 3). Many text mining approaches [51,52,53] use such trees to generate features for machine learning algorithms and these approaches are discussed in later sections. In this section we focus on approaches that use rule based extraction as a primary strategy to detect sentences that allude to a relationship.

Grammatical patterns can simplify sentences for easy extraction [40,54]. Jonnalagadda et al. used a set of grammar rules inspired by constituency trees to reshape complex sentences with simpler versions [40] and these simplified versions were manually curated to determine the presence of a relationship. By simplifying sentences this approach achieved high recall, but had low precision [40]. Other approach used simplification techniques to make extraction easier [55,56,57,58]. Tudor et al. simplified sentences to detect protein phosphorylation events [57]. Their sentence simplifier broke complex sentences that contain multiple protein events into smaller sentences that contain only one

distinct event. By breaking these sentences down the authors were able to increase their recall; however, sentences that contained ambiguous directionality or multiple phosphorylation events were too complex for the simplifier. As a consequence the simplifier missed some relevant sentences [57]. These errors highlight a crucial need for future algorithms to be generalizable enough to handle various forms of complex sentences.

Pattern matching is a fundamental approach used to detect relationship asserting sentences. These patterns can consist of phrases from constituency trees, a set of keywords or some combination of both [28,59,60,61,62,63]. Xu et al. designed a pattern matcher system to detect sentences in PubMed abstracts that indicate drug-disease treatments [62]. This system matched drug-disease pairs from ClinicalTrials.gov to drug-disease pairs mentioned in abstracts. This matching process aided the authors in identifying sentences that can be used to create simple patterns, such as "Drug in the treatment of Disease" [62], to match other sentences in a wide variety of abstracts. The authors hand curated two datasets for evaluation and achieved a high precision score of 0.904 and a low recall score of 0.131 [62]. This low recall score was based on constructed patterns being too specific to detect infrequent drug pairs. Besides constituency trees, some approaches used dependency trees to construct patterns [51,64]. Depending upon the nature of the algorithm and text, dependency trees could be more appropriate than constituency trees and vise versa. The performance difference between the two trees remains as an open question for future exploration.

Rules based methods provide a basis for many relationship extraction systems. Approaches in this category range from simplifying sentences for easy extraction to identifying sentences based on matched key phrases or grammatical patterns. Both require a significant amount of manual effort and expert knowledge to perform well. A future direction is to develop ways to automatically construct these hand-crafted patterns, which would accelerate the process of creating these rule-based systems.

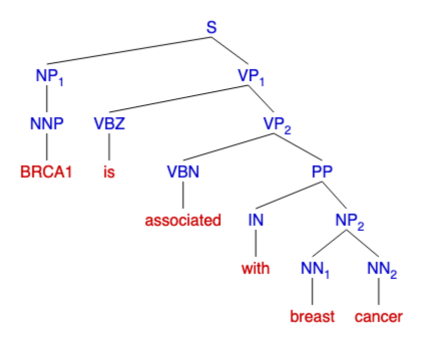


Figure 2: A visualization of a constituency parse tree using the following sentence: "BRCA1 is associated with breast cancer" [65]. This type of tree has the root start at the beginning of the sentence. Each word is grouped into subphrases depending its correlating part of speech tag. For example, the word "associated" is a past participle verb (VBN) that belongs to the verb phrase (VP) subgroup.

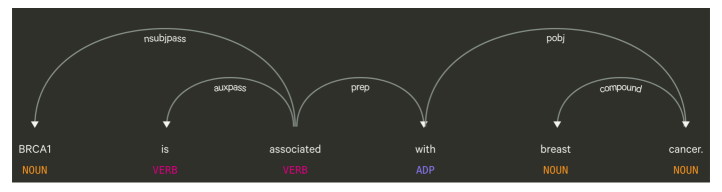


Figure 3: A visualization of a dependency parse tree using the following sentence: "BRCA1 is associated with breast cancer" [66]. For these type of trees the root begins with the main verb of the sentence. Each arrows represents the dependency shared between two words. For example, the dependency between BRCA1 and associated is nsubjpass, which stands for passive nominal subject. This means that "BRCA1" is the subject of the sentence and it is being referred to by the word "associated".

Extracting Relationships Without Labels

Unsupervised extractors draw inferences from textual data without the use of annotated labels. These methods involve some form of clustering or statistical calculations. In this section we focus on methods that use unsupervised learning to extract relationships from text.

An unsupervised extractor can exploit the fact that two entities may appear together in text. This event is referred to as co-occurrence and studies that use this phenomenon can be found in Table 2. Two databases DISEASES [12] and STRING [67] were populated using a co-occurrence scoring method on PubMed abstracts, which measured the frequency of co-mention pairs within individual sentences as well as the abstracts themselves. This technique assumes that each individual co-occurring pair is independent from one another. Under this assumption mention pairs that occur more than expected were presumed to implicate the presence of an association or interaction. This approach was able to identify 543,405 disease gene associations [12] and 792,730 high confidence protein protein interactions [67], but is limited to only using PubMed abstracts.

Full text articles are able to dramatically enhance relationship detection [68,69]. Westergaard et al. used a co-occurrence approach, similar to DISEASES [12] and STRING [67], to mine full articles for protein-protein interactions and other protein related information [68]. The authors discovered that full text provided better prediction power than using abstracts alone, which suggests that future text mining approaches should consider using full text to increase detection power.

Unsupervised extractors often treat different biomedical relationships as multiple isolated problems. An alternative to this perspective is to capture all different types at once. Clustering is an approach that accomplish this concept of simultaneous extraction. Percha et al. used a biclustering algorithm on generated dependency parse trees to group sentences within PubMed abstract [70]. Each cluster was manually curated to determine which relationship each group represented. This approach captured 4,451,661 dependency paths for 36 different groups [70]. Despite the success, this approach suffered from technical issues such as dependency tree parsing errors. These errors resulted in some sentences not being captured by the clustering algorithm [70] and future clustering approaches should consider simplifying sentences to prevent this type of issue.

Overall unsupervised methods provide a means to rapidly extract relationship asserting sentences without the need of annotated text. Approaches in this category range from calculating co-occurrence scores to clustering sentences and provide a generalizable framework that can be used on large repositories of text. Full text has already been show to meaningfully improve the performance of methods that aim to infer relationships using cooccurrences [68], and we should expect similar

benefits for machine learning approaches. Furthermore, we expect that simplifying sentences would improve unsupervised methods and should considered as an initial preprocessing step.

Table 2: Table of approaches that mainly use a form of co-occurrence.

Study	Relationship of Interest	
CoCoScore [71]	Protein-Protein Interactions, Disease-Gene and Tissue- Gene Associations	
Rastegar-Mojarad et al. [72]	Drug Disease Treatments	
CoPub Discovery [73]	Drug, Gene and Disease interactions	
Westergaard et al. [68]	Protein-Protein Interactions	
DISEASES [12]	Disease-Gene associations	
STRING [74]	Protein-Protein Interactions	
Singhal et al. [75]	Genotype-Phenotype Relationships	

Supervised Relationship Extraction

Supervised extractors use labeled sentences to construct generalized patterns that bisect positive examples (sentences that allude to a relationship) from negative ones (sentences that do not allude to a relationship). Most of these approaches have flourished due to pre-labelled publicly available datasets (Table 3). These datasets were constructed by curators for shared open tasks [76,77] or as a means to provide the scientific community with a gold standard [77,78,79]. Approaches that use these available datasets range from using linear classifiers such as support vector machines (SVMs) to non-linear classifiers such as deep learning techniques. The rest of this section discuss approaches that use supervised extractors to detect relationship asserting sentences.

Some supervised extractors involve mapping textual input onto a high dimensional space. SVMs are a type of classifier that can accomplish this task with a mapping function called a kernel [53,80]. These kernels take information such as a sentence's dependency tree [51,52], part of speech tags [53] or even word counts [80] and map them onto a dense feature space. Within this space, these methods construct a hyperplane that separates sentences in the positive class (illustrates a relationship) from the negative class (does not illustrate a relationship). Kernels can be manually constructed or selected to cater to the relationship of interest [52,53,80,80]. Determining the correct kernel is a nontrivial task that requires expert knowledge to be successful. In addition to single kernel methods, a recent study used an ensemble of SVMs to extract disease-gene associations [81]. This ensemble outperformed notable disease-gene association extractors [64,82] in terms of precision, recall and F1 score. Overall, SVMs have been shown to be beneficial in terms of relationship mining; however, major focus have shifted to utilizing deep learning techniques as these approaches can perform nonlinear mappings of high dimensional data.

Deep learning is an increasingly popular class of techniques that can construct their own features within a high dimensional space [83,84]. These methods amount to different forms of neural networks, such as recurrent or convolutional neural networks, to perform classification.

Recurrent neural networks (RNN) are designed for sequential analysis and use a repeatedly updating hidden state to make predictions. An example of a recurrent neural network is a long short term memory (LSTM) network [85]. Cocos et al. [86] used a LSTM to extract drug side effects from deidentified twitter posts, while Yadav et al. [87] used an LSTM to extract protein-protein interactions. Others have also embraced LSTMs to perform relationship extraction [86,88,89,90,91]. Despite the success of these networks, training can be difficult as these networks are highly susceptible to vanishing and exploding gradients [92,93]. One proposed solution to this problem is to clip the

gradients while the neural network trains [94]. Besides the gradient problem, these approaches only peak in performance when the datasets reach at least a tens of thousand of data points [95].

Convolutional neural networks (CNNs), which are widely applied for image analysis, use multiple kernel filters to capture small subsets of an overall image [84]. In the context of text mining an image is replaced with words within a sentence mapped to dense vectors (i.e., word embeddings) [96,97]. Peng et al. used a CNN to extract sentences that mentioned protein-protein interactions [98] and Zhou et al. used a CNN to extract chemical-disease relations [99]. Others have used CNNs and variants of CNNs to extract relationships from text [100,101,102]. Just like RNNs, these networks perform well when millions of labeled examples are present [95]; however, obtaining these large datasets is a non-trivial task. Future approaches that use CNNs or RNNs should consider solutions to obtaining these large quantities of data through means such as weak supervision [103], semi-supervised learning [104] or using pre-trained networks via transfer learning [105,106].

Semi-supervised learning [104] and weak supervision [103] are techniques that can rapidly construct large datasets for machine learning classifiers. Semi-supervised learning trains classifiers by combining labeled data with unlabeled data. For example, one study used a variational auto encoder with a LSTM network to extract protein-protein interactions from PubMed abstracts and full text [107]. This is an elegant solution for the small dataset problem, but requires labeled data to start. This dependency makes finding under-studied relationships difficult as one would need to find or construct examples of the missing relationships at the start.

Weak or distant supervision takes a different approach that uses noisy or even erroneous labels to train classifiers [103,108,109,110]. Under this paradigm sentences are labeled based on their mention pair being present (positive) or absent (negative) in a database and, once labeled, machine learning classifier can now be trained to extract relationships from text [103]. For example, Thomas et al. [111] used distant supervision to train a SVM to extract sentences mentioning protein-protein interactions (ppi). Their SVM model achieved comparable performance against a baseline model; however, the noise generated via distant supervision was difficult to eradicate [111]. A number of efforts have focused on combining distant supervision with other types of labeling strategies to mitigate the negative impacts of noisy knowledge bases [112,113,114]. Nicholson et al. [102] found that, in some circumstances, these strategies can be reused across different types of biomedical relationships to learn a heterogeneous knowledge graph in cases where those relationships describe similar physical concepts. Combining distant supervision with other types of labeling strategies remains an active area of investigation with numerous associated challenges and opportunities. Overall, semi-supervised learning and weak supervision provide promising results in terms of relationship extraction and future approaches should consider using these paradigms to train machine learning classifiers.

Table 3: A set of publicly available datasets for supervised text mining.

Dataset	Type of Sentences
AlMed [<u>42</u>]	PPI
BioInfer [<u>115</u>]	PPI
LLL [<u>116</u>]	PPI
IEPA [117]	PPI
HPRD5 [<u>78</u>]	PPI
EU-ADR [41]	DaG
BeFree [<u>82</u>]	DaG
CoMAGC [79]	DaG

Dataset	Type of Sentences	
CRAFT [118]	DaG	
Biocreative V CDR [77]	Compound induces Disease	
Biocreative IV ChemProt [76]	CbG	

Applying Knowledge Graphs to Biomedical Challenges

Knowledge graphs can help researchers tackle many biomedical tasks such as finding new treatments for existing drugs [9], aiding efforts to diagnose patients [119] and predicting associations between diseases and biomolecules [120]. In many cases, solutions rely on representing knowledges graphs in a low dimensional space, which is a process called representation learning. This space preserves a knowledge graph's local and/or global structure and can support efforts to apply machine learning methods to make predictions. In the next sections we review the unifying techniques that construct this low dimensional space and unifying applications that use this space to solve biomedical problems.

Unifying Techniques

Mapping high dimensional data into a low dimensional space greatly improves modeling performance in fields such as natural language processing [96,97] and image analysis [121]. The success of these approaches provides rationale for representing knowledge graphs into a low dimensional space [122]. Techniques that construct this representation often require information on how nodes are connected with one another [123,124,125,126], while other approaches can work directly with the edges themselves [127]. We group these methods into the following three categories: matrix factorization, translational methods, and deep learning (Figure 4).

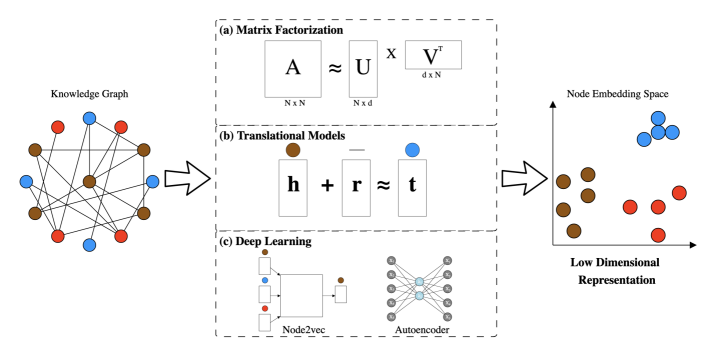


Figure 4: Pipeline for representing knowledge graphs in a low dimensional space. Starting with a knowledge graph, this space can be generated using one of the following options: Matrix Factorization (a), Translational Models (b) or Deep Learning (c). The output of this pipeline is an embedding space that clusters similar node types together.

Matrix Factorization

Matrix factorization is a technique that uses linear algebra to map high dimensional data onto a low dimensional space. This projection is accomplished by decomposing a matrix into a set of small rectangular matrices (Figure 4 (a)). Notable methods for matrix decomposition include Isomap [128], Laplacian eigenmaps [129] and Principal Component Analysis (PCA) [130]/Singular Vector Decomposition (SVD) [131]. These methods were designed to be used on many different types of data; however, we discuss their use in the context of representing a knowledge graphs in a low dimensional space.

SVD [131] is an algorithm that uses matrix factorization to portray knowledge graphs in a low dimensional space. The input for this algorithm is an adjacency matrix (A), which is a square matrix where rows and columns represent nodes and each entry represents the presence of an edge between two nodes. This matrix (A) gets decomposed into three parts: a square matrix Σ and a set of two small rectangular matrices U and V^T . This values within Σ are called singular values, which are akin to eigenvalues [131]. Each row in U and each column in V^T represents nodes within a low dimensional space [130,131]. In practice U is usually used to represent nodes in a knowledge graph, but V^T can also be used [131,132]. Typically, SVD appears in recommendation systems via collaborative filtering [133]; however, this technique can also be used as a standalone baseline to compare to other approaches [134].

Laplacian eigenmaps is a technique that assumes there is low dimensional structure in a high dimensional space [129]. This algorithm preserves this structure while projecting data into a low dimensional space. Typically, the first step of this algorithm is to construct a figurative knowledge graph where nodes represent datapoints and edges are constructed based on similarity of two datapoints; however, in this context, the knowledge graph has already been defined. The next step in this algorithm is to obtain both an adjacency matrix (A) and a degree matrix (D) from the knowledge graph. A degree matrix is a diagonal matrix where each entry represents the number of edges connected to a node. The adjacency and degree matrices are converted into a laplacian matrix (L), which is a matrix that shares the same properties as the adjacency matrix. The laplacian matrix is generated by subtracting the adjacency matrix from the degree matrix (L=D-A) and, once constructed, the algorithm uses linear algebra to calculate the laplacian's eigenvalues and eigenvectors ($Lx = \lambda Dx$). The generated eigenvectors represent the knowledge graph's nodes represented in a low dimensional space [129]. Other efforts have used variants of this algorithm to construct their own low dimensional representations of knowledge graphs [123,124,135]. Typically, eigenmaps work well when knowledge graphs have a sparse number of edges between nodes but struggle when presented with denser networks [134,135,136]. An open area of exploration is to adapt these methods to accommodate knowledge graphs that have a large number of edges.

Matrix factorization is a powerful technique that uses a matrices such as an adjacency matrix as input. Common approaches involve using SVD, Laplacian eigenmaps or variants of the two to construct low dimensional representations. Despite reported success, the dependence on matrices like an adjacency matrix creates an issue of scalability as matrices of large networks would take too much memory for a regular computer to operate well. Furthermore, these methods treat all edge types the same, but a possible extension is to incorporate node and edge types as sources of input.

Translational Distance Models

Translational distance models treat edges in a knowledge graph as linear transformations. For example, one such algorithm, TransE [137], treats every node-edge pair as a triplet with head nodes represented as \mathbf{h} , edges represented as \mathbf{r} , and tail nodes represented as \mathbf{t} . These representations are combined into an equation that mimics the iconic word vectors translations ($\mathbf{king} - \mathbf{man} + \mathbf{woman} \approx \mathbf{queen}$) from the word2vec model [97]. The described equation is shown as follows: $\mathbf{h} + \mathbf{r} \approx \mathbf{t}$. Starting at the head node (\mathbf{h}), add the edge vector (\mathbf{r}) and the result should be the tail node (\mathbf{t}). TransE optimizes vectors for \mathbf{h} , \mathbf{r} , \mathbf{t} , while guaranteeing the global

equation ($\mathbf{h} + \mathbf{r} \approx \mathbf{t}$) is satisfied [137]. A caveat to the TransE approach is that it the training steps force relationships to have a one to one mapping, which may not be appropriate for all relationship types.

Wang et al. attempted to resolve the one to one mapping issue by developing the TransH model [138]. TransH treats relations as hyperplanes rather than a regular vector and projects the head (\mathbf{h}) and tail (\mathbf{t}) nodes onto this hyperplane. Following this projection, a distance vector (\mathbf{d}_r) is calculated between the projected head and tail nodes. Finally, each vector is optimized while preserving the global equation: $\mathbf{h} + \mathbf{d}_r \approx \mathbf{t}$ [138]. Other effots have built off of the TransE and TransH models [139,140]. In the future, it may be beneficial for these models is to incorporate other types of information such as edge confidence scores, textual information, or edge type information when optimizing these vectors.

Deep Learning

Deep learning is a paradigm that uses multiple non-linear transformations to map high dimensional data into a low dimensional space. Many techniques that use deep learning on knowledge graphs are based on word2vec [96,97], a set of approaches that are widely used for natural language processing. The goal of word2vec is to project words onto a low dimensional space that preserves their semantic meaning. Strategies for training word2vec models use one of two neural network architectures: skipgram and continuous bag of words (CBOW). Both models are feed-forward neural networks, but CBOW models are trained to predict a word given its context while skip-gram models are trained to predict the context given a word [96,97]. Once training has finished, words are now associated with dense vectors that downstream models, such as feed forward networks or recurrent networks, can use for input.

Deepwalk is an early method that represents knowledge graphs in a low dimensional space [141]. The first step of this method is to perform a random walk along a knowledge graph. During the random walk, every generated sequence of nodes is recorded and treated as a sentence in word2vec [96,97]. After every node has been processed, a skip-gram model is trained to predict the context of each node thereby constructing a low dimensional representation of a knowledge graph [141]. A limitation for deepwalk is that the random walk cannot be controlled, so every node has an equal chance to be reached. Grover and Leskovec demonstrated that this limitation can hurt performance when classifying edges between nodes and developed node2vec as a result [142]. Node2vec operates the in the same fashion as deepwalk; however, this algorithm specifies a parameter that lets the random walk be biased when traversing nodes [142]. A caveat to both deepwalk and node2vec is that they ignore information such as edge type and node type. Various approaches have evolved to fix this limitation by incorporating node, edge and even path types when representing knowledge graphs in a low dimensional space [143,144,145,146]. An emerging area of work is to develop approaches that capture both the local and global structure of a graph when constructing this low dimensional space.

Instead of using the word2vec framework, some deep learning approaches use an adjacency matrix as input [96,97]. These approaches use neural networks called autoencoders to generate this low dimensional space [147,148,149]. Autoencoders map input such as an adjacency matrices into a low dimensional space and then determines how to construct this space via reconstructing the same input [150,151]. This generated space represents the nodes and their connectivity structure within a knowledge graph [147,148,149]. Despite the high potential of this approach, this method relies on an adjacency matrix for input which can run into scalability issues as a knowledge graph asymptotically increases in size [152]. Plus, Khosla et al. discovered that approaches akin to node2vec outperformed algorithms using autoencoders when undergoing link prediction and node classification [152]. Overall, the performance of deep learning techniques largely depends upon the structure of nodes and edges within a knowledge graph [152]. Future work should include hybrid models that use both

node2vec and autoencoders to construct complementary low dimensional representations of knowledge graphs.

Unifying Applications

Knowledge graphs have been applied to many biomedical challenges ranging from identifying proteins' functions [153] to prioritizing cancer genes [154] to recommending safer drugs to patients [155,156] (Figure 5). In this section we review how knowledge graphs are applied in biomedical settings and put particular emphasis on an emerging set of techniques that represent knowledge graphs in a low dimensional space.

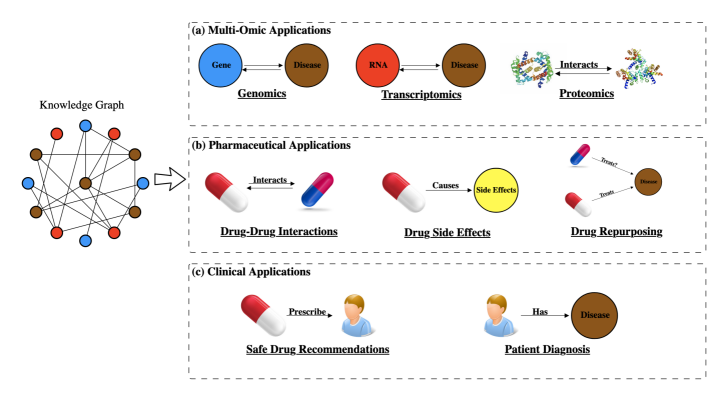


Figure 5: Overview of biomedical applications that make use of knowledge graphs. Categories consist of: (a) Multi-Omic applications, (b) Pharmaceutical Applications and (c) Clinical Applications.

Multi-Omic Applications

Multi-omic applications employ knowledge graphs to study the genome, how genes are expressed in the transcriptome, and how the products of those transcripts interact in the proteome. These graphs are used to establish connections between -omic entities as well as diseases. Tasks in this context include gene-symptom prioritization [157], protein-protein interaction prediction [158,159], and detecting miRNA-disease associations [120]. We focus specifically on multi-omic applications that represent knowledge graphs in a low dimensional space to make connections.

Recommendation systems make use of knowledge graphs to establish links between RNA with disease and proteins with other proteins. Shen et al. used an algorithm called collaborative filtering to establish an association between miRNA and diseases [120]. The authors constructed an miRNA-Disease network using the Human MicroRNA Disease database (HMDD) [160] and generated an adjacency matrix with the rows representing miRNA and the columns representing diseases. This matrix was decomposed into small rectangular matrices using SVD, then these small matrices were used to calculate similarity scores between miRNAs and diseases. High scores implied a high likelihood that a given miRNA had an association with a given disease [120]. Other approaches have built off of Shen et al.'s work by incorporating novel ways to perform matrix factorization [161,162,163] or by integrating machine learning models in conjunction with matrix factorization

[164]. These approaches achieved high area under the receiver operating curve (AUROC), but new discoveries have been hard to validate as experiments in this space are costly and time consuming at best [120]. Apart from miRNA, collaborative filtering has been used to predict protein-protein interactions [158,159,165]. Although extensive validation of newly generated candidates may be impractical, it would be helpful to see future efforts in this space include a blinded literature search for prioritized and randomly selected candidates as part of the standard evaluation pipeline.

Applications of deep learning techniques have mainly used the node2vec model [142] or variants of it. Yang et al. used node2vec to create a recommendation system to infer associations between genes and disease symptoms [157]. The authors constructed a gene-disease symptom knowledge graph by combining two bipartite graphs: genes with diseases and diseases with disease symptoms. The generated graph was embedded via node2vec and similarity scores were calculated for every gene-symptom pair in the graph. High scores implied a high likelihood of an association [157]. This approach outperformed methods that didn't use a knowledge graph; however, validation was difficult as it involved manual curation of the literature [157]. Similar approaches used variants of node2vec to predict gene-disease associations [8,166,167] analyze RNA-seq data [168] and infer novel protein information [153,169,170,171].

Knowledge graphs benefited the multi-omics field as a resource for generating novel discoveries. Most approaches to date use matrix factorization and node2vec to project knowledge graph into a low dimensional space, while translational models may be an untapped resource that could aid future efforts. Another area of exploration could be incorporating multiple sources of information such as compounds, anatomic locations or genetic pathways to improve the specificity of findings (i.e., to predict that a protein-protein interaction happens in a specific cell type or tissue).

Pharmaceutical Applications

There are a multitude of examples where knowledge graphs have been applied to identify new properties of drugs. Tasks in this field involve predicting drugs interacting with other drugs [172], identifying molecular targets a drug might interact with [173] and identifying new disease treatments for previously established drugs [174]. In this section we concentrate on applications that apply these graphs to discover new properties of drugs and focus on approaches that use these graphs in a low-dimensional space.

Similar to multi-omic applications recommendation systems have utilized knowledge graphs to infer novel links between drugs and diseases. Dai et al. used collaborative filtering to infer drug-disease associations [173]. The authors constructed a drug-disease network by integrating two bipartite networks: a drug-gene interaction network and a disease-gene interaction network. They integrated both networks under the assumption that drugs associated with a disease interact with the same gene of interest. Following construction, the authors generated an adjacency matrix where rows represent drugs and columns represent diseases. This matrix was decomposed into two small rectangular matrices and these matrices were used to calculate similarity scores between all drugs and all diseases. High values implied a high chance of an association [173]. Related approaches used this technique to infer drug-target interactions [175,176,177] and drug-disease treatments [178,179,180,181,182]. In spite of reported success, these approaches are limited to the drugs and diseases contained in the graph. Combining these approaches with representations of chemical structures might make it possible to one day make predictions about novel compounds.

Applications that use deep learning techniques have used node2vec [183,184] and autoencoders [185,186] approaches to represent knowledge graphs in a low dimensional space. Zong et al. used a node2vec-like model to predict drug-target associations [183]. The authors constructed a disease-target-disease network using drug centered databases: Drugbank [187] and Diseasome [188]. Next, the authors applied a random walk to the graph and trained a skip-gram model to generate a low

dimensional representation of the graph. Lastly, the authors construct a similarity metric that used this space to rank how similar drugs are to their targets [183]. A limitation to this approach is that their graph is missing information such as pharmacological class or drug chemical structure that could improve prediction performance. Overall, deep learning provides a robust set of techniques that have been shown to outperform most linear approaches in this context [189,190].

Applications that discover new properties of drugs have benefited from using knowledge graphs as a resource. Most methods to date use matrix factorization and deep learning techniques to produce a low-dimensional representation. Due to the success of deep learning [189,190] much of the field's focus has shifted to these techniques; however, a possible extension is to use an ensemble of deep learning techniques and linear methods to improve performance. Plus, another area of exploration is to incorporate information such as pharmaceutical classes for drugs or chemical structure to improve detection power.

Clinical applications

Clinical applications that use knowledge graphs are in early stages of development, but the long-term goal is to use analyses of these graphs to aid patient care. Typically, graphs for these applications are constructed from electronic health records (EHR) and nodes represent patients, drugs and diseases while edges represent relationships such as a patient being prescribed a treatment or a patient being diagnosed with a disease [18,191,192,193]. Tasks within this field range from improving patient diagnoses [194,195] to recommending safer drugs for patients [155,195] and we briefly discuss efforts that use knowledge graphs to accomplish such tasks.

Early work in this field applied translational models (Figure $\underline{4}$ (b)) to knowledge graphs with the goal of recommend safe drugs. Wang et al. used a variant of the TransH [138] model to create such a system for patients [155]. They constructed a disease-patient-drug network by integrating a patient-disease bipartite network with a patient-drug bipartite network. Together with all translational models every node in the graph was embedded while satisfying the following equation: $\mathbf{h} - \mathbf{r} \approx \mathbf{t}$. Following the embedding step, the authors formulated their own similarity metric that selected drug combinations with a low number of interactions [155]. Researchers in [140] applied a similar variant of the TransH model to a medical knowledge graph and evaluated their model for link prediction rather than patient recommendation.

In contrast with most applications where node2vec and autoencoder models have become established, deep learning methods in this field have focused on using graph attention models [196]. These models mimic machine translation models [197] and aim to simultaneously represent knowledge graphs in a low dimensional space and perform the task at hand. Choi et al. used a graph attention model to predict patient diagnoses [119]. The authors constructed a directed graph using medical concepts from patient EHR data. This directed graph was fed into a graph attention network and then used to predict a patient's likelihood of heart failure [119]. Other approaches have used graph attention models to perform clinical tasks such as drug safety recommendations [156] and patient diagnoses [198].

Knowledge graphs have shown promising results when used for clinical applications; however, there is still room for improvement. Most approaches have run into the common problem of missing data within EHR [119,155,156]. Future directions for the field consist of designing algorithms that can fill in this missing data gap or construct models that can take missing data into account.

Conclusion

Knowledge graphs are becoming widely used in biomedicine, and we expect their use to continue to grow. At the moment, most are constructed from databases derived from manual curation or from

co-occurrences in text. However, we expect that machine learning approaches will play a key role in quickly bringing new findings into these graphs. Representing these knowledge graphs in a low dimensional space that captures a graph's local and global structure can enable many downstream machine learning analyses, and methods to capture this structure are an active area of research.

As with any field, rigorous evaluation that can identify key factors that drive success is critical to moving the field forward. In regard to knowledge graphs, evaluation remains hard. Experiments in this context require a significant amount of time and consequently resources [120,157]. Moving from open ended and uncontrolled evaluations that consist of describing findings that are consistent with the literature to blinded evaluations of the literature that corroborate predictions and non-predictions would be a valuable first step. There are also well-documented biases related to node degree and degree distribution that must be considered for accurate evaluation [199]. Furthermore, the diversity of applications hinders the development of a standardized set of expected evaluations.

We anticipate that a fruitful avenue of research will be techniques that can produce low dimensional representations of knowledge graphs that distinguish between multple node and edge types. There are also many different sources of bias that lead to spurious edges or incompleteness, and modeling these may support better representations. It is a promising time for research into the construction and application of knowledge graphs. The peer reviewed literature is growing at an increasing rate and maintaining a complete understanding is becoming increasingly challenging for scientists. One path that scientists can take to maintain complete awareness is to become hyper-focused on specific areas of knowledge graph literature. If advances in how these graphs are constructed, represented, and applied can enable the linking of fields, we may be able to savor the benefits of this detailed knowledge without losing the broader contextual links.

References

1. Node Classification in Social Networks

Smriti Bhagat, Graham Cormode, S. Muthukrishnan *Social Network Data Analytics* (2011) https://doi.org/fjj48w

DOI: <u>10.1007/978-1-4419-8462-3_5</u>

2. Network Embedding Based Recommendation Method in Social Networks

Yufei Wen, Lei Guo, Zhumin Chen, Jun Ma

Companion of the The Web Conference 2018 on The Web Conference 2018 - WWW '18 (2018)

https://doi.org/gf6rtt

DOI: 10.1145/3184558.3186904

3. Open Question Answering with Weakly Supervised Embedding Models

Antoine Bordes, Jason Weston, Nicolas Usunier *arXiv* (2014-04-16) https://arxiv.org/abs/1404.4326v1

4. Neural Network-based Question Answering over Knowledge Graphs on Word and Character Level

Denis Lukovnikov, Asja Fischer, Jens Lehmann, Sören Auer *Proceedings of the 26th International Conference on World Wide Web* (2017-04-03) https://doi.org/gfv8hp

DOI: 10.1145/3038912.3052675

5. Towards integrative gene prioritization in Alzheimer's disease.

Jang H Lee, Graciela H Gonzalez

Pacific Symposium on Biocomputing. Pacific Symposium on Biocomputing (2011)

https://www.ncbi.nlm.nih.gov/pubmed/21121028

DOI: 10.1142/9789814335058 0002 · PMID: 21121028

6. PhenoGeneRanker: A Tool for Gene Prioritization Using Complete Multiplex Heterogeneous Networks

Cagatay Dursun, Naoki Shimoyama, Mary Shimoyama, Michael Schläppi, Serdar Bozdag *Cold Spring Harbor Laboratory* (2019-05-27) https://doi.org/gf6rtr

DOI: <u>10.1101/651000</u>

7. Biological Random Walks: Integrating heterogeneous data in disease gene prioritization

Michele Gentili, Leonardo Martini, Manuela Petti, Lorenzo Farina, Luca Becchetti 2019 IEEE Conference on Computational Intelligence in Bioinformatics and Computational Biology (CIBCB) (2019-07) https://doi.org/gf6rts

DOI: <u>10.1109/cibcb.2019.8791472</u>

8. Semantic Disease Gene Embeddings (SmuDGE): phenotype-based disease gene prioritization without phenotypes

Mona Alshahrani, Robert Hoehndorf

Bioinformatics (2018-09-01) https://doi.org/gd9k8n

DOI: <u>10.1093/bioinformatics/bty559</u> · PMID: <u>30423077</u> · PMCID: <u>PMC6129260</u>

9. Systematic integration of biomedical knowledge prioritizes drugs for repurposing

Daniel Scott Himmelstein, Antoine Lizee, Christine Hessler, Leo Brueggeman, Sabrina L Chen, Dexter Hadley, Ari Green, Pouya Khankhanian, Sergio E Baranzini

eLife (2017-09-22) https://doi.org/cdfk

DOI: 10.7554/elife.26726 · PMID: 28936969 · PMCID: PMC5640425

10. Assessing Drug Target Association Using Semantic Linked Data

Bin Chen, Ying Ding, David J. Wild

PLoS Computational Biology (2012-07-05) https://doi.org/rn6

DOI: 10.1371/journal.pcbi.1002574 · PMID: 22859915 · PMCID: PMC3390390

11. Towards a definition of knowledge graphs

Lisa Ehrlinger, Wolfram Wöß SEMANTiCS (2016)

12. DISEASES: Text mining and data integration of disease-gene associations

Sune Pletscher-Frankild, Albert Pallejà, Kalliopi Tsafou, Janos X. Binder, Lars Juhl Jensen *Methods* (2015-03) https://doi.org/f3mn6s

DOI: 10.1016/j.ymeth.2014.11.020 · PMID: 25484339

13. DrugBank 5.0: a major update to the DrugBank database for 2018

David S Wishart, Yannick D Feunang, An C Guo, Elvis J Lo, Ana Marcu, Jason R Grant, Tanvir Sajed, Daniel Johnson, Carin Li, Zinat Sayeeda, ... Michael Wilson

Nucleic Acids Research (2017-11-08) https://doi.org/gcwtzk

DOI: 10.1093/nar/gkx1037 · PMID: 29126136 · PMCID: PMC5753335

14. A network integration approach for drug-target interaction prediction and computational drug repositioning from heterogeneous information

Yunan Luo, Xinbin Zhao, Jingtian Zhou, Jinglin Yang, Yanqing Zhang, Wenhua Kuang, Jian Peng, Ligong Chen, Jianyang Zeng

Nature Communications (2017-09-18) https://doi.org/gbxwrc

DOI: <u>10.1038/s41467-017-00680-8</u> · PMID: <u>28924171</u> · PMCID: <u>PMC5603535</u>

15. Inferring new indications for approved drugs via random walk on drug-disease heterogenous networks

Hui Liu, Yinglong Song, Jihong Guan, Libo Luo, Ziheng Zhuang

BMC Bioinformatics (2016-12) https://doi.org/gf6v27

DOI: 10.1186/s12859-016-1336-7 · PMID: 28155639 · PMCID: PMC5259862

16. Finding disease similarity based on implicit semantic similarity

Sachin Mathur, Deendayal Dinakarpandian

Journal of Biomedical Informatics (2012-04) https://doi.org/b7b3tw

DOI: <u>10.1016/j.jbi.2011.11.017</u> · PMID: <u>22166490</u>

17. Bio2RDF: Towards a mashup to build bioinformatics knowledge systems

François Belleau, Marc-Alexandre Nolin, Nicole Tourigny, Philippe Rigault, Jean Morissette *Journal of Biomedical Informatics* (2008-10) https://doi.org/frqkq5

DOI: 10.1016/j.jbi.2008.03.004 · PMID: 18472304

18. KnowLife: a versatile approach for constructing a large knowledge graph for biomedical sciences

Patrick Ernst, Amy Siu, Gerhard Weikum

BMC Bioinformatics (2015-05-14) https://doi.org/gb8w8d

DOI: 10.1186/s12859-015-0549-5 · PMID: 25971816 · PMCID: PMC4448285

19. Constructing biomedical domain-specific knowledge graph with minimum supervision

Jianbo Yuan, Zhiwei Jin, Han Guo, Hongxia Jin, Xianchao Zhang, Tristram Smith, Jiebo Luo *Knowledge and Information Systems* (2019-03-23) https://doi.org/gf6v26

DOI: 10.1007/s10115-019-01351-4

20. Feature assisted stacked attentive shortest dependency path based Bi-LSTM model for protein–protein interaction

Shweta Yadav, Asif Ekbal, Sriparna Saha, Ankit Kumar, Pushpak Bhattacharyya *Knowledge-Based Systems* (2019-02) https://doi.org/gf4788

DOI: 10.1016/j.knosys.2018.11.020

21. Biological Databases- Integration of Life Science Data

Nishant Toomula, Arun Kumar, Sathish Kumar D, Vijaya Shanti Bheemidi Journal of Computer Science & Systems Biology (2012) https://doi.org/gf8qcb

DOI: <u>10.4172/jcsb.1000081</u>

22. COSMIC: somatic cancer genetics at high-resolution

Simon A. Forbes, David Beare, Harry Boutselakis, Sally Bamford, Nidhi Bindal, John Tate, Charlotte G. Cole, Sari Ward, Elisabeth Dawson, Laura Ponting, ... Peter J. Campbell

Nucleic Acids Research (2016-11-28) https://doi.org/f9v865

DOI: <u>10.1093/nar/gkw1121</u> · PMID: <u>27899578</u> · PMCID: <u>PMC5210583</u>

23. COSMIC: the Catalogue Of Somatic Mutations In Cancer

John G Tate, Sally Bamford, Harry C Jubb, Zbyslaw Sondka, David M Beare, Nidhi Bindal, Harry Boutselakis, Charlotte G Cole, Celestino Creatore, Elisabeth Dawson, ... Simon A Forbes *Nucleic Acids Research* (2018-10-29) https://doi.org/gf9hxg

DOI: 10.1093/nar/gky1015 · PMID: 30371878 · PMCID: PMC6323903

24. Recurated protein interaction datasets

Lukasz Salwinski, Luana Licata, Andrew Winter, David Thorneycroft, Jyoti Khadake, Arnaud Ceol, Andrew Chatr Aryamontri, Rose Oughtred, Michael Livstone, Lorrie Boucher, ... Henning Hermjakob *Nature Methods* (2009-12) https://doi.org/fgvkmf

DOI: 10.1038/nmeth1209-860 · PMID: 19935838

25. Literature-curated protein interaction datasets

Michael E Cusick, Haiyuan Yu, Alex Smolyar, Kavitha Venkatesan, Anne-Ruxandra Carvunis, Nicolas Simonis, Jean-François Rual, Heather Borick, Pascal Braun, Matija Dreze, ... Marc Vidal *Nature Methods* (2008-12-30) https://doi.org/d4j62p

DOI: <u>10.1038/nmeth.1284</u> · PMID: <u>19116613</u> · PMCID: <u>PMC2683745</u>

26. Curation accuracy of model organism databases

I. M. Keseler, M. Skrzypek, D. Weerasinghe, A. Y. Chen, C. Fulcher, G.-W. Li, K. C. Lemmer, K. M. Mladinich, E. D. Chow, G. Sherlock, P. D. Karp

Database (2014-06-12) https://doi.org/gf63jz

DOI: 10.1093/database/bau058 · PMID: 24923819 · PMCID: PMC4207230

27. OMIM.org: Online Mendelian Inheritance in Man (OMIM®), an online catalog of human genes and genetic disorders

Joanna S. Amberger, Carol A. Bocchini, François Schiettecatte, Alan F. Scott, Ada Hamosh *Nucleic Acids Research* (2014-11-26) https://doi.org/gf8qb6

DOI: 10.1093/nar/gku1205 · PMID: 25428349 · PMCID: PMC4383985

28. Textpresso Central: a customizable platform for searching, text mining, viewing, and curating biomedical literature

H.-M. Müller, K. M. Van Auken, Y. Li, P. W. Sternberg

BMC Bioinformatics (2018-03-09) https://doi.org/gf7rbz

DOI: <u>10.1186/s12859-018-2103-8</u> · PMID: <u>29523070</u> · PMCID: <u>PMC5845379</u>

29. Text mining and expert curation to develop a database on psychiatric diseases and their genes

Alba Gutiérrez-Sacristán, Àlex Bravo, Marta Portero-Tresserra, Olga Valverde, Antonio Armario, M. C. Blanco-Gandía, Adriana Farré, Lierni Fernández-Ibarrondo, Francina Fonseca, Jesús Giraldo, ... Laura I. Furlong

Database (2017-01-01) https://doi.org/gf8qb5

DOI: <u>10.1093/database/bax043</u> · PMID: <u>29220439</u> · PMCID: <u>PMC5502359</u>

30. Manual curation is not sufficient for annotation of genomic databases

William A. Baumgartner Jr, K. Bretonnel Cohen, Lynne M. Fox, George Acquaah-Mensah, Lawrence Hunter

Bioinformatics (2007-07-01) https://doi.org/dtck86

DOI: <u>10.1093/bioinformatics/btm229</u> · PMID: <u>17646325</u> · PMCID: <u>PMC2516305</u>

31. The rate of growth in scientific publication and the decline in coverage provided by Science Citation Index

Peder Olesen Larsen, Markus von Ins

Scientometrics (2010-03-10) https://doi.org/c4hb8r

DOI: <u>10.1007/s11192-010-0202-z</u> · PMID: <u>20700371</u> · PMCID: <u>PMC2909426</u>

32. Semi-automatic semantic annotation of PubMed queries: A study on quality, efficiency, satisfaction

Aurélie Névéol, Rezarta Islamaj Doğan, Zhiyong Lu

Journal of Biomedical Informatics (2011-04) https://doi.org/bq34sj

DOI: <u>10.1016/j.jbi.2010.11.001</u> · PMID: <u>21094696</u> · PMCID: <u>PMC3063330</u>

33. Assisting manual literature curation for protein-protein interactions using BioQRator

D. Kwon, S. Kim, S.-Y. Shin, A. Chatr-aryamontri, W. J. Wilbur

Database (2014-07-22) https://doi.org/gf7hm3

DOI: 10.1093/database/bau067 · PMID: 25052701 · PMCID: PMC4105708

34. Argo: an integrative, interactive, text mining-based workbench supporting curation

R. Rak, A. Rowley, W. Black, S. Ananiadou

Database (2012-03-20) https://doi.org/h5d

DOI: <u>10.1093/database/bas010</u> · PMID: <u>22434844</u> · PMCID: <u>PMC3308166</u>

35. **CurEx**

Michael Loster, Felix Naumann, Jan Ehmueller, Benjamin Feldmann

Proceedings of the 27th ACM International Conference on Information and Knowledge Management (2018-10-17) https://doi.org/gf8qb8

DOI: 10.1145/3269206.3269229

36. Re-curation and rational enrichment of knowledge graphs in Biological Expression Language

Charles Tapley Hoyt, Daniel Domingo-Fernández, Rana Aldisi, Lingling Xu, Kristian Kolpeja, Sandra Spalek, Esther Wollert, John Bachman, Benjamin M Gyori, Patrick Greene, Martin Hofmann-Apitius

Database (2019-01-01) https://doi.org/gf7hm4

DOI: 10.1093/database/baz068 · PMID: 31225582 · PMCID: PMC6587072

37. LocText: relation extraction of protein localizations to assist database curation

Juan Miguel Cejuela, Shrikant Vinchurkar, Tatyana Goldberg, Madhukar Sollepura Prabhu Shankar, Ashish Baghudana, Aleksandar Bojchevski, Carsten Uhlig, André Ofner, Pandu Raharja-Liu, Lars Juhl Jensen, Burkhard Rost

BMC Bioinformatics (2018-01-17) https://doi.org/gf8qb9

DOI: <u>10.1186/s12859-018-2021-9</u> · PMID: <u>29343218</u> · PMCID: <u>PMC5773052</u>

38. Evaluating the impact of pre-annotation on annotation speed and potential bias: natural language processing gold standard development for clinical named entity recognition in clinical trial announcements

Todd Lingren, Louise Deleger, Katalin Molnar, Haijun Zhai, Jareen Meinzen-Derr, Megan Kaiser, Laura Stoutenborough, Qi Li, Imre Solti

Journal of the American Medical Informatics Association (2014-05) https://doi.org/f5zggh

DOI: 10.1136/amiajnl-2013-001837 · PMID: 24001514 · PMCID: PMC3994857

39. iSimp in BioC standard format: enhancing the interoperability of a sentence simplification system

Y. Peng, C. O. Tudor, M. Torii, C. H. Wu, K. Vijay-Shanker

Database (2014-05-21) https://doi.org/gf9hxf

DOI: 10.1093/database/bau038 · PMID: 24850848 · PMCID: PMC4028706

40. BioSimplify: an open source sentence simplification engine to improve recall in automatic biomedical information extraction.

Siddhartha Jonnalagadda, Graciela Gonzalez

AMIA ... Annual Symposium proceedings. AMIA Symposium (2010-11-13)

https://www.ncbi.nlm.nih.gov/pubmed/21346999

PMID: 21346999 · PMCID: PMC3041388

41. The EU-ADR corpus: Annotated drugs, diseases, targets, and their relationships

Erik M. van Mulligen, Annie Fourrier-Reglat, David Gurwitz, Mariam Molokhia, Ainhoa Nieto, Gianluca Trifiro, Jan A. Kors, Laura I. Furlong

Journal of Biomedical Informatics (2012-10) https://doi.org/f36vn6

DOI: 10.1016/j.jbi.2012.04.004 · PMID: 22554700

42. Comparative experiments on learning information extractors for proteins and their interactions

Razvan Bunescu, Ruifang Ge, Rohit J. Kate, Edward M. Marcotte, Raymond J. Mooney, Arun K. Ramani, Yuk Wah Wong

Artificial Intelligence in Medicine (2005-02) https://doi.org/dhztpn

DOI: <u>10.1016/j.artmed.2004.07.016</u> · PMID: <u>15811782</u>

43. A Unified Active Learning Framework for Biomedical Relation Extraction

Hong-Tao Zhang, Min-Lie Huang, Xiao-Yan Zhu

Journal of Computer Science and Technology (2012-11) https://doi.org/gf8qb4

DOI: <u>10.1007/s11390-012-1306-0</u>

44. The BioGRID interaction database: 2013 update

Andrew Chatr-aryamontri, Bobby-Joe Breitkreutz, Sven Heinicke, Lorrie Boucher, Andrew Winter, Chris Stark, Julie Nixon, Lindsay Ramage, Nadine Kolas, Lara O'Donnell, ... Mike Tyers

Nucleic Acids Research (2012-11-30) https://doi.org/f4jmz4

DOI: 10.1093/nar/gks1158 · PMID: 23203989 · PMCID: PMC3531226

45. The Comparative Toxicogenomics Database: update 2019

Allan Peter Davis, Cynthia J Grondin, Robin J Johnson, Daniela Sciaky, Roy McMorran, Jolene Wiegers, Thomas C Wiegers, Carolyn J Mattingly

Nucleic Acids Research (2018-09-24) https://doi.org/gf8qb7

DOI: <u>10.1093/nar/gky868</u> · PMID: <u>30247620</u> · PMCID: <u>PMC6323936</u>

46. CARD 2017: expansion and model-centric curation of the comprehensive antibiotic resistance database

Baofeng Jia, Amogelang R. Raphenya, Brian Alcock, Nicholas Waglechner, Peiyao Guo, Kara K. Tsang, Briony A. Lago, Biren M. Dave, Sheldon Pereira, Arjun N. Sharma, ... Andrew G. McArthur *Nucleic Acids Research* (2016-10-26) https://doi.org/f9wbjs

DOI: <u>10.1093/nar/gkw1004</u> · PMID: <u>27789705</u> · PMCID: <u>PMC5210516</u>

47. Entrez Gene: gene-centered information at NCBI

D. Maglott, J. Ostell, K. D. Pruitt, T. Tatusova

Nucleic Acids Research (2010-11-28) https://doi.org/fsjcqz

DOI: 10.1093/nar/gkq1237 · PMID: 21115458 · PMCID: PMC3013746

48. OMIM.org: leveraging knowledge across phenotype-gene relationships.

Joanna S Amberger, Carol A Bocchini, Alan F Scott, Ada Hamosh

Nucleic acids research (2019-01-08) https://www.ncbi.nlm.nih.gov/pubmed/30445645

DOI: 10.1093/nar/gky1151 · PMID: 30445645 · PMCID: PMC6323937

49. Pharmacogenomics Knowledge for Personalized Medicine

M Whirl-Carrillo, EM McDonagh, JM Hebert, L Gong, K Sangkuhl, CF Thorn, RB Altman, TE Klein *Clinical Pharmacology & Therapeutics* (2012-10) https://doi.org/gdnfzr
DOI: 10.1038/clpt.2012.96 · PMID: 22992668 · PMCID: PMC3660037

50. UniProt: a worldwide hub of protein knowledge Nucleic Acids Research (2018-11-05)

https://doi.org/gfwqck

DOI: <u>10.1093/nar/gky1049</u> · PMID: <u>30395287</u> · PMCID: <u>PMC6323992</u>

51. LPTK: a linguistic pattern-aware dependency tree kernel approach for the BioCreative VI CHEMPROT task

Neha Warikoo, Yung-Chun Chang, Wen-Lian Hsu

Database (2018-01-01) https://doi.org/gfhjr6

DOI: <u>10.1093/database/bay108</u> · PMID: <u>30346607</u> · PMCID: <u>PMC6196310</u>

52. DTMiner: identification of potential disease targets through biomedical literature mining

Dong Xu, Meizhuo Zhang, Yanping Xie, Fan Wang, Ming Chen, Kenny Q. Zhu, Jia Wei *Bioinformatics* (2016-08-09) https://doi.org/f9nw36

DOI: 10.1093/bioinformatics/btw503 · PMID: 27506226 · PMCID: PMC5181534

53. Exploiting graph kernels for high performance biomedical relation extraction

Nagesh C. Panyam, Karin Verspoor, Trevor Cohn, Kotagiri Ramamohanarao *Journal of Biomedical Semantics* (2018-01-30) https://doi.org/gf49nn

DOI: <u>10.1186/s13326-017-0168-3</u> · PMID: <u>29382397</u> · PMCID: <u>PMC5791373</u>

54. iSimp in BioC standard format: enhancing the interoperability of a sentence simplification system.

Yifan Peng, Catalina O Tudor, Manabu Torii, Cathy H Wu, K Vijay-Shanker *Database: the journal of biological databases and curation* (2014-05-21)

https://www.ncbi.nlm.nih.gov/pubmed/24850848

DOI: 10.1093/database/bau038 · PMID: 24850848 · PMCID: PMC4028706

55. BELMiner: adapting a rule-based relation extraction system to extract biological expression language statements from bio-medical literature evidence sentences

K. E. Ravikumar, Majid Rastegar-Mojarad, Hongfang Liu

Database (2017-01-01) https://doi.org/gf7rbx

DOI: 10.1093/database/baw156 · PMID: 28365720 · PMCID: PMC5467463

56. A generalizable NLP framework for fast development of pattern-based biomedical relation extraction systems

Yifan Peng, Manabu Torii, Cathy H Wu, K Vijay-Shanker

BMC Bioinformatics (2014-08-23) https://doi.org/f6rndz

DOI: 10.1186/1471-2105-15-285 · PMID: 25149151 · PMCID: PMC4262219

57. Construction of phosphorylation interaction networks by text mining of full-length articles using the eFIP system

Catalina O. Tudor, Karen E. Ross, Gang Li, K. Vijay-Shanker, Cathy H. Wu, Cecilia N. Arighi

Database (2015-01-01) https://doi.org/gf8fpt

DOI: 10.1093/database/bav020 · PMID: 25833953 · PMCID: PMC4381107

58. miRTex: A Text Mining System for miRNA-Gene Relation Extraction

Gang Li, Karen E. Ross, Cecilia N. Arighi, Yifan Peng, Cathy H. Wu, K. Vijay-Shanker

PLOS Computational Biology (2015-09-25) https://doi.org/f75mwb

DOI: <u>10.1371/journal.pcbi.1004391</u> · PMID: <u>26407127</u> · PMCID: <u>PMC4583433</u>

59. LimTox: a web tool for applied text mining of adverse event and toxicity associations of compounds, drugs and genes

Andres Cañada, Salvador Capella-Gutierrez, Obdulia Rabal, Julen Oyarzabal, Alfonso Valencia, Martin Krallinger

Nucleic Acids Research (2017-05-22) https://doi.org/gf479h

DOI: <u>10.1093/nar/gkx462</u> · PMID: <u>28531339</u> · PMCID: <u>PMC5570141</u>

60. DiMeX: A Text Mining System for Mutation-Disease Association Extraction

A. S. M. Ashique Mahmood, Tsung-Jung Wu, Raja Mazumder, K. Vijay-Shanker

PLOS ONE (2016-04-13) https://doi.org/f8xktj

DOI: <u>10.1371/journal.pone.0152725</u> · PMID: <u>27073839</u> · PMCID: <u>PMC4830514</u>

61. Automated extraction of mutation data from the literature: application of MuteXt to G protein-coupled receptors and nuclear hormone receptors

F. Horn, A. L. Lau, F. E. Cohen

Bioinformatics (2004-01-22) https://doi.org/d7cjgj

DOI: 10.1093/bioinformatics/btg449 · PMID: 14990452

62. Large-scale extraction of accurate drug-disease treatment pairs from biomedical literature for drug repurposing

Rong Xu, QuanQiu Wang

BMC Bioinformatics (2013-06-06) https://doi.org/gb8v3k

DOI: <u>10.1186/1471-2105-14-181</u> · PMID: <u>23742147</u> · PMCID: <u>PMC3702428</u>

63. RLIMS-P 2.0: A Generalizable Rule-Based Information Extraction System for Literature Mining of Protein Phosphorylation Information

Manabu Torii, Cecilia N. Arighi, Gang Li, Qinghua Wang, Cathy H. Wu, K. Vijay-Shanker *IEEE/ACM Transactions on Computational Biology and Bioinformatics* (2015-01-01) https://doi.org/gf8fpv

DOI: <u>10.1109/tcbb.2014.2372765</u> · PMID: <u>26357075</u> · PMCID: <u>PMC4568560</u>

64. PKDE4J: Entity and relation extraction for public knowledge discovery

Min Song, Won Chul Kim, Dahee Lee, Go Eun Heo, Keun Young Kang *Journal of Biomedical Informatics* (2015-10) https://doi.org/f7v7jj DOI: 10.1016/j.jbi.2015.08.008 • PMID: 26277115

65. PhpSyntaxTree tool

A Eisenbach, M Eisenbach (2006)

66. Spacy 2: Natural language understanding with bloom embeddings, convolutional neural networks and incremental parsing

Matthew Honnibal, Ines Montani *To appear* (2017)

67. STRING v9.1: protein-protein interaction networks, with increased coverage and integration

Andrea Franceschini, Damian Szklarczyk, Sune Frankild, Michael Kuhn, Milan Simonovic, Alexander Roth, Jianyi Lin, Pablo Minguez, Peer Bork, Christian von Mering, Lars J. Jensen *Nucleic Acids Research* (2012-11-29) https://doi.org/gf5kcd

DOI: 10.1093/nar/gks1094 · PMID: 23203871 · PMCID: PMC3531103

68. A comprehensive and quantitative comparison of text-mining in 15 million full-text articles versus their corresponding abstracts

David Westergaard, Hans-Henrik Stærfeldt, Christian Tønsberg, Lars Juhl Jensen, Søren Brunak *PLOS Computational Biology* (2018-02-15) https://doi.org/gcx747

DOI: <u>10.1371/journal.pcbi.1005962</u> · PMID: <u>29447159</u> · PMCID: <u>PMC5831415</u>

69. STITCH 4: integration of protein-chemical interactions with user data

Michael Kuhn, Damian Szklarczyk, Sune Pletscher-Frankild, Thomas H. Blicher, Christian von Mering, Lars J. Jensen, Peer Bork

Nucleic Acids Research (2013-11-28) https://doi.org/f5shb4

DOI: 10.1093/nar/gkt1207 · PMID: 24293645 · PMCID: PMC3964996

70. A global network of biomedical relationships derived from text

Bethany Percha, Russ B Altman

Bioinformatics (2018-02-27) https://doi.org/gc3ndk

DOI: 10.1093/bioinformatics/bty114 · PMID: 29490008 · PMCID: PMC6061699

71. CoCoScore: context-aware co-occurrence scoring for text mining applications using distant supervision

Alexander Junge, Lars Juhl Jensen

Bioinformatics (2019-06-14) https://doi.org/gf4789

DOI: 10.1093/bioinformatics/btz490 · PMID: 31199464 · PMCID: PMC6956794

72. A new method for prioritizing drug repositioning candidates extracted by literature-based discovery

Majid Rastegar-Mojarad, Ravikumar Komandur Elayavilli, Dingcheng Li, Rashmi Prasad, Hongfang Liu

https://doi.org/gf479j

DOI: 10.1109/bibm.2015.7359766

73. Literature Mining for the Discovery of Hidden Connections between Drugs, Genes and Diseases

Raoul Frijters, Marianne van Vugt, Ruben Smeets, René van Schaik, Jacob de Vlieg, Wynand Alkema *PLoS Computational Biology* (2010-09-23) https://doi.org/bhrw7x

DOI: <u>10.1371/journal.pcbi.1000943</u> · PMID: <u>20885778</u> · PMCID: <u>PMC2944780</u>

74. STRING v10: protein-protein interaction networks, integrated over the tree of life

Damian Szklarczyk, Andrea Franceschini, Stefan Wyder, Kristoffer Forslund, Davide Heller, Jaime Huerta-Cepas, Milan Simonovic, Alexander Roth, Alberto Santos, Kalliopi P. Tsafou, ... Christian von Mering

Nucleic Acids Research (2014-10-28) https://doi.org/f64rfn

DOI: 10.1093/nar/gku1003 · PMID: 25352553 · PMCID: PMC4383874

75. Text Mining Genotype-Phenotype Relationships from Biomedical Literature for Database Curation and Precision Medicine

Ayush Singhal, Michael Simmons, Zhiyong Lu

PLOS Computational Biology (2016-11-30) https://doi.org/f9gz4b

DOI: 10.1371/journal.pcbi.1005017 · PMID: 27902695 · PMCID: PMC5130168

76. Overview of the biocreative vi chemical-protein interaction track

Martin Krallinger, Obdulia Rabal, Saber A Akhondi, others

Proceedings of the sixth biocreative challenge evaluation workshop (2017)

https://www.semanticscholar.org/paper/Overview-of-the-BioCreative-VI-chemical-protein-Krallinger-Rabal/eed781f498b563df5a9e8a241c67d63dd1d92ad5

77. BioCreative V CDR task corpus: a resource for chemical disease relation extraction

Jiao Li, Yueping Sun, Robin J. Johnson, Daniela Sciaky, Chih-Hsuan Wei, Robert Leaman, Allan Peter Davis, Carolyn J. Mattingly, Thomas C. Wiegers, Zhiyong Lu

Database (2016) https://doi.org/gf5hfw

DOI: <u>10.1093/database/baw068</u> · PMID: <u>27161011</u> · PMCID: <u>PMC4860626</u>

78. RelEx-Relation extraction using dependency parse trees

K. Fundel, R. Kuffner, R. Zimmer

Bioinformatics (2006-12-01) https://doi.org/cz7q4d

DOI: 10.1093/bioinformatics/btl616 · PMID: 17142812

79. CoMAGC: a corpus with multi-faceted annotations of gene-cancer relations

Hee-Jin Lee, Sang-Hyung Shim, Mi-Ryoung Song, Hyunju Lee, Jong C Park

BMC Bioinformatics (2013) https://doi.org/gb8v5s

DOI: 10.1186/1471-2105-14-323 · PMID: 24225062 · PMCID: PMC3833657

80. Text Mining for Protein Docking

Varsha D. Badal, Petras J. Kundrotas, Ilya A. Vakser

PLOS Computational Biology (2015-12-09) https://doi.org/gcvj3b

DOI: 10.1371/journal.pcbi.1004630 · PMID: 26650466 · PMCID: PMC4674139

81. Automatic extraction of gene-disease associations from literature using joint ensemble learning

Balu Bhasuran, Jeyakumar Natarajan

PLOS ONE (2018-07-26) https://doi.org/gdx63f

DOI: 10.1371/journal.pone.0200699 · PMID: 30048465 · PMCID: PMC6061985

82. Extraction of relations between genes and diseases from text and large-scale data analysis: implications for translational research

Àlex Bravo, Janet Piñero, Núria Queralt-Rosinach, Michael Rautschka, Laura I Furlong

BMC Bioinformatics (2015-02-21) https://doi.org/f7kn8s

DOI: <u>10.1186/s12859-015-0472-9</u> · PMID: <u>25886734</u> · PMCID: <u>PMC4466840</u>

83. Deep learning

lan Goodfellow, Yoshua Bengio, Aaron Courville *The MIT Press* (2016)

ISBN: 0262035618, 9780262035613

84. Deep learning

Yann LeCun, Yoshua Bengio, Geoffrey Hinton

Nature (2015-05) https://doi.org/bmgp

DOI: 10.1038/nature14539 · PMID: 26017442

85. Long Short-Term Memory

Sepp Hochreiter, Jürgen Schmidhuber

Neural Computation (1997-11) https://doi.org/bxd65w

DOI: <u>10.1162/neco.1997.9.8.1735</u> · PMID: <u>9377276</u>

86. Deep learning for pharmacovigilance: recurrent neural network architectures for labeling adverse drug reactions in Twitter posts

Anne Cocos, Alexander G Fiks, Aaron J Masino

Journal of the American Medical Informatics Association (2017-02-22) https://doi.org/gbp9nj

DOI: <u>10.1093/jamia/ocw180</u> · PMID: <u>28339747</u>

87. Semantic Relations in Compound Nouns: Perspectives from Inter-Annotator Agreement

Yadav Prabha, Jezek Elisabetta, Bouillon Pierrette, Callahan Tiffany J., Bada Michael, Hunter Lawrence E., Cohen K. Bretonnel

Studies in Health Technology and Informatics (2017) https://doi.org/ggmk8t

DOI: 10.3233/978-1-61499-830-3-644

88. Cross-Sentence N-ary Relation Extraction with Graph LSTMs

Nanyun Peng, Hoifung Poon, Chris Quirk, Kristina Toutanova, Wen-tau Yih *arXiv* (2017-08-12) https://arxiv.org/abs/1708.03743v1

89. Drug drug interaction extraction from biomedical literature using syntax convolutional neural network

Zhehuan Zhao, Zhihao Yang, Ling Luo, Hongfei Lin, Jian Wang

Bioinformatics (2016-07-27) https://doi.org/f9nsq7

DOI: <u>10.1093/bioinformatics/btw486</u> · PMID: <u>27466626</u> · PMCID: <u>PMC5181565</u>

90. N-ary Relation Extraction using Graph State LSTM

Linfeng Song, Yue Zhang, Zhiguo Wang, Daniel Gildea *arXiv* (2018-08-28) https://arxiv.org/abs/1808.09101v1

91. A neural joint model for entity and relation extraction from biomedical text

Fei Li, Meishan Zhang, Guohong Fu, Donghong Ji

BMC Bioinformatics (2017-03-31) https://doi.org/gcgnx2

DOI: <u>10.1186/s12859-017-1609-9</u> · PMID: <u>28359255</u> · PMCID: <u>PMC5374588</u>

92. The problem of learning long-term dependencies in recurrent networks

Y. Bengio, P. Frasconi, P. Simard

IEEE International Conference on Neural Networks https://doi.org/d7zs24

DOI: 10.1109/icnn.1993.298725

93. Fundamentals of Recurrent Neural Network (RNN) and Long Short-Term Memory (LSTM) network

Alex Sherstinsky

Physica D: Nonlinear Phenomena (2020-03) https://doi.org/ggmzpd

DOI: 10.1016/j.physd.2019.132306

94. On the difficulty of training Recurrent Neural Networks

Razvan Pascanu, Tomas Mikolov, Yoshua Bengio *arXiv* (2012-11-21) https://arxiv.org/abs/1211.5063v2

95. Revisiting Unreasonable Effectiveness of Data in Deep Learning Era

Chen Sun, Abhinav Shrivastava, Saurabh Singh, Abhinav Gupta *arXiv* (2017-07-10) https://arxiv.org/abs/1707.02968v2

96. Efficient Estimation of Word Representations in Vector Space

Tomas Mikolov, Kai Chen, Greg Corrado, Jeffrey Dean *arXiv* (2013-01-16) https://arxiv.org/abs/1301.3781v3

97. Distributed Representations of Words and Phrases and their Compositionality

Tomas Mikolov, Ilya Sutskever, Kai Chen, Greg Corrado, Jeffrey Dean *arXiv* (2013-10-16) https://arxiv.org/abs/1310.4546v1

98. Deep learning for extracting protein-protein interactions from biomedical literature

Yifan Peng, Zhiyong Lu

arXiv(2017-06-05) https://arxiv.org/abs/1706.01556v2

99. Knowledge-guided convolutional networks for chemical-disease relation extraction

Huiwei Zhou, Chengkun Lang, Zhuang Liu, Shixian Ning, Yingyu Lin, Lei Du

BMC Bioinformatics (2019-05-21) https://doi.org/gf45zn

DOI: <u>10.1186/s12859-019-2873-7</u> · PMID: <u>31113357</u> · PMCID: <u>PMC6528333</u>

100. Extraction of protein-protein interactions (PPIs) from the literature by deep convolutional neural networks with various feature embeddings

Sung-Pil Choi

Journal of Information Science (2016-11-01) https://doi.org/gcv8bn

DOI: 10.1177/0165551516673485

101. Extracting chemical-protein relations with ensembles of SVM and deep learning models

Yifan Peng, Anthony Rios, Ramakanth Kavuluru, Zhiyong Lu

Database (2018-01-01) https://doi.org/gf479f

DOI: <u>10.1093/database/bay073</u> · PMID: <u>30020437</u> · PMCID: <u>PMC6051439</u>

102. Expanding a Database-derived Biomedical Knowledge Graph via Multi-relation Extraction from Biomedical Abstracts

David N. Nicholson, Daniel S. Himmelstein, Casey S. Greene

Cold Spring Harbor Laboratory (2019-08-08) https://doi.org/gf6qxh

DOI: <u>10.1101/730085</u>

103. Distant supervision for relation extraction without labeled data

Mike Mintz, Steven Bills, Rion Snow, Dan Jurafsky

Proceedings of the Joint Conference of the 47th Annual Meeting of the ACL and the 4th International Joint Conference on Natural Language Processing of the AFNLP: Volume 2 - ACL-IJCNLP '09 (2009)

https://doi.org/fg9q43

DOI: <u>10.3115/1690219.1690287</u>

104. Introduction to Semi-Supervised Learning

Xiaojin Zhu, Andrew B. Goldberg

Synthesis Lectures on Artificial Intelligence and Machine Learning (2009-01) https://doi.org/bq7pm2

DOI: <u>10.2200/s00196ed1v01y200906aim006</u>

105. A Survey on Transfer Learning

Sinno Jialin Pan, Qiang Yang

IEEE Transactions on Knowledge and Data Engineering (2010-10) https://doi.org/bc4vws

DOI: 10.1109/tkde.2009.191

106. A survey of transfer learning

Karl Weiss, Taghi M. Khoshgoftaar, DingDing Wang Journal of Big Data (2016-05-28) https://doi.org/gfkr2w

DOI: 10.1186/s40537-016-0043-6

107. Exploring Semi-supervised Variational Autoencoders for Biomedical Relation Extraction

Yijia Zhang, Zhiyong Lu

arXiv(2019-01-18) https://arxiv.org/abs/1901.06103v1

108. Large-scale extraction of gene interactions from full-text literature using DeepDive

Emily K. Mallory, Ce Zhang, Christopher Ré, Russ B. Altman

Bioinformatics (2015-09-03) https://doi.org/gb5g7b

DOI: 10.1093/bioinformatics/btv476 · PMID: 26338771 · PMCID: PMC4681986

109. Snorkel

Alexander Ratner, Stephen H. Bach, Henry Ehrenberg, Jason Fries, Sen Wu, Christopher Ré *Proceedings of the VLDB Endowment* (2017-11-01) https://doi.org/ch44

DOI: <u>10.14778/3157794.3157797</u> · PMID: <u>29770249</u> · PMCID: <u>PMC5951191</u>

110. Snorkel MeTaL

Alex Ratner, Braden Hancock, Jared Dunnmon, Roger Goldman, Christopher Ré

Proceedings of the Second Workshop on Data Management for End-To-End Machine Learning -

DEEM'18 (2018) https://doi.org/gf3xk7

DOI: 10.1145/3209889.3209898 · PMID: 30931438 · PMCID: PMC6436830

111. Learning protein protein interaction extraction using distant supervision

Philippe Thomas, Illés Solt, Roman Klinger, Ulf Leser (2011-01)

112. Robust Distant Supervision Relation Extraction via Deep Reinforcement Learning

Pengda Qin, Weiran Xu, William Yang Wang

arXiv (2018-05-24) https://arxiv.org/abs/1805.09927v1

113. DSGAN: Generative Adversarial Training for Distant Supervision Relation Extraction

Pengda Qin, Weiran Xu, William Yang Wang *arXiv* (2018-05-24) https://arxiv.org/abs/1805.09929v1

114. Noise Reduction Methods for Distantly Supervised Biomedical Relation Extraction

Gang Li, Cathy Wu, K. Vijay-Shanker

BioNLP 2017 (2017) https://doi.org/ggmk8s

DOI: 10.18653/v1/w17-2323

115. BioInfer: a corpus for information extraction in the biomedical domain

Sampo Pyysalo, Filip Ginter, Juho Heimonen, Jari Björne, Jorma Boberg, Jouni Järvinen, Tapio Salakoski *BMC Bioinformatics* (2007-02-09) https://doi.org/b7bhhc

DOI: 10.1186/1471-2105-8-50 · PMID: 17291334 · PMCID: PMC1808065

116. Learning language in logic - genic interaction extraction challenge

C. Nédellec

Proceedings of the learning language in logic 2005 workshop at the international conference on machine learning (2005)

117. Mining medline: Abstracts, sentences, or phrases?

Jing Ding, Daniel Berleant, Dan Nettleton, Eve Syrkin Wurtele *Pacific symposium on biocomputing* (2002) http://helix-web.stanford.edu/psb02/ding.pdf

118. Concept annotation in the CRAFT corpus

Michael Bada, Miriam Eckert, Donald Evans, Kristin Garcia, Krista Shipley, Dmitry Sitnikov, William A Baumgartner Jr, K Bretonnel Cohen, Karin Verspoor, Judith A Blake, Lawrence E Hunter *BMC Bioinformatics* (2012-07-09) https://doi.org/gb8vdr

DOI: <u>10.1186/1471-2105-13-161</u> · PMID: <u>22776079</u> · PMCID: <u>PMC3476437</u>

119. GRAM: Graph-based Attention Model for Healthcare Representation Learning

Edward Choi, Mohammad Taha Bahadori, Le Song, Walter F. Stewart, Jimeng Sun *arXiv* (2016-11-21) https://arxiv.org/abs/1611.07012v3

120. miRNA-Disease Association Prediction with Collaborative Matrix Factorization

Zhen Shen, You-Hua Zhang, Kyungsook Han, Asoke K. Nandi, Barry Honig, De-Shuang Huang *Complexity* (2017) https://doi.org/ggmrpm

DOI: <u>10.1155/2017/2498957</u>

121. Deep Residual Learning for Image Recognition

Kaiming He, Xiangyu Zhang, Shaoqing Ren, Jian Sun *arXiv* (2015-12-10) https://arxiv.org/abs/1512.03385v1

122. Representation Learning on Graphs: Methods and Applications

William L. Hamilton, Rex Ying, Jure Leskovec *arXiv* (2017-09-17) https://arxiv.org/abs/1709.05584v3

123. Signed laplacian embedding for supervised dimension reduction

Chen Gong, Dacheng Tao, Jie Yang, Keren Fu *Proceedings of the twenty-eighth aaai conference on artificial intelligence* (2014) <u>http://dl.acm.org/citation.cfm?id=2892753.2892809</u>

124. A Semi-NMF-PCA Unified Framework for Data Clustering

Kais Allab, Lazhar Labiod, Mohamed Nadif

IEEE Transactions on Knowledge and Data Engineering (2017-01-01) https://doi.org/f9hm9g

DOI: 10.1109/tkde.2016.2606098

125. Partially supervised graph embedding for positive unlabelled feature selection

Yufei Han, Yun Shen

Proceedings of the twenty-fifth international joint conference on artificial intelligence (2016)

http://dl.acm.org/citation.cfm?id=3060832.3060837

ISBN: <u>978-1-57735-770-4</u>

126. **GraRep**

Shaosheng Cao, Wei Lu, Qiongkai Xu

Proceedings of the 24th ACM International on Conference on Information and Knowledge

Management - CIKM '15 (2015) https://doi.org/gf8rgf

DOI: 10.1145/2806416.2806512

127. Improved Knowledge Base Completion by Path-Augmented TransR Model

Wenhao Huang, Ge Li, Zhi Jin

arXiv (2016-10-06) https://arxiv.org/abs/1610.04073v1

128. A Global Geometric Framework for Nonlinear Dimensionality Reduction

J. B. Tenenbaum

Science (2000-12-22) https://doi.org/cz8wgk

DOI: 10.1126/science.290.5500.2319 · PMID: 11125149

129. Laplacian Eigenmaps for Dimensionality Reduction and Data Representation

Mikhail Belkin, Partha Niyogi

Neural Computation (2003-06) https://doi.org/bbr9cw

DOI: <u>10.1162/089976603321780317</u>

130. Principal component analysis

Svante Wold, Kim Esbensen, Paul Geladi

Chemometrics and Intelligent Laboratory Systems (1987-08) https://doi.org/bm8dnf

DOI: 10.1016/0169-7439(87)80084-9

131. The approximation of one matrix by another of lower rank

Carl Eckart, Gale Young

Psychometrika (1936-09) https://doi.org/c2frtd

DOI: 10.1007/bf02288367

132. Network embedding as matrix factorization: Unifying deepwalk, line, pte, and node2vec

Jiezhong Qiu, Yuxiao Dong, Hao Ma, Jian Li, Kuansan Wang, Jie Tang

Proceedings of the eleventh acm international conference on web search and data mining (2018)

https://doi.org/10.1145/3159652.3159706

DOI: 10.1145/3159652.3159706 · ISBN: 9781450355810

133. A Survey of Collaborative Filtering Techniques

Xiaoyuan Su, Taghi M. Khoshgoftaar

Advances in Artificial Intelligence (2009) https://doi.org/fk9jjg

DOI: 10.1155/2009/421425

134. Graph embedding on biomedical networks: methods, applications and evaluations

Xiang Yue, Zhen Wang, Jingong Huang, Srinivasan Parthasarathy, Soheil Moosavinasab, Yungui Huang, Simon M Lin, Wen Zhang, Ping Zhang, Huan Sun

Bioinformatics (2019-10-04) https://doi.org/ggmzpf
DOI: 10.1093/bioinformatics/btz718 · PMID: 31584634

135. GLEE: Geometric Laplacian Eigenmap Embedding

Leo Torres, Kevin S Chan, Tina Eliassi-Rad

arXiv(2019-05-23) https://arxiv.org/abs/1905.09763v2

DOI: 10.1093/comnet/cnaa007

136. Vicus: Exploiting local structures to improve network-based analysis of biological data

Bo Wang, Lin Huang, Yuke Zhu, Anshul Kundaje, Serafim Batzoglou, Anna Goldenberg

PLOS Computational Biology (2017-10-12) https://doi.org/gb368p

DOI: <u>10.1371/journal.pcbi.1005621</u> · PMID: <u>29023470</u> · PMCID: <u>PMC5638230</u>

137. Translating embeddings for modeling multi-relational data

Antoine Bordes, Nicolas Usunier, Alberto García-Durán, Jason Weston, Oksana Yakhnenko *NIPS* (2013)

138. Knowledge graph embedding by translating on hyperplanes

Zhen Wang, Jianwen Zhang, Jianlin Feng, Zheng Chen *Proceedings of the twenty-eighth aaai conference on artificial intelligence* (2014) http://dl.acm.org/citation.cfm?id=2893873.2894046

139. Learning entity and relation embeddings for knowledge graph completion

Yankai Lin, Zhiyuan Liu, Maosong Sun, Yang Liu, Xuan Zhu

Proceedings of the twenty-ninth aaai conference on artificial intelligence (2015)

http://dl.acm.org/citation.cfm?id=2886521.2886624

ISBN: <u>0-262-51129-0</u>

140. PrTransH: Embedding Probabilistic Medical Knowledge from Real World EMR Data

Linfeng Li, Peng Wang, Yao Wang, Jinpeng Jiang, Buzhou Tang, Jun Yan, Shenghui Wang, Yuting Liu *arXiv* (2019-09-02) https://arxiv.org/abs/1909.00672v1

141. DeepWalk

Bryan Perozzi, Rami Al-Rfou, Steven Skiena

Proceedings of the 20th ACM SIGKDD international conference on Knowledge discovery and data mining - KDD '14 (2014) https://doi.org/gfkpqt

DOI: <u>10.1145/2623330.2623732</u>

142. node2vec: Scalable Feature Learning for Networks

Aditya Grover, Jure Leskovec

arXiv (2016-07-03) https://arxiv.org/abs/1607.00653v1

143. **struc2vec**

Leonardo F. R. Ribeiro, Pedro H. P. Saverese, Daniel R. Figueiredo

Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data Mining - KDD '17 (2017) https://doi.org/gd874b

DOI: 10.1145/3097983.3098061

144. metapath2vec

Yuxiao Dong, Nitesh V. Chawla, Ananthram Swami

Proceedings of the 23rd ACM SIGKDD International Conference on Knowledge Discovery and Data

Mining - KDD '17 (2017) https://doi.org/gfsqzn

DOI: <u>10.1145/3097983.3098036</u>

145. edge2vec: Representation learning using edge semantics for biomedical knowledge discovery

Zheng Gao, Gang Fu, Chunping Ouyang, Satoshi Tsutsui, Xiaozhong Liu, Jeremy Yang, Christopher Gessner, Brian Foote, David Wild, Qi Yu, Ying Ding arXiv (2018-09-07) https://arxiv.org/abs/1809.02269v3

146. Learning Graph Embeddings from WordNet-based Similarity Measures

Andrey Kutuzov, Mohammad Dorgham, Oleksiy Oliynyk, Chris Biemann, Alexander Panchenko *arXiv* (2018-08-16) https://arxiv.org/abs/1808.05611v4

147. Learning to Make Predictions on Graphs with Autoencoders

Phi Vu Tran

2018 IEEE 5th International Conference on Data Science and Advanced Analytics (DSAA) (2018-10)

https://doi.org/ggmzpg

DOI: 10.1109/dsaa.2018.00034

148. Variational Graph Auto-Encoders

Thomas N. Kipf, Max Welling arXiv (2016-11-21) https://arxiv.org/abs/1611.07308v1

149. Adversarially Regularized Graph Autoencoder for Graph Embedding

Shirui Pan, Ruiqi Hu, Guodong Long, Jing Jiang, Lina Yao, Chengqi Zhang *arXiv* (2018-02-13) https://arxiv.org/abs/1802.04407v2

150. Deep learning in neural networks: An overview

Jürgen Schmidhuber

Neural Networks (2015-01) https://doi.org/f6v78n

DOI: 10.1016/j.neunet.2014.09.003 · PMID: 25462637

151. Autoencoders, unsupervised learning and deep architectures

Pierre Baldi

Proceedings of the 2011 international conference on unsupervised and transfer learning workshop - volume 27 (2011)

152. A Comparative Study for Unsupervised Network Representation Learning

Megha Khosla, Vinay Setty, Avishek Anand

IEEE Transactions on Knowledge and Data Engineering (2019) https://doi.org/ggmzph

DOI: 10.1109/tkde.2019.2951398

153. Neural networks for link prediction in realistic biomedical graphs: a multi-dimensional evaluation of graph embedding-based approaches

Gamal Crichton, Yufan Guo, Sampo Pyysalo, Anna Korhonen

BMC Bioinformatics (2018-05-21) https://doi.org/ggkm7q

DOI: <u>10.1186/s12859-018-2163-9</u> · PMID: <u>29783926</u> · PMCID: <u>PMC5963080</u>

154. Network-based integration of multi-omics data for prioritizing cancer genes

Christos Dimitrakopoulos, Sravanth Kumar Hindupur, Luca Häfliger, Jonas Behr, Hesam Montazeri, Michael N Hall, Niko Beerenwinkel

Bioinformatics (2018-03-14) https://doi.org/gc6953

DOI: <u>10.1093/bioinformatics/bty148</u> · PMID: <u>29547932</u> · PMCID: <u>PMC6041755</u>

155. Safe Medicine Recommendation via Medical Knowledge Graph Embedding

Meng Wang, Mengyue Liu, Jun Liu, Sen Wang, Guodong Long, Buyue Qian

156. GAMENet: Graph Augmented MEmory Networks for Recommending Medication Combination

Junyuan Shang, Cao Xiao, Tengfei Ma, Hongyan Li, Jimeng Sun

Proceedings of the AAAI Conference on Artificial Intelligence (2019-07-17) https://doi.org/ggkm7r

DOI: 10.1609/aaai.v33i01.33011126

157. Heterogeneous network embedding for identifying symptom candidate genes

Kuo Yang, Ning Wang, Guangming Liu, Ruyu Wang, Jian Yu, Runshun Zhang, Jianxin Chen, Xuezhong Zhou

Journal of the American Medical Informatics Association (2018-10-23) https://doi.org/gfg6nr

DOI: 10.1093/jamia/ocy117 · PMID: 30357378

158. Predicting Protein-Protein Interactions from Multimodal Biological Data Sources via Nonnegative Matrix Tri-Factorization

Hua Wang, Heng Huang, Chris Ding, Feiping Nie

Journal of Computational Biology (2013-04) https://doi.org/f4thrx

DOI: 10.1089/cmb.2012.0273 · PMID: 23509857

159. Protein functional properties prediction in sparsely-label PPI networks through regularized non-negative matrix factorization

Qingyao Wu, Zhenyu Wang, Chunshan Li, Yunming Ye, Yueping Li, Ning Sun

BMC Systems Biology (2015) https://doi.org/gb5tvr

DOI: <u>10.1186/1752-0509-9-s1-s9</u> · PMID: <u>25708164</u> · PMCID: <u>PMC4331684</u>

160. HMDD v3.0: a database for experimentally supported human microRNA-disease associations

Zhou Huang, Jiangcheng Shi, Yuanxu Gao, Chunmei Cui, Shan Zhang, Jianwei Li, Yuan Zhou, Qinghua Cui

Nucleic Acids Research (2018-10-26) https://doi.org/ggmrph

DOI: 10.1093/nar/gky1010 · PMID: 30364956 · PMCID: PMC6323994

161. Predicting MiRNA-Disease Association by Latent Feature Extraction with Positive Samples

Kai Che, Maozu Guo, Chunyu Wang, Xiaoyan Liu, Xi Chen

Genes (2019-01-24) https://doi.org/ggmrpr

DOI: <u>10.3390/genes10020080</u> · PMID: <u>30682853</u> · PMCID: <u>PMC6410147</u>

162. NPCMF: Nearest Profile-based Collaborative Matrix Factorization method for predicting miRNA-disease associations

Ying-Lian Gao, Zhen Cui, Jin-Xing Liu, Juan Wang, Chun-Hou Zheng

BMC Bioinformatics (2019-06-24) https://doi.org/ggmrpn

DOI: <u>10.1186/s12859-019-2956-5</u> · PMID: <u>31234797</u> · PMCID: <u>PMC6591872</u>

163. RCMF: a robust collaborative matrix factorization method to predict miRNA-disease associations

Zhen Cui, Jin-Xing Liu, Ying-Lian Gao, Chun-Hou Zheng, Juan Wang

BMC Bioinformatics (2019-12) https://doi.org/ggmrpp

DOI: <u>10.1186/s12859-019-3260-0</u> · PMID: <u>31874608</u> · PMCID: <u>PMC6929455</u>

164. LWPCMF: Logistic Weighted Profile-based Collaborative Matrix Factorization for Predicting MiRNA-Disease Associations

Meng-Meng Yin, Zhen Cui, Ming-Ming Gao, Jin-Xing Liu, Ying-Lian Gao

IEEE/ACM Transactions on Computational Biology and Bioinformatics (2019) https://doi.org/ggmrpk

DOI: 10.1109/tcbb.2019.2937774 · PMID: 31478868

165. Protein-protein interaction prediction via Collective Matrix Factorization

Qian Xu, Evan Wei Xiang, Qiang Yang

2010 IEEE International Conference on Bioinformatics and Biomedicine (BIBM) (2010-12)

https://doi.org/csnv5m

DOI: 10.1109/bibm.2010.5706537

166. A network embedding model for pathogenic genes prediction by multi-path random walking on heterogeneous network

Bo Xu, Yu Liu, Shuo Yu, Lei Wang, Jie Dong, Hongfei Lin, Zhihao Yang, Jian Wang, Feng Xia *BMC Medical Genomics* (2019-12) https://doi.org/ggmrpq

DOI: <u>10.1186/s12920-019-0627-z</u> · PMID: <u>31865919</u> · PMCID: <u>PMC6927107</u>

167. Predicting gene-disease associations from the heterogeneous network using graph embedding

Xiaochan Wang, Yuchong Gong, Jing Yi, Wen Zhang 2019 IEEE International Conference on Bioinformatics and Biomedicine (BIBM) (2019-11)

https://doi.org/ggmrpj

DOI: 10.1109/bibm47256.2019.8983134

168. Network embedding-based representation learning for single cell RNA-seq data

Xiangyu Li, Weizheng Chen, Yang Chen, Xuegong Zhang, Jin Gu, Michael Q. Zhang

Nucleic Acids Research (2017-08-28) https://doi.org/ggmrpg

DOI: 10.1093/nar/gkx750 · PMID: 28977434 · PMCID: PMC5737094

169. Neuro-symbolic representation learning on biological knowledge graphs

Mona Alshahrani, Mohammad Asif Khan, Omar Maddouri, Akira R Kinjo, Núria Queralt-Rosinach, Robert Hoehndorf

Bioinformatics (2017-04-25) https://doi.org/gbv6vm

DOI: 10.1093/bioinformatics/btx275 · PMID: 28449114 · PMCID: PMC5860058

170. Deep Learning the Protein Function in Protein Interaction Networks

Kire Trivodaliev, Martin Josifoski, Slobodan Kalajdziski

Communications in Computer and Information Science (2018) https://doi.org/ggmrpd

DOI: <u>10.1007/978-3-030-00825-3 16</u>

171. Detection of protein complexes from multiple protein interaction networks using graph embedding

Xiaoxia Liu, Zhihao Yang, Shengtian Sang, Hongfei Lin, Jian Wang, Bo Xu

Artificial Intelligence in Medicine (2019-05) https://doi.org/ggmrpf

DOI: 10.1016/j.artmed.2019.04.001 · PMID: 31164203

172. Large-scale structural and textual similarity-based mining of knowledge graph to predict drug-drug interactions

Ibrahim Abdelaziz, Achille Fokoue, Oktie Hassanzadeh, Ping Zhang, Mohammad Sadoghi *Journal of Web Semantics* (2017-05) https://doi.org/gcrwk3

DOI: <u>10.1016/j.websem.2017.06.002</u>

173. Matrix Factorization-Based Prediction of Novel Drug Indications by Integrating Genomic Space

Wen Dai, Xi Liu, Yibo Gao, Lin Chen, Jianglong Song, Di Chen, Kuo Gao, Yongshi Jiang, Yiping Yang,

Jianxin Chen, Peng Lu

Computational and Mathematical Methods in Medicine (2015) https://doi.org/gb58g8

DOI: 10.1155/2015/275045 · PMID: 26078775 · PMCID: PMC4452507

174. Abstract

eLife Sciences Publications, Ltd

https://doi.org/gf4fdb

DOI: 10.7554/elife.26726.001

175. Drug-Target Interaction Prediction with Graph Regularized Matrix Factorization

Ali Ezzat, Peilin Zhao, Min Wu, Xiao-Li Li, Chee-Keong Kwoh

IEEE/ACM Transactions on Computational Biology and Bioinformatics (2017-05-01)

https://doi.org/ggmrrp

DOI: 10.1109/tcbb.2016.2530062 · PMID: 26890921

176. Predicting Drug-Target Interaction Using Deep Matrix Factorization

Hafez Eslami Manoochehri, Mehrdad Nourani

2018 IEEE Biomedical Circuits and Systems Conference (BioCAS) (2018-10) https://doi.org/ggmrrn

DOI: <u>10.1109/biocas.2018.8584817</u>

177. Network-based prediction of drug-target interactions using an arbitrary-order proximity embedded deep forest

Xiangxiang Zeng, Siyi Zhu, Yuan Hou, Pengyue Zhang, Lang Li, Jing Li, L Frank Huang, Stephen J Lewis, Ruth Nussinov, Feixiong Cheng

Bioinformatics (2020-01-23) https://doi.org/ggmrrk

DOI: <u>10.1093/bioinformatics/btaa010</u> · PMID: <u>31971579</u>

178. DrPOCS: Drug Repositioning Based on Projection Onto Convex Sets

Yin-Ying Wang, Chunfeng Cui, Ligun Qi, Hong Yan, Xing-Ming Zhao

IEEE/ACM Transactions on Computational Biology and Bioinformatics (2019-01-01)

https://doi.org/ggmrrq

DOI: 10.1109/tcbb.2018.2830384 · PMID: 29993698

179. Neighborhood Regularized Logistic Matrix Factorization for Drug-Target Interaction Prediction

Yong Liu, Min Wu, Chunyan Miao, Peilin Zhao, Xiao-Li Li

PLOS Computational Biology (2016-02-12) https://doi.org/ggmrrw

DOI: <u>10.1371/journal.pcbi.1004760</u> · PMID: <u>26872142</u> · PMCID: <u>PMC4752318</u>

180. Predicting drug-target interactions by dual-network integrated logistic matrix factorization

Ming Hao, Stephen H. Bryant, Yanli Wang

Scientific Reports (2017-01-12) https://doi.org/ggmrrj

DOI: 10.1038/srep40376 · PMID: 28079135 · PMCID: PMC5227688

181. Drug-Disease Association and Drug-Repositioning Predictions in Complex Diseases Using Causal Inference-Probabilistic Matrix Factorization

Jihong Yang, Zheng Li, Xiaohui Fan, Yiyu Cheng

Journal of Chemical Information and Modeling (2014-08-22) https://doi.org/f6hpb4

DOI: <u>10.1021/ci500340n</u> · PMID: <u>25116798</u>

182. Predicting drug-disease associations by using similarity constrained matrix factorization

Wen Zhang, Xiang Yue, Weiran Lin, Wenjian Wu, Ruoqi Liu, Feng Huang, Feng Liu

BMC Bioinformatics (2018-06-19) https://doi.org/ggmrrt

DOI: 10.1186/s12859-018-2220-4 · PMID: 29914348 · PMCID: PMC6006580

183. Deep mining heterogeneous networks of biomedical linked data to predict novel drugtarget associations

Nansu Zong, Hyeoneui Kim, Victoria Ngo, Olivier Harismendy

Bioinformatics (2017-04-18) https://doi.org/gbqjgx

DOI: <u>10.1093/bioinformatics/btx160</u> · PMID: <u>28430977</u> · PMCID: <u>PMC5860112</u>

184. Scalable and Accurate Drug-target Prediction Based on Heterogeneous Bio-linked Network Mining

Nansu Zong, Rachael Sze Nga Wong, Victoria Ngo, Yue Yu, Ning Li

Cold Spring Harbor Laboratory (2019-02-03) https://doi.org/ggmrrm

DOI: 10.1101/539643

185. Drug Similarity Integration Through Attentive Multi-view Graph Auto-Encoders

Tengfei Ma, Cao Xiao, Jiayu Zhou, Fei Wang

arXiv (2018-04-28) https://arxiv.org/abs/1804.10850v1

186. Modeling polypharmacy side effects with graph convolutional networks

Marinka Zitnik, Monica Agrawal, Jure Leskovec

Bioinformatics (2018-06-27) https://doi.org/gfgn55

DOI: 10.1093/bioinformatics/bty294 · PMID: 29949996 · PMCID: PMC6022705

187. DrugBank: a knowledgebase for drugs, drug actions and drug targets

David S. Wishart, Craig Knox, An Chi Guo, Dean Cheng, Savita Shrivastava, Dan Tzur, Bijaya Gautam, Murtaza Hassanali

Nucleic Acids Research (2007-11-29) https://doi.org/d3qqpj

DOI: 10.1093/nar/gkm958 · PMID: 18048412 · PMCID: PMC2238889

188. The human disease network

K.-I. Goh, M. E. Cusick, D. Valle, B. Childs, M. Vidal, A.-L. Barabasi

Proceedings of the National Academy of Sciences (2007-05-14) https://doi.org/bt6qvc

DOI: <u>10.1073/pnas.0701361104</u> · PMID: <u>17502601</u> · PMCID: <u>PMC1885563</u>

189. Evaluation of knowledge graph embedding approaches for drug-drug interaction prediction in realistic settings

Remzi Celebi, Huseyin Uyar, Erkan Yasar, Ozgur Gumus, Oguz Dikenelli, Michel Dumontier *BMC Bioinformatics* (2019-12) https://doi.org/ggmrrv

DOI: <u>10.1186/s12859-019-3284-5</u> · PMID: <u>31852427</u> · PMCID: <u>PMC6921491</u>

190. Drug-Drug Interaction Prediction Based on Knowledge Graph Embeddings and Convolutional-LSTM Network

Md. Rezaul Karim, Michael Cochez, Joao Bosco Jares, Mamtaz Uddin, Oya Beyan, Stefan Decker *Proceedings of the 10th ACM International Conference on Bioinformatics, Computational Biology and Health Informatics - BCB '19* (2019) https://doi.org/ggmrrs

DOI: 10.1145/3307339.3342161

191. Mining Electronic Health Records using Linked Data.

David J Odgers, Michel Dumontier

AMIA Joint Summits on Translational Science proceedings. AMIA Joint Summits on Translational Science (2015-03-23) https://www.ncbi.nlm.nih.gov/pubmed/26306276

PMID: <u>26306276</u> · PMCID: <u>PMC4525267</u>

192. Applying linked data principles to represent patient's electronic health records at Mayo clinic

Jyotishman Pathak, Richard C. Kiefer, Christopher G. Chute

Proceedings of the 2nd ACM SIGHIT symposium on International health informatics - IHI '12 (2012) https://doi.org/fzm2p7

DOI: 10.1145/2110363.2110415

193. PDD Graph: Bridging Electronic Medical Records and Biomedical Knowledge Graphs via Entity Linking

Meng Wang, Jiaheng Zhang, Jun Liu, Wei Hu, Sen Wang, Xue Li, Wenqiang Liu *arXiv* (2017-07-17) https://arxiv.org/abs/1707.05340v2

194. Diagnosis Code Assignment Using Sparsity-Based Disease Correlation Embedding

Sen Wang, Xiaojun Chang, Xue Li, Guodong Long, Lina Yao, Quan Z. Sheng

IEEE Transactions on Knowledge and Data Engineering (2016-12-01) https://doi.org/f9cgtv

DOI: 10.1109/tkde.2016.2605687

195. EMR-based medical knowledge representation and inference via Markov random fields and distributed representation learning

Chao Zhao, Jingchi Jiang, Yi Guan *arXiv* (2017-09-20) https://arxiv.org/abs/1709.06908v1

196. Attention Is All You Need

Ashish Vaswani, Noam Shazeer, Niki Parmar, Jakob Uszkoreit, Llion Jones, Aidan N. Gomez, Lukasz Kaiser, Illia Polosukhin

arXiv (2017-06-12) https://arxiv.org/abs/1706.03762v5

197. Neural Machine Translation by Jointly Learning to Align and Translate

Dzmitry Bahdanau, Kyunghyun Cho, Yoshua Bengio *arXiv* (2014-09-01) https://arxiv.org/abs/1409.0473v7

198. Learning the Graphical Structure of Electronic Health Records with Graph Convolutional Transformer

Edward Choi, Zhen Xu, Yujia Li, Michael W. Dusenberry, Gerardo Flores, Yuan Xue, Andrew M. Dai *arXiv* (2019-06-11) https://arxiv.org/abs/1906.04716v3

199. The probability of edge existence due to node degree: a baseline for network-based predictions

Michael Zietz, Daniel S. Himmelstein, Kyle Kloster, Christopher Williams, Michael W. Nagle, Blair D. Sullivan, Casey S. Greene

Manubot (2020-03-05) https://greenelab.github.io/xswap-manuscript/