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

A DOI-citable version of this manuscript is available at https://doi.org/10.1101/730085.

This manuscript (<u>permalink</u>) was automatically generated from <u>greenelab/text_mined_hetnet_manuscript@8e28137</u> on March 29, 2022.

Authors

David N. Nicholson

Department of Systems Pharmacology and Translational Therapeutics, University of Pennsylvania · Funded by GBMF4552

Daniel S. Himmelstein

D 0000-0002-3012-7446 · ○ dhimmel · У dhimmel

Department of Systems Pharmacology and Translational Therapeutics, University of Pennsylvania · Funded by GBMF4552

Casey S. Greene

© 0000-0001-8713-9213 • ☐ cgreene • У GreeneScientist

Department of Systems Pharmacology and Translational Therapeutics, University of Pennsylvania · Funded by GBMF4552 and R01 HG010067

Abstract

Knowledge graphs support multiple research efforts by providing contextual information for biomedical entities, constructing networks, and supporting the interpretation of high-throughput analyses. These databases are populated via some form of manual curation, which is difficult to scale in the context of an increasing publication rate. Data programming is a paradigm that circumvents this arduous manual process by combining databases with simple rules and heuristics written as label functions, which are programs designed to automatically annotate textual data. Unfortunately, writing a useful label function requires substantial error analysis and is a nontrivial task that takes multiple days per function. This makes populating a knowledge graph with multiple nodes and edge types practically infeasible. We sought to accelerate the label function creation process by evaluating the extent to which label functions could be re-used across multiple edge types. We used a subset of an existing knowledge graph centered on disease, compound, and gene entities to evaluate label function re-use. We determined the best label function combination by comparing a baseline database-only model with the same model but added edge-specific or edge-mismatch label functions. We confirmed that adding additional edge-specific rather than edge-mismatch label functions often improves text annotation and shows that this approach can incorporate novel edges into our source knowledge graph. We expect that continued development of this strategy has the potential to swiftly populate knowledge graphs with new discoveries, ensuring that these resources include cutting-edge results.

Introduction

Knowledge bases are important resources that hold complex structured and unstructured information. These resources have been used in important tasks such as network analysis for drug repurposing discovery [1]–[3] or as a source of training labels for text mining systems [4]–[6]. Populating knowledge bases often requires highly trained scientists to read biomedical literature and summarize the results [7]. This time-consuming process is referred to as manual curation. In 2007, researchers estimated that filling a knowledge base via manual curation would require approximately 8.4 years to complete [8]. The rate of publications continues to exponentially increase [9], so using only manual curation to fully populate a knowledge base has become impractical.

Relationship extraction has been studied as a solution towards handling the challenge posed by an exponentially growing body of literature [7]. This process consists of creating an expert system to automatically scan, detect and extract relationships from textual sources. Typically, these systems utilize machine learning techniques that require extensive corpora of well-labeled training data. These corpora are difficult to obtain, because they are constructed via extensive manual curation pipelines.

Distant supervision is a technique also designed to sidestep the dependence on manual curation and quickly generate large training datasets. This technique assumes that positive examples established in selected databases can be applied to any sentence that contains them [4]. The central problem with this technique is that generated labels are often of low quality which results in an expansive amount of false positives raw:Jiang2018RevisitingDS?.

Ratner et al. [10] recently introduced "data programming" as a solution. Data programming is a paradigm that combines distant supervision with simple rules and heuristics written as small programs called label functions. These label functions are consolidated via a noise aware generative model that is designed to produce training labels for large datasets. Using this paradigm can dramatically reduce the time required to obtain sufficient training data; however, writing a useful label function requires a significant amount of time and error analysis. This dependency makes

constructing a knowledge base with a myriad of heterogenous relationships nearly impossible as tens or possibly hundreds of label functions are required per relationship type.

In this paper, we seek to accelerate the label function creation process by measuring the extent to which label functions can be re-used across different relationship types. We hypothesize that sentences describing one relationship type may share linguistic features such as keywords or sentence structure with sentences describing other relationship types. We conducted a series of experiments to determine the degree to which label function re-use enhanced performance over distant supervision alone. We focus on relationships that indicate similar types of physical interactions (i.e., gene-binds-gene and compound-binds-gene) as well as different types (i.e., disease-associates-gene and compound-treats-disease). Re-using label functions could dramatically reduce the time required to populate a knowledge base with a multitude of heterogeneous relationships.

Related Work

Relationship extraction is the process of detecting semantic relationships from a collection of text. This process can be broken down into three different categories: (1) the use of natural language processing techniques such as manually crafted rules and heuristics for relationship extraction (Rule Based Extractors), (2) the use of unsupervised methods such as co-occurrence scores or clustering to find patterns within sentences and documents (Unsupervised Extractors), and (3) the use of supervised or semi-supervised machine learning for classifying the presence of a relation within documents or sentences (Supervised Extractors). In this section, we briefly discuss selected efforts under each category.

Rule Based Extractors

Rule based extractors rely heavily on expert knowledge to perform extraction. Typically, these systems use linguistic rules and heuristics to identify key sentences or phrases. For example, a hypothetical extractor focused on protein phosphorylation events would identify sentences containing the phrase "gene X phosphorylates gene Y" [11]. This phrase is a straightforward indication that two genes have a fundamental role in protein phosphorylation. Other phrase extractors have been used to identify drug-disease treatments tag:ctd medline?, pharmcogenomic events tag:pharmpresso? and protein-protein interactions tag:ppinterfinder?, tag:hpiminer?. These extractors provide a simple and effective way to extract sentences; however, they depend on extensive knowledge about the text to be properly constructed.

A sentence's grammatical structure can also support relationship extraction via dependency trees. Dependency trees are data structures that depict a sentence's grammatical relation structure in the form of nodes and edges. Nodes represent words and edges represent the dependency type each word shares between one another. For example, a possible extractor would classify sentences as a positive if a sentence contained the following dependency tree path: "gene X (subject)-> promotes (verb)<- cell death (direct object) <- in (preposition) <-tumors (object of preposition)" tag:pkde4j?. This approach provides extremely precise results, but the quantity of positive results remains modest as sentences appear in distinct forms and structure. Because of this limitation, recent approaches have incorporated methods on top of rule based extractors such as co-occurrence and machine learning systems [12], tag:limtox?. We discuss the pros and cons of added methods in a later section. For this project, we constructed our label functions without the aid of these works; however, approaches discussed in this section provide substantial inspiration for novel label functions in future endeavors.

Unsupervised Extractors

Unsupervised extractors detect relationships without the need of annotated text. Notable approaches exploit the fact that two entities can occur together in text. This event is referred to as co-occurrence. Extractors utilize these events by generating statistics on the frequency of entity pairs occurring in text. For example, a possible extractor would say gene X is associated with disease Y, because gene X and disease Y appear together more often than individually tag:diseases?. This approach has been used to establish the following relationship types: disease-gene relationships tag:diseases?, tag:polysearch?, tag:dg text pubmed?, tag:lgscore?, tag:full text co abstracts?, tag:copub discovery?, protein-protein interactions [13], tag:protein protein co network?, tag:full text co abstracts?, drug-disease treatments tag:abc drugs?, and tissue-gene relations [14]. Extractors using the co-occurrence strategy provide exceptional recall results; however, these methods may fail to detect underreported relationships, because they depend on entity-pair frequency for detection. Junge et al. created a hybrid approach to account for this issue using distant supervision to train a classifier to learn the context of each sentence tag:cocoscore?. Once the classifier was trained, they scored every sentence within their corpus, and each sentence's score was incorporated into calculating co-occurrence frequencies to establish relationship existence tag:cocoscore?. Co-occurrence approaches are powerful in establishing edges on the global scale; however, they cannot identify individual sentences without the need for supervised methods.

Clustering is an unsupervised approach that extracts relationships from text by grouping similar sentences together. Percha et al. used this technique to group sentences based on their grammatical structure tag;global network? Using Stanford's Core NLP Parser [15], a dependency tree was generated for every sentence in each Pubmed abstract tag;global network? Each tree was clustered based on similarity and each cluster was manually annotated to determine which relationship each group represented tag;global network? For our project we incorporated the results of this work as domain heuristic label functions. Overall, unsupervised approaches are desirable since they do not require well-annotated training data. Such approaches provide excellent recall; however, performance can be limited in terms of precision when compared to supervised machine learning methods [16], [17].

Supervised Extractors

Supervised extractors consist of training a machine learning classifier to predict the existence of a relationship within text. These classifiers require access to well-annotated datasets, which are usually created via some form of manual curation. Previous work consists of research experts curating their own datasets to train classifiers tag:befree?, tag:bioinfer?, tag:hprd50?; however, there have been community-wide efforts to create datasets for shared tasks [18], [19]], tag:biocreative v?. Shared tasks are open challenges that aim to build the best classifier for natural language processing tasks such as named entity tagging or relationship extraction. A notable example is the BioCreative community that hosted a number of shared tasks such as predicting compound-protein interactions (BioCreative VI track 5) [18] and compound induced diseases [19]. Often these datasets are well annotated, but are modest in size (2,432 abstracts for BioCreative VI [18] and 1500 abstracts for BioCreative V [19]). As machine learning classifiers become increasingly complex, these small dataset sizes cannot suffice. Plus, these multitude of datasets are uniquely annotated which can generate noticeable differences in terms of classifier performance [19]). Overall, obtaining large well-annotated datasets still remains as an open non-trivial task.

Before the rise of deep learning, a classifier that was most frequently used was support vector machines. This classifier uses a projection function called a kernel to map data onto a high dimensional space so datapoints can be easily discerned between classes [20]. This method was used to extract disease-gene associations tag:befree?, tag:dtminer?, tag:ensemble_svm?, protein-protein interactionstag:ppi_graph_kernels?, tag:limtox?, tag:lptk? and protein docking information tag:protein_docking?. Generally, support vector machines perform well on small datasets with large feature spaces but are slow to train as the number of datapoints becomes asymptotically large.

Deep learning has been increasingly popular as these methods can outperform common machine learning methods [21]. Approaches in this field consist of using various neural network architectures, such as recurrent neural networks tag:ppi bilstm?, tag:cbg ensemble dl?, tag:cbg neural attention?, tag:recursive nn?, tag:semi supervised vae?, tag:biobert? and convolutional neural networks tag:ppi deep conv?, tag:mcdepcnn?, tag:cbg ensemble dl?, tag:semi supervised vae?, tag:cbg transfer learning?, to extract relationships from text. In fact approaches in this field were the winning model within the BioCreative VI shared task [18], raw:chemprot winner?. Despite the substantial success of these models, they often require large amounts of data to perform well. Obtaining large datasets is a time-consuming task, which makes training these models a non-trivial challenge. Distant supervision has been used as a solution to fix the barren amount of large datasets [4]. Approaches have used this paradigm to extract chemicalgene interactions tag:semi supervised vae?, disease-gene associations tag:cocoscore? and protein-protein interactions tag:deep dive?, tag:cocoscore?, tag:semi supervised vae?. In fact, efforts done in tag:deep dive? served as one of the motivating rationales for our work.

Overall, deep learning has provided exceptional results in terms of relationships extraction. Thus, we decided to use a deep neural network as our discriminative model.

Methods and Materials

Hetionet

Hetionet v1 [3] is a heterogeneous network that contains pharmacological and biological information. This network depicts information in the form of nodes and edges of different types. Nodes in this network represent biological and pharmacological entities, while edges represent relationships between entities. Hetionet v1 contains 47,031 nodes with 11 different data types and 2,250,197 edges that represent 24 different relationship types (Figure 1). Edges in Hetionet v1 were obtained from open databases, such as the GWAS Catalog [22], Human Interaction database [23] and DrugBank [24]. For this project, we analyzed performance over a subset of the Hetionet v1 edge types: disease associates with a gene (DaG), compound binds to a gene (CbG), compound treating a disease (CtD), and gene interacts with gene (GiG) (bolded in Figure 1).

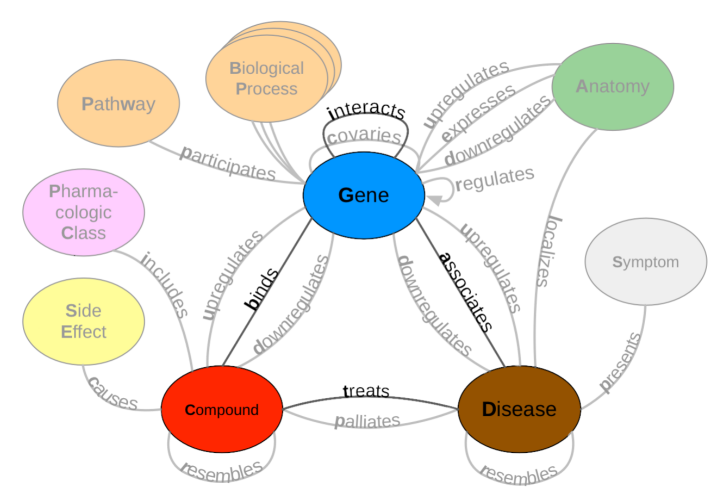


Figure 1: A metagraph (schema) of Hetionet v1 where biomedical entities are represented as nodes and the relationships between them are represented as edges. We examined performance on the highlighted subgraph; however, the long-term vision is to capture edges for the entire graph.

Dataset

We used PubTator Central [25] as input to our analysis. PubTator Central provides MEDLINE abstracts that have been annotated with well-established entity recognition tools including Tagger One [26] for disease, chemical and cell line entities, tmVar [27] for genetic variation tagging, GNormPlus [28] for gene entities and SR4GN [29] for species entities. We downloaded PubTator Central on March 1, 2020, at which point it contained approximately 30,000,000 documents. After downloading, we filtered out annotated entities that were not contained in Hetionet v1. We extracted sentences with two or more annotations and termed these sentences as candidate sentences. We used the Spacy's English natural language processing (NLP) pipeline (en_core_web_sm) [30] to generate dependency trees and parts of speech tags for every extracted candidate sentence. Each candidate sentence was stratified by their corresponding abstract ID to produce a training set, tuning set, and a testing set. We used random assortment to assign dataset labels to each abstract. Every abstract had a 70% chance of being labeled training, 20% chance of being labeled tuning, and 10% chance of being labeled testing. Despite the power of data programming, all text mining systems need to have ground truth labels to be well-calibrated. We hand-labeled five hundred to a thousand candidate sentences of each edge type to obtain a ground truth set (Table 1).

Table 1: Statistics of Candidate Sentences. We sorted each abstract into a training, tuning and testing set. Numbers in parentheses show the number of positives and negatives that resulted from the hand-labeling process.

| Relationship | Train | Tune | Test |
|-------------------------|--------|-------------------|-------------------|
| Disease Associates Gene | 2.49 M | 696K (397+, 603-) | 348K (351+, 649-) |

| Relationship | Train | Tune | Test |
|-------------------------|-------|-------------------|-------------------|
| Compound Binds Gene | 2.4M | 684K (37+, 463-) | 341k (31+, 469-) |
| Compound Treats Disease | 1.5M | 441K (96+, 404-) | 223K (112+, 388-) |
| Gene Interacts Gene | 11.2M | 2.19M (60+, 440-) | 1.62M (76+, 424-) |

Label Functions for Annotating Sentences

The challenge of having too few ground truth annotations is familiar to many natural language processing applications, even when unannotated text is abundant. Data programming circumvents this issue by quickly annotating large datasets using multiple noisy signals emitted by label functions [10]. Label functions are simple pythonic functions that emit: a positive label (1), a negative label (0), or abstain from emitting a label (-1). These functions can use different approaches or techniques to emit a label; however, these functions can be grouped into simple categories discussed below. Once constructed, these functions are combined using a generative model to output a single annotation. This single annotation is a consensus probability score bounded between 0 (low chance of mentioning a relationship) and 1 (high chance of mentioning a relationship). We used these annotations to train a discriminative model for the final classification step.

Label Function Categories

Label functions can be constructed in various ways; however, they also share similar characteristics. We grouped functions into databases and text patterns. The majority of our label functions fall into the text pattern category (Supplemental Table 2). Further, we described each label function category and provided an example that refers to the following candidate sentence: "PTK6 may be a novel therapeutic target for pancreatic cancer".

Databases: These label functions incorporate existing databases to generate a signal, as seen in distant supervision [4]. These functions detect if a candidate sentence's co-mention pair is present in a given database. Our label function emits a positive label if the pair is present and abstains otherwise. If the pair is not present in any existing database, a separate label function emits a negative label. We used a separate label function to prevent a label imbalance problem, which can occur when a single function labels every possible sentence despite being correct or not. If this problem isn't handled correctly, the generative model could become biased and only emit one prediction (solely positive or solely negative) for every sentence.

$$\Lambda_{DB}(extstyle{D}, extstyle{G}) = \left\{egin{array}{ll} 1 & (extstyle{D}, extstyle{G}) \in DB \ 0 & otherwise \end{array}
ight.$$

$$\Lambda_{
eg DB}(extbf{ extit{D}}, extbf{ extit{G}}) = egin{cases} -1 & (extbf{ extit{D}}, extbf{ extit{G}})
otherwise \end{cases}$$

Text Patterns: These label functions are designed to use keywords or sentence context to generate a signal. For example, a label function could focus on the number of words between two mentions and emit a label if two mentions are too close. Alternatively, a label function could focus on the parts of speech contained within a sentence and ensures a verb is present. Besides parts of speech, a label function could exploit dependency parse trees to emit a label. These trees are akin to the tree data structure where words are nodes and edges are how each word modifies each other. Label functions that use these parse trees will test if the generated tree matches a pattern and emits a positive label if true. For our analysis, we used previously identified patterns designed for biomedical text to generate our label functions tag; global network?.

$$\Lambda_{TP}(D, G) = egin{cases} 1 & "target" \in Candidate\ Sentence \ -1 & otherwise \end{cases}$$

$$\Lambda_{TP}(extbf{ extit{D}}, extbf{ extit{G}}) = egin{cases} 0 & "VB"
otin pos_tags(Candidate Sentence) \ -1 & otherwise \end{cases}$$

$$\Lambda_{TP}(extstyle{D}, extstyle{G}) = egin{cases} 1 & dep(Candidate\ Sentence) \in Cluster\ Theme \ -1 & otherwise \end{cases}$$

Each text pattern label function was constructed via manual examination of sentences within the training set. For example, using the candidate sentence above, one would identify the phrase "novel therapeutic target" and incorporate this phrase into a global list that a label function would use to check if present in a sentence. After initial construction, we tested and augmented the label function using sentences in the tune set. We repeated this process for every label function in our repertoire.

Table 2: The distribution of each label function per relationship.

| Relationship | Databases (DB) | Text Patterns (TP) |
|--------------|----------------|--------------------|
| DaG | 7 | 30 |
| CtD | 3 | 22 |
| CbG | 9 | 20 |
| GiG | 9 | 28 |

Experimental Design

Being able to re-use label functions across edge types would substantially reduce the number of label functions required to extract multiple relationships from biomedical literature. We first established a baseline by training a generative model using only distant supervision label functions designed for the target edge type (see Supplemental Methods). For example, in the Gene interacts Gene (GiG) edge type we used label functions that returned a 1 if the pair of genes were included in the Human Interaction database [23], the iRefIndex database [31] or in the Incomplete Interactome database [32]. Then we compared the baseline model with models that also included text and domain-heuristic label functions. Using a sampling with replacement approach, we sampled these text and domain-heuristic label functions separately within edge types, across edge types, and from a pool of all label functions. We compared within-edge-type performance to across-edge-type and all-edge-type performance. For each edge type we sampled a fixed number of label functions consisting of five evenly spaced numbers between one and the total number of possible label functions. We repeated this sampling process 50 times for each point. Furthermore, at each point we also trained the discriminative model using annotations from the generative model trained on edge-specific label functions (see Supplemental Methods). We report performance of both models in terms of the area under the receiver operating characteristic curve (AUROC) and the area under the precision-recall curve (AUPR). Ensuing model evaluations, we quantified the number of edges we could incorporate into Hetionet v1. Using a calibrated discriminative model (see Supplemental Methods), we scored every candidate sentence within our dataset and grouped candidates based on their mention pair. We took the max score within each candidate group and this score represents the probability of the existence of an edge. We established edges by using a cutoff score that produced an equal error rate between the false positives and false negatives. We report the number of preexisting edges we could recall as well as the number of novel edges we can incorporate. Lastly, we compared our framework with a previously established unsupervised approach tag:cocoscore?.

Results

Generative Model Using Randomly Sampled Label Functions

Creating label functions is a labor-intensive process that can take days to accomplish. We sought to accelerate this process by measuring the extent to which label functions can be reused. Our hypothesis was that certain edge types share similar linguistic features such as keywords and/or sentence structure. This shared characteristic would make certain edge types amenable to label function reuse. We designed a set of experiments to test this hypothesis on an individual level (edge vs edge) as well as a global level (collective pool of sources). We observed that performance increased when edge-specific label functions were added to an edge-specific baseline model, while label function reuse usually provided less benefit (AUROC Figure 2, AUPR Supplemental Figure 5). We also evaluated randomly selecting label functions from among all sets and observed similar performance (AUROC Supplemental Figure 6, AUPR Supplemental Figure 7) The quintessential example of this overarching trend is the Compound treats Disease (CtD) edge type, where edge-specific label functions always outperformed transferred label functions. However, there are hints of label function transferability for selected edge types and label function sources. Performance increases as more CbG label functions are incorporated to the GiG baseline model and vice versa. This suggests that sentences for GiG and CbG may share similar linguistic features or terminology that allows for label functions to be reused. Perplexingly, edge-specific Disease associates Gene (DaG) label functions did not improve performance over label functions drawn from other edge types. Overall, only CbG and GiG showed significant signs of reusability which suggests label functions could be shared between the two edge types.

Test Set AUROC of Predicted Relations

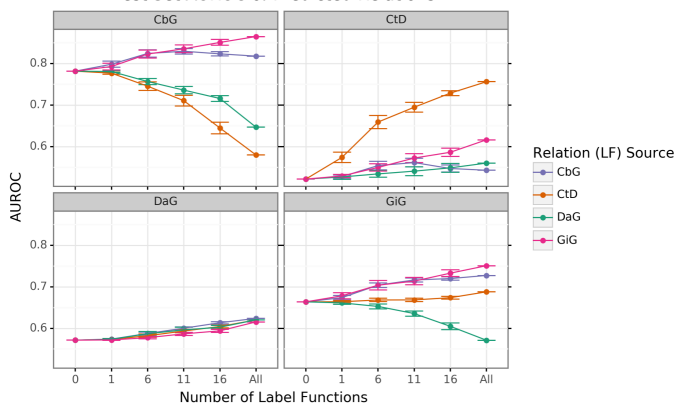


Figure 2: Edge-specific label functions are better performing than edge-mismatch label functions, but certain mismatch situations show signs of successful transfer. Each line plot header depicts the edge type the generative model is trying to predict, while the colors represent the source of label functions. For example, orange represents sampling label functions designed to predict the Compound treats Disease (CtD) edge type. The x axis shows the number of randomly sampled label functions being incorporated into the database-only baseline model (point at 0). The y axis shows area under the receiver operating curve (AUROC). Each point on the plot shows the average of 50 sample runs, while the

error bars show the 95% confidence intervals of all runs. The baseline and "All" data points consist of sampling from the entire fixed set of label functions.

We found that sampling from all label function sources at once usually underperformed relative to edge-specific label functions (Supplemental Figures 6 and 7). As more label functions were sampled, the gap between edge-specific sources and all sources widened. CbG is a prime example of this trend (Supplemental Figures 6 and 7), while CtD and GiG show a similar but milder trend. DaG was the exception to the general rule: the pooled set of label functions improved performance over the edge-specific ones, which aligns with the previously observed results for individual edge types (Figure 2). The decreasing trend when pooling all label functions supports the notion that label functions cannot easily transfer between edge types (exception being CbG on GiG and vice versa).

Discriminative Model Performance

The discriminative model is designed to augment performance over the generative model by incorporating textual features along with estimated training labels. The discriminative model is a piecewise convolutional neural network trained over word embeddings (See Methods and Materials). We found that the discriminative model generally out-performed the generative model as more edgespecific label functions are incorporated (Figure 3 and Supplemental Figure 8). The discriminative model's performance is often poorest when very few edge-specific label functions are added to the baseline model (seen in Disease associates Gene (DaG), Compound binds Gene (CbG) and Gene interacts Gene (GiG)). This suggests that generative models trained with more label functions produce outputs that are more suitable for training discriminative models. An exception to this trend is Compound treats Disease (CtD) where the discriminative model out-performs the generative model at all levels of sampling. We observed the opposite trend with the Compound-binds-Gene (CbG) edges: the discriminative model was always poorer or indistinguishable from the generative model. Interestingly, the AUPR for CbG plateaus below the generative model and decreases when all edgespecific label functions are used (Supplemental Figure 8). This suggests that the discriminative model might be predicting more false positives in this setting. Incorporating more edge-specific label functions usually improves performance for the discriminative model over the generative model.

Discriminative Model Performance (Test Set)

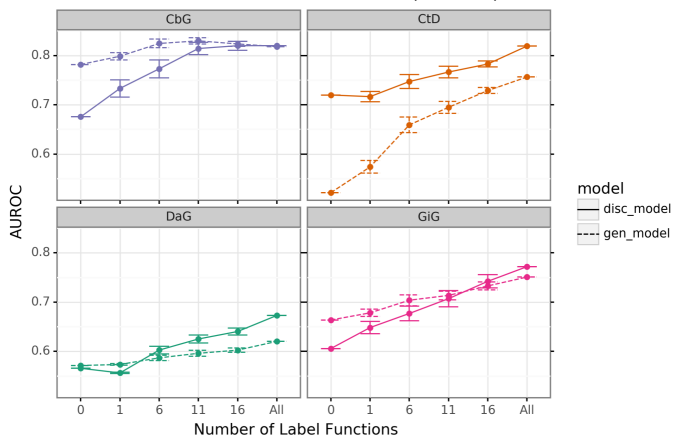


Figure 3: The discriminative model usually improves at a faster rate than the generative model as more edge-specific label function are included. The line plot headers represent the specific edge type the discriminative model is trying to predict. The x-axis shows the number of randomly sampled label functions that are incorporated into the baseline model (point at 0). The y axis shows the area under the receiver operating curve (AUROC). Each datapoint represents the average of 50 sample runs and the error bars represent the 95% confidence interval of each run. The baseline and "All" data points consist of sampling from the entire fixed set of label functions.

Discussion

We measured the extent to which label functions can be re-used across multiple edge types to extract relationships from literature. Through our sampling experiment, we found that adding edge-specific label functions increases performance for the generative model (Figure 2). We found that label functions designed from relatively related edge types can increase performance (Gene interacts Gene (GiG) label functions predicting the Compound binds Gene (CbG) edge and vice versa), while the Disease associates Gene (DaG) edge type remained agnostic to label function sources (Figure 2 and Supplemental Figure 5). Furthermore, we found that using all label functions at once generally hurts performance with the exception being the DaG edge type (Supplemental Figures 6 and 7). One possibility for this observation is that DaG is a broadly defined edge type. For example, DaG may contain many concepts related to other edge types such as Disease (up/down) regulating a Gene, which makes it more agnostic to label function sources (examples highlighted in our annotated sentences).

Regarding the discriminative model, adding edge-specific label function substantially improved performance for two out of the four edge types (Compound treats Disease (CtD) and Disease associates Gene (DaG)) (Figure 3 and Supplemental Figure 8). Gene interacts Gene (GiG) and Compound binds Gene (CbG) discriminative models showed minor improvements compared to the generative model, but only when nearly all edge-specific label functions are included (Figure 3 and Supplemental Figure 8). We came across a large amount of spurious gene mentions when working

with the discriminative model and believe that these mentions contributed to CbG and GiG's hindered performance. We encountered difficulty in calibrating each discriminative model (Supplemental Figure 9). The temperature scaling algorithm appears to improve calibration for the highest scores for each model but did not successfully calibrate throughout the entire range of predictions. Improving performance for all predictions may require more labeled examples or may be a limitation of the approach in this setting. Even with these limitations, this early-stage approach could recall many existing edges from an existing knowledge base, Hetionet v1, and suggest many new high-confidence edges for inclusion (Supplemental Figure 10). Our findings suggest that further work, including an expansion of edge types and a move to full text from abstracts, may make this approach suitable for building continuously updated knowledge bases to address drug repositioning and other biomedical challenges.

Conclusion and Future Direction

Filling out knowledge bases via manual curation can be an arduous and erroneous task [8]. As the rate of publications increases, relying on manual curation alone becomes impractical. Data programming, a paradigm that uses label functions as a means to speed up the annotation process, can be used as a solution for this problem. An obstacle for this paradigm, however, is creating useful label functions, which takes a considerable amount of time. We tested the feasibility of reusing label functions as a way to reduce the total number of label functions required for strong prediction performance. We conclude that label functions may be re-used with closely related edge types, but that re-use does not improve performance for most pairings. The discriminative model's performance improves as more edge-specific label functions are incorporated into the generative model; however, we did notice that performance greatly depends on the annotations provided by the generative model.

This work sets up the foundation for creating a common framework that mines text to create edges. Within this framework we would continuously incorporate new knowledge as novel findings are published, while providing a single confidence score for an edge via sentence score consolidation. As opposed to many existing knowledge graphs (for example, Hetionet v1 where text-derived edges generally cannot be exactly attributed to excerpts from literature [3], [33]), our approach has the potential to annotate each edge based on its source sentences. In addition, edges generated with this approach would be unencumbered from upstream licensing or copyright restrictions, enabling openly licensed hetnets at a scale not previously possible [34]–[36]. New multitask learning [37] strategies may make it even more practical to reuse label functions to construct continuously updating literature-derived knowledge graphs.

Supplemental Information

An online version of this manuscript is available at

https://greenelab.github.io/text mined hetnet manuscript/. Labeled sentences are available at https://github.com/greenelab/text mined hetnet manuscript/tree/master/supplementary materials/annotated sentences. Source code for this work is available under open licenses at: https://github.com/greenelab/snorkeling/.

Acknowledgements

The authors would like to thank Christopher Ré's group at Stanford University, especially Alex Ratner and Steven Bach, for their assistance with this project. We also want to thank Graciela Gonzalez-Hernandez for her advice and input with this project. This work was support by <u>Grant GBMF4552</u> from the Gordon Betty Moore Foundation.

References

- [1] R. Gramatica, T. Di Matteo, S. Giorgetti, M. Barbiani, D. Bevec, and T. Aste, "Graph Theory Enables Drug Repurposing How a Mathematical Model Can Drive the Discovery of Hidden Mechanisms of Action," *PLoS ONE*, vol. 9, no. 1, p. e84912, Jan. 2014 [Online]. Available: https://doi.org/gf45zp. [Accessed: 29-Mar-2022]
- [2] M. Alshahrani and R. Hoehndorf, "Drug repurposing through joint learning on knowledge graphs and literature," Bioinformatics, preprint, Aug. 2018 [Online]. Available: https://doi.org/gf45zk. [Accessed: 29-Mar-2022]
- [3] D. S. Himmelstein *et al.*, "Systematic integration of biomedical knowledge prioritizes drugs for repurposing," *eLife*, vol. 6, p. e26726, Sep. 2017 [Online]. Available: https://doi.org/cdfk. [Accessed: 29-Mar-2022]
- [4] M. Mintz, S. Bills, R. Snow, and D. Jurafsky, "Distant supervision for relation extraction without labeled data," in *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*, Suntec, Singapore, 2009, vol. 2, p. 1003 [Online]. Available: https://doi.org/fg9q43. [Accessed: 29-Mar-2022]
- [5] A. Junge and L. J. Jensen, "CoCoScore: Context-aware co-occurrence scoring for text mining applications using distant supervision," Bioinformatics, preprint, Oct. 2018 [Online]. Available: https://doi.org/gf45zm. [Accessed: 29-Mar-2022]
- [6] H. Zhou, C. Lang, Z. Liu, S. Ning, Y. Lin, and L. Du, "Knowledge-guided convolutional networks for chemical-disease relation extraction," *BMC Bioinformatics*, vol. 20, no. 1, p. 260, Dec. 2019 [Online]. Available: https://doi.org/gf45zn. [Accessed: 29-Mar-2022]
- [7] R. Winnenburg, T. Wachter, C. Plake, A. Doms, and M. Schroeder, "Facts from text: can text mining help to scale-up high-quality manual curation of gene products with ontologies?" *Briefings in Bioinformatics*, vol. 9, no. 6, pp. 466–478, Jul. 2008 [Online]. Available: https://doi.org/bfsnwg. [Accessed: 29-Mar-2022]
- [8] W. A. Baumgartner, K. B. Cohen, L. M. Fox, G. Acquaah-Mensah, and L. Hunter, "Manual curation is not sufficient for annotation of genomic databases," *Bioinformatics*, vol. 23, no. 13, pp. i41–i48, Jul. 2007 [Online]. Available: https://doi.org/dtck86. [Accessed: 29-Mar-2022]
- [9] L. Bornmann and R. Mutz, "Growth rates of modern science: A bibliometric analysis based on the number of publications and cited references: Growth Rates of Modern Science: A Bibliometric Analysis Based on the Number of Publications and Cited References," *Journal of the Association for Information Science and Technology*, vol. 66, no. 11, pp. 2215–2222, Nov. 2015 [Online]. Available: https://doi.org/gfj5zc. [Accessed: 29-Mar-2022]
- [10] A. Ratner, C. De Sa, S. Wu, D. Selsam, and C. Ré, "Data Programming: Creating Large Training Sets, Quickly," arXiv, 1605.07723, Dec. 2018 [Online]. Available: https://arxiv.org/abs/1605.07723
- [11] M. Torii, C. N. Arighi, G. Li, Q. Wang, C. H. Wu, and K. Vijay-Shanker, "RLIMS-P 2.0: A Generalizable Rule-Based Information Extraction System for Literature Mining of Protein Phosphorylation Information," *IEEE/ACM Transactions on Computational Biology and Bioinformatics*, vol. 12, no. 1, pp. 17–29, Jan. 2015 [Online]. Available: https://doi.org/gf8fpv. [Accessed: 29-Mar-2022]

- [12] H.-M. Müller, K. M. Van Auken, Y. Li, and P. W. Sternberg, "Textpresso Central: a customizable platform for searching, text mining, viewing, and curating biomedical literature," *BMC Bioinformatics*, vol. 19, no. 1, p. 94, Dec. 2018 [Online]. Available: https://doi.org/gf7rbz. [Accessed: 29-Mar-2022]
- [13] J. X. Binder *et al.*, "COMPARTMENTS: unification and visualization of protein subcellular localization evidence," *Database*, vol. 2014, no. 0, pp. bau012–bau012, Feb. 2014 [Online]. Available: https://doi.org/btbm. [Accessed: 29-Mar-2022]
- [14] A. Santos, K. Tsafou, C. Stolte, S. Pletscher-Frankild, S. I. O'Donoghue, and L. J. Jensen, "Comprehensive comparison of large-scale tissue expression datasets," *PeerJ*, vol. 3, p. e1054, Jun. 2015 [Online]. Available: https://doi.org/f3mn6p. [Accessed: 29-Mar-2022]
- [15] C. Manning, M. Surdeanu, J. Bauer, J. Finkel, S. Bethard, and D. McClosky, "The Stanford CoreNLP Natural Language Processing Toolkit," in *Proceedings of 52nd Annual Meeting of the Association for Computational Linguistics: System Demonstrations*, Baltimore, Maryland, 2014, pp. 55–60 [Online]. Available: https://doi.org/gf3xhp. [Accessed: 29-Mar-2022]
- [16] L. J. Jensen, J. Saric, and P. Bork, "Literature mining for the biologist: from information retrieval to biological discovery," *Nature Reviews Genetics*, vol. 7, no. 2, pp. 119–129, Feb. 2006 [Online]. Available: https://doi.org/bgq7q9. [Accessed: 29-Mar-2022]
- [17] W. W. M. Fleuren and W. Alkema, "Application of text mining in the biomedical domain," *Methods*, vol. 74, pp. 97–106, Mar. 2015 [Online]. Available: https://doi.org/f64p6n. [Accessed: 29-Mar-2022]
- [18] M. Krallinger, O. Rabal, S. A. Akhondi, et al., "Overview of the biocreative vi chemical-protein interaction track," in *Proceedings of the sixth biocreative challenge evaluation workshop*, 2017, vol. 1, pp. 141–146 [Online]. Available: https://www.semanticscholar.org/paper/Overview-of-the-BioCreative-VI-chemical-protein-Krallinger-Rabal/eed781f498b563df5a9e8a241c67d63dd1d92ad5
- [19] S. Pyysalo, A. Airola, J. Heimonen, J. Björne, F. Ginter, and T. Salakoski, "Comparative analysis of five protein-protein interaction corpora," *BMC Bioinformatics*, vol. 9, no. S3, p. S6, Apr. 2008 [Online]. Available: https://doi.org/fh3df7. [Accessed: 29-Mar-2022]
- [20] M. A. Hearst, S. T. Dumais, E. Osuna, J. Platt, and B. Scholkopf, "Support vector machines," *IEEE Intelligent Systems and their Applications*, vol. 13, no. 4, pp. 18–28, Jul. 1998 [Online]. Available: https://doi.org/fwgxrj. [Accessed: 29-Mar-2022]
- [21] J. Schmidhuber, "Deep learning in neural networks: An overview," *Neural Networks*, vol. 61, pp. 85–117, Jan. 2015 [Online]. Available: https://doi.org/f6v78n. [Accessed: 29-Mar-2022]
- [22] J. MacArthur *et al.*, "The new NHGRI-EBI Catalog of published genome-wide association studies (GWAS Catalog)," *Nucleic Acids Research*, vol. 45, no. D1, pp. D896–D901, Jan. 2017 [Online]. Available: https://doi.org/f9v7cp. [Accessed: 29-Mar-2022]
- [23] T. Rolland *et al.*, "A Proteome-Scale Map of the Human Interactome Network," *Cell*, vol. 159, no. 5, pp. 1212–1226, Nov. 2014 [Online]. Available: https://doi.org/f3mn6x. [Accessed: 29-Mar-2022]
- [24] D. S. Wishart *et al.*, "DrugBank 5.0: a major update to the DrugBank database for 2018," *Nucleic Acids Research*, vol. 46, no. D1, pp. D1074–D1082, Jan. 2018 [Online]. Available: https://doi.org/gcwtzk. [Accessed: 29-Mar-2022]

- [25] C.-H. Wei, A. Allot, R. Leaman, and Z. Lu, "PubTator central: automated concept annotation for biomedical full text articles," *Nucleic Acids Research*, vol. 47, no. W1, pp. W587–W593, Jul. 2019 [Online]. Available: https://doi.org/ggzfsc. [Accessed: 29-Mar-2022]
- [26] R. Leaman and Z. Lu, "TaggerOne: joint named entity recognition and normalization with semi-Markov Models," *Bioinformatics*, vol. 32, no. 18, pp. 2839–2846, Sep. 2016 [Online]. Available: https://doi.org/f855dg. [Accessed: 29-Mar-2022]
- [27] C.-H. Wei, L. Phan, J. Feltz, R. Maiti, T. Hefferon, and Z. Lu, "tmVar 2.0: integrating genomic variant information from literature with dbSNP and ClinVar for precision medicine," *Bioinformatics*, vol. 34, no. 1, pp. 80–87, Jan. 2018 [Online]. Available: https://doi.org/gbzsmc. [Accessed: 29-Mar-2022]
- [28] C.-H. Wei, H.-Y. Kao, and Z. Lu, "GNormPlus: An Integrative Approach for Tagging Genes, Gene Families, and Protein Domains," *BioMed Research International*, vol. 2015, pp. 1–7, 2015 [Online]. Available: https://doi.org/gb85jb. [Accessed: 29-Mar-2022]
- [29] C.-H. Wei, H.-Y. Kao, and Z. Lu, "SR4GN: A Species Recognition Software Tool for Gene Normalization," *PLoS ONE*, vol. 7, no. 6, p. e38460, Jun. 2012 [Online]. Available: https://doi.org/gpq498. [Accessed: 29-Mar-2022]
- [30] M. Honnibal and I. Montani, "spaCy 2: Natural language understanding with Bloom embeddings, convolutional neural networks and incremental parsing," 2017.
- [31] S. Razick, G. Magklaras, and I. M. Donaldson, "iRefIndex: A consolidated protein interaction database with provenance," *BMC Bioinformatics*, vol. 9, no. 1, p. 405, Dec. 2008 [Online]. Available: https://doi.org/b99bjj. [Accessed: 29-Mar-2022]
- [32] J. Menche *et al.*, "Uncovering disease-disease relationships through the incomplete interactome," *Science*, vol. 347, no. 6224, pp. 1257601–1257601, Feb. 2015 [Online]. Available: https://doi.org/f3mn6z. [Accessed: 29-Mar-2022]
- [33] "[No title found]." [Online]. Available: https://doi.org/f3mqwp
- [34] "[No title found]." [Online]. Available: https://doi.org/bfmk
- [35] S. Oxenham, "Legal confusion threatens to slow data science," *Nature*, vol. 536, no. 7614, pp. 16–17, Aug. 2016 [Online]. Available: https://doi.org/bndt. [Accessed: 29-Mar-2022]
- [36] S. Carbon, R. Champieux, J. A. McMurry, L. Winfree, L. R. Wyatt, and M. A. Haendel, "An analysis and metric of reusable data licensing practices for biomedical resources," *PLOS ONE*, vol. 14, no. 3, p. e0213090, Mar. 2019 [Online]. Available: https://doi.org/gf5m8v. [Accessed: 29-Mar-2022]
- [37] A. Ratner, B. Hancock, J. Dunnmon, R. Goldman, and C. Ré, "Snorkel MeTaL: Weak Supervision for Multi-Task Learning," in *Proceedings of the Second Workshop on Data Management for End-To-End Machine Learning*, Houston TX USA, 2018, pp. 1–4 [Online]. Available: https://doi.org/gf3xk7. [Accessed: 29-Mar-2022]
- [38] A. Ratner, S. H. Bach, H. Ehrenberg, J. Fries, S. Wu, and C. Ré, "Snorkel: rapid training data creation with weak supervision," *Proceedings of the VLDB Endowment*, vol. 11, no. 3, pp. 269–282, Nov. 2017 [Online]. Available: https://doi.org/ch44. [Accessed: 29-Mar-2022]
- [39] Y. Zhang and B. Wallace, "A Sensitivity Analysis of (and Practitioners' Guide to) Convolutional Neural Networks for Sentence Classification," arXiv, 1510.03820, Apr. 2016 [Online]. Available: https://arxiv.org/abs/1510.03820

- [40] D. P. Kingma and J. Ba, "Adam: A Method for Stochastic Optimization," arXiv, 1412.6980, Jan. 2017 [Online]. Available: https://arxiv.org/abs/1412.6980
- [41] T. Mikolov, I. Sutskever, K. Chen, G. Corrado, and J. Dean, "Distributed Representations of Words and Phrases and their Compositionality," arXiv, 1310.4546, Oct. 2013 [Online]. Available: https://arxiv.org/abs/1310.4546
- [42] P. Bojanowski, E. Grave, A. Joulin, and T. Mikolov, "Enriching Word Vectors with Subword Information," arXiv, 1607.04606, Jun. 2017 [Online]. Available: https://arxiv.org/abs/1607.04606
- [43] T. Mikolov, K. Chen, G. Corrado, and J. Dean, "Efficient Estimation of Word Representations in Vector Space," arXiv, 1301.3781, Sep. 2013 [Online]. Available: https://arxiv.org/abs/1301.3781
- [44] C. Guo, G. Pleiss, Y. Sun, and K. Q. Weinberger, "On Calibration of Modern Neural Networks," arXiv, 1706.04599, Aug. 2017 [Online]. Available: https://arxiv.org/abs/1706.04599
- [45] V. Kuleshov, N. Fenner, and S. Ermon, "Accurate Uncertainties for Deep Learning Using Calibrated Regression," arXiv, 1807.00263, Jul. 2018 [Online]. Available: https://arxiv.org/abs/1807.00263

Supplemental Methods

Training Models

Generative Model

The generative model is a core part of this automatic annotation framework. It integrates multiple signals emitted by label functions and assigns a training class to each candidate sentence. This model assigns training classes by estimating the joint probability distribution of the latent true class (Y) and label function signals (Λ) , $(P_{\theta}(\Lambda,Y))$. Assuming each label function is conditionally independent, the joint distribution is defined as follows:

$$P_{ heta}(\Lambda,Y) = rac{\exp(\sum_{i=1}^m heta^T F_i(\Lambda,y))}{\sum_{\Lambda'} \sum_{y'} \exp(\sum_{i=1}^m heta^T F_i(\Lambda',y'))}$$

where m is the number of candidate sentences, F is the vector of summary statistics and θ is a vector of weights for each summary statistic. The summary statistics used by the generative model are as follows:

$$egin{aligned} F_{i,j}^{Lab}(\Lambda,Y) &= \mathbb{1}\{\Lambda_{i,j}
eq 0\} \ F_{i,j}^{Acc}(\Lambda,Y) &= \mathbb{1}\{\Lambda_{i,j} = y_{i,j}\} \end{aligned}$$

Lab is the label function's propensity (the frequency of a label function emitting a signal). Acc is the individual label function's accuracy given the training class. This model optimizes the weights (θ) by minimizing the negative log likelihood:

$$\hat{ heta} = argmin_{ heta} - \sum_{\Lambda} \sum_{Y} log P_{ heta}(\Lambda, Y)$$

In the framework we used predictions from the generative model, $\hat{Y}=P_{\hat{\theta}}(Y\mid\Lambda)$, as training classes for our dataset [37], [38].

Discriminative Model

The discriminative model is a neural network trained to produce classification labels by integrating predicted probabilities from the generative model along with sentence representations via word embeddings. The goal of this combined approach is to develop models that learn text features associated with the overall task, beyond the supplied label functions. We used a piecewise convolutional neural network that contains multiple kernel filters as our discriminative model. We built a network with multiple filters using a fixed width of 300 (size of word embeddings) and a fixed height of 7 (Figure 4). We chose a fixed height of 7 because this height was previously reported to optimize performance in relationship classification [39]. We trained this model for 15 epochs using the Adam optimizer [40] with PyTorch's default parameter settings and a learning rate of 0.001 that decreases by half every epoch until the lower bound of 1e-5 is reached, which we observed was often sufficient for convergence. We added a L2 penalty (lambda=0.002) on the network weights to prevent overfitting. Lastly, we added a dropout layer (p=0.25) between the fully connected layer and the softmax layer.

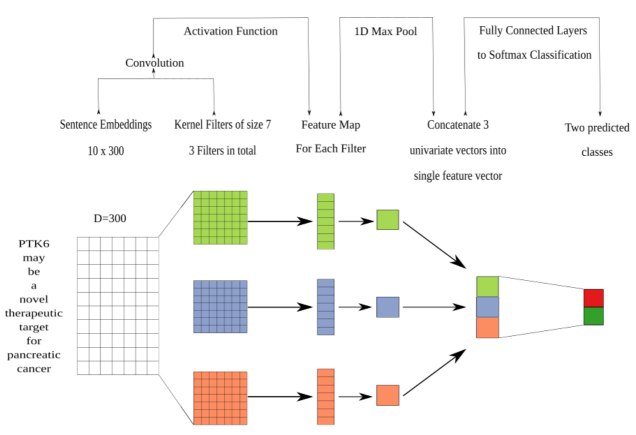


Figure 4: The architecture of the discriminative model was a convolutional neural network. We performed a convolution step using multiple filters. The filters generated a feature map that was sent into a maximum pooling layer that was designed to extract the largest feature in each map. The extracted features were concatenated into a singular vector that was passed into a fully connected network. The fully connected network had 300 neurons for the first layer, 100 neurons for the second layer and 50 neurons for the last layer. The last step of the fully connected network was to generate predictions using a softmax layer.

Word Embeddings

Word embeddings are representations that map individual words to real valued vectors of user-specified dimensions. These embeddings have been shown to capture the semantic and syntactic information between words [41]. We trained Facebook's fastText [42] using all candidate sentences for each individual relationship pair to generate word embeddings. FastText uses a skip-gram model [43] that aims to predict the surrounding context for a candidate word and pairs the model with a novel scoring function that treats each word as a bag of character n-grams. We trained this model for 20

epochs using a window size of 2 and generated 300-dimensional word embeddings. We use the optimized word embeddings as input to our discriminative model.

Calibration of the Discriminative Model

Often many tasks require a machine learning model to output reliable probability predictions. A model is well calibrated if the probabilities emitted from the model match the observed probabilities. For example, a well-calibrated model that assigns a class label with 80% probability should have that class appear 80% of the time. Deep neural network models can often be poorly calibrated [44], [45]. These models are usually over-confident in their predictions. For this reason, we calibrated our convolutional neural network using temperature scaling [44]. Temperature scaling uses a parameter T to scale each value of the logit vector (z) before being passed into the softmax (SM) function.

$$\sigma_{SM}(rac{z_i}{T}) = rac{\exp(rac{z_i}{T})}{\sum_i \exp(rac{z_i}{T})}$$

We found the optimal T by minimizing the negative log likelihood (NLL) of the tune set.

Supplemental Figures

Generative Model Using Randomly Sampled Label Functions

Individual Sources

Test Set AUPR of Predicted Relations

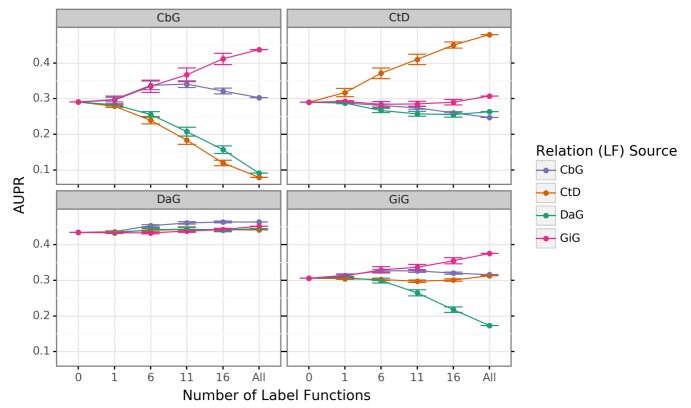


Figure 5: Edge-specific label functions improves performance over edge-mismatch label functions. Each line plot header depicts the edge type the generative model is trying to predict, while the colors represent the source of label functions. For example orange represents sampling label functions designed to predict the Compound treats Disease (CtD) edge type. The x axis shows the number of randomly sampled label functions being incorporated into the

database-only baseline model (point at 0). The y axis shows area under the precision recall curve (AUPR). Each point on the plot shows the average of 50 sample runs, while the error bars show the 95% confidence intervals of all runs. The baseline and "All" data points consist of sampling from the entire fixed set of label functions.

Collective Pool of Sources

Test Set AUROC of Predicted Relations

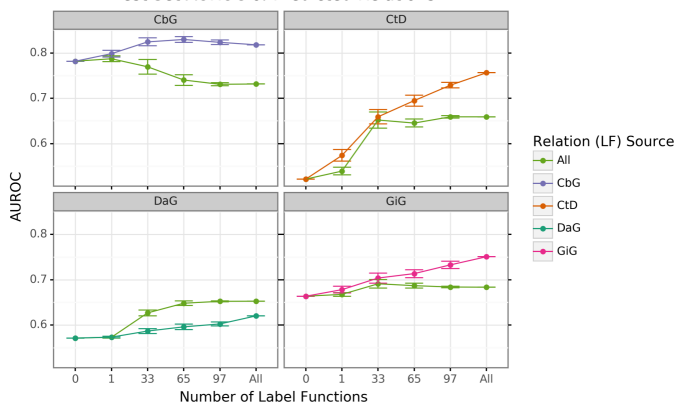


Figure 6: Using all label functions generally hinders generative model performance. Each line plot header depicts the edge type the generative model is trying to predict, while the colors represent the source of label functions. For example, orange represents sampling label functions designed to predict the Compound treats Disease (CtD) edge type. The x axis shows the number of randomly sampled label functions being incorporated into the database-only baseline model (point at 0). The y axis shows area under the receiver operating curve (AUROC). Each point on the plot shows the average of 50 sample runs, while the error bars show the 95% confidence intervals of all runs. The baseline and "All" data points consist of sampling from the entire fixed set of label functions.

Test Set AUPR of Predicted Relations

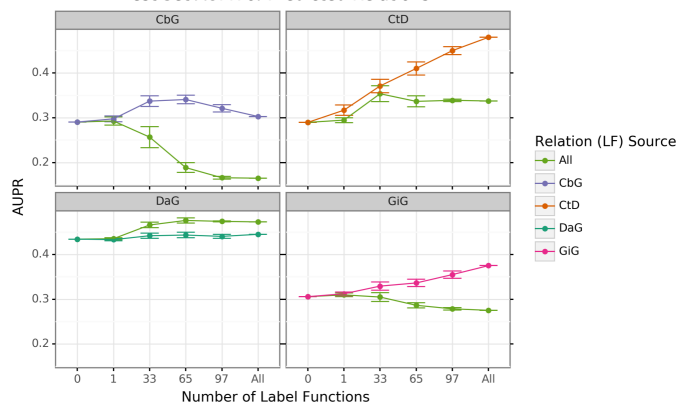


Figure 7: Using all label functions generally hinders generative model performance. Each line plot header depicts the edge type the generative model is trying to predict, while the colors represent the source of label functions. For example, orange represents sampling label functions designed to predict the Compound treats Disease (CtD) edge type. The x axis shows the number of randomly sampled label functions being incorporated into the database-only baseline model (point at 0). The y axis shows area under the precision recall curve (AUPR). Each point on the plot shows the average of 50 sample runs, while the error bars show the 95% confidence intervals of all runs. The baseline and "All" data points consist of sampling from the entire fixed set of label functions.

Discriminative Model Performance

Discriminative Model Performance (Test Set)

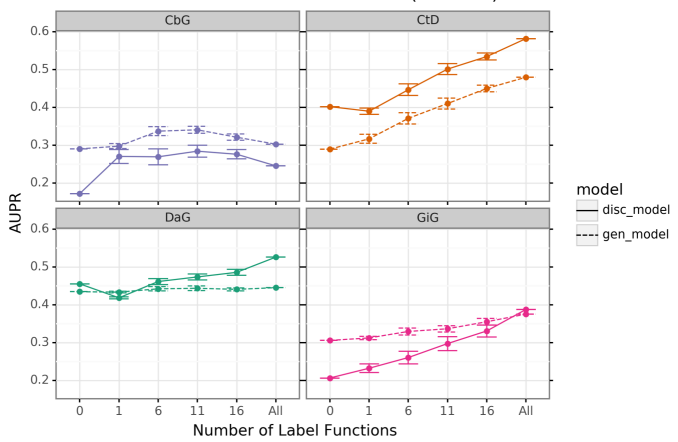


Figure 8: The discriminator model improves performance as the number of edge-specific label functions is added to the baseline model. The line plot headers represents the specific edge type the discriminator model is trying to predict. The x-axis shows the number of randomly sampled label functions incorporated on top of the baseline model (point at 0). The y axis shows the area under the precision recall curve (AUPR). Each datapoint shows the average of 50 sample runs, while the error bars represents the 95% confidence interval at each point. The baseline and "All" data points consist of sampling from the entire fixed set of label functions.

Discriminative Model Calibration

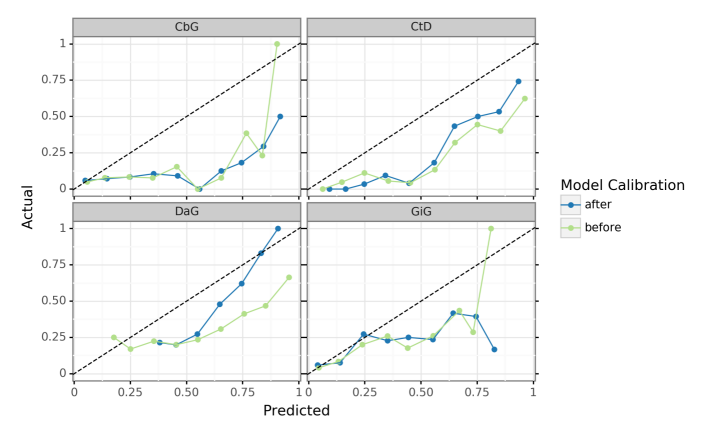


Figure 9: Deep learning models are overconfident in their predictions and need to be calibrated after training. These are calibration plots for the discriminative model, where the green line represents the predictions before calibration and the blue line shows predictions after calibration. Data points that lie closer to the diagonal line show better model calibration, while data points far from the diagonal line show poor performance. A perfectly calibrated model would align straight along the diagonal line.

Even deep learning models with impressive AUROC and AUPR statistics can be subject to poor calibration. Typically, these models are overconfident in their predictions [44], [45]. We attempted to use temperature scaling to fix the calibration of the best performing discriminative models (Figure 9). Before calibration (green lines), our models were aligned with the ideal calibration only when predicting low probability scores (close to 0.25). Applying the temperature scaling calibration algorithm (blue lines) did not substantially improve the calibration of the model in most cases. The exception to this pattern is the Disease associates Gene (DaG) model where high confidence scores are shown to be better calibrated. Overall, calbrating deep learning models is a nontrivial task that requires more complex approaches to accomplish.

Text Mined Edges Can Expand a Database-derived Knowledge Graph

Reconstructing Edges in Hetionet

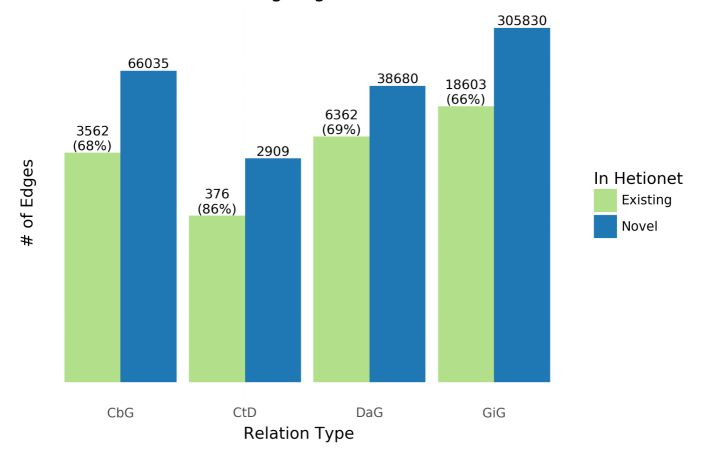


Figure 10: Text-mined edges recreate a substantial fraction of an existing knowledge graph and include new predictions. This bar chart shows the number of edges we can successfully recall in green and shows the number of new edges that can be added in blue.

The recall for the Hetionet v1 knowledge graph is shown as a percentage in parentheses. For example, for the Compound treats Disease (CtD) edge our method recalls 85% of existing edges and adds 6,088 new edges.

One of the goals in our work is to measure the extent to which learning multiple edge types could construct a biomedical knowledge graph. Using Hetionet v1 as an evaluation set, we measured this framework's recall and quantified how many new edges could be added with high confidence. Overall, we were able to recall more than half of preexisting edges for all edge types (Figure 10) and report our top ten scoring sentences for each edge type in Supplemental Table 11. Our best recall is with the Compound treats Disease (CtD) edge type, where we retain 85% of preexisting edges. Plus, we can add over 6,000 new edges to that category. In contrast, we could only recall close to 70% of existing edges for the other categories; however, we can add over 40,000 novel edges to each category. This highlights the fact that Hetionet v1 is missing a compelling amount of biomedical information and this framework is a viable way to close the information gap.

Comparison with CoCoScore using Hetionet v1 as an Evaluation Set

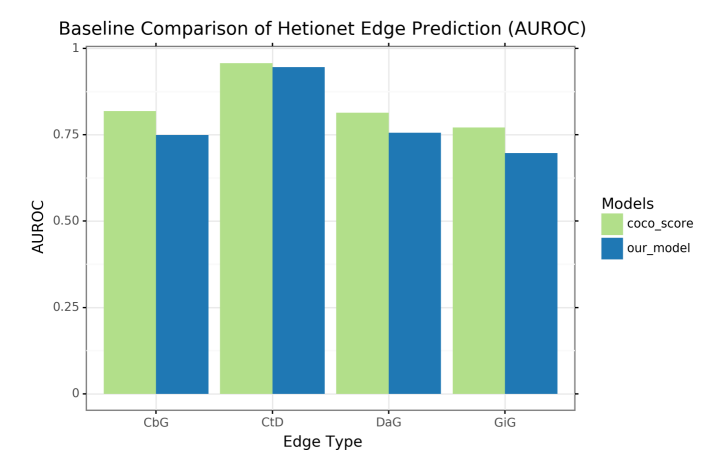


Figure 11: Our extractor shows similar performance to a previously published method when using Hetionet v1 as an evaluation set. We compared our model (blue) with the CoCoScore model <u>tag:cocoscore?</u> (green). The y axis represents AUROC and the x axis represents the edge type both models are trying to predict.

Our model showed promising performance in terms of recalling edges in Hetionet v1. We assessed our model's performance relative to a recently published method tag:cocoscore?. Though our method is primarily designed to predict assertions, not edges, we compared performance at an edge level because this was available for CoCoScore. We found that a simple summary approach, max sentence score, provided comparable performance to the CoCoScore for the compound treats disease (CtD) edge type and slightly poorer performance for other edge types (Supplemental Figure 11). Sentence-level scores can be integrated in multiple ways, and approaches that consider more complexity (e.g., the number of sentences with high-probability) should be evaluated in future work.

Supplemental Tables

Discriminative Model Calibration Tables

Table 3: Contains the top ten Disease-associates-Gene confidence scores before and after model calbration. Disease mentions are highlighted in brown and Gene mentions are highlighted in blue.

| Disease Name | Gene Symbol | Text | Before Calibration | After Calibration |
|-----------------|----------------|---|-----------------------|----------------------|
| prostate cancer | DKK1 | conclusion: high dkk-1 serum levels are associated with a poor survival in patients with prostate cancer. | 0.999 | 0.916 |
| breast cancer | ERBB2 | conclusion: her-2 / neu overexpression in primary breast carcinoma is correlated with patients ' age (under age 50) and calcifications at mammography . | 0.998 | 0.906 |

| Disease Name | Gene Symbol | Text | Before Calibration | After Calibration |
|---------------------------|----------------|--|-----------------------|----------------------|
| breast cancer | ERBB2 | the results of multiple linear regression analysis , with her2 as the dependent variable , showed that family history of breast cancer was significantly associated with elevated her2 levels in the tumors (p = 0.0038) , after controlling for the effects of age , tumor estrogen receptor , and dna index . | 0.998 | 0.904 |
| colon cancer | SP3 | ba also decreased expression of sp1 , sp3 and sp4 transcription factors which are overexpressed in colon cancer cells and decreased levels of several sp-regulated genes including survivin , vascular endothelial growth factor , p65 sub-unit of nfkb , epidermal growth factor receptor , cyclin d1 , and pituitary tumor transforming gene-1 . | 0.998 | 0.902 |
| breast cancer | ERBB2 | in breast cancer , overexpression of her2 is associated with an aggressive tumor phenotype and poor prognosis . | 0.998 | 0.898 |
| breast cancer | BCL2 | in clinical breast cancer samples , high bcl2 expression was associated with poor prognosis . | 0.997 | 0.886 |
| adrenal gland cancer | TP53 | the mechanisms of adrenal tumorigenesis remain poorly established; the r337h germline mutation in the p53 gene has previously been associated with acts in brazilian children. | 0.996 | 0.883 |
| prostate cancer | AR | the androgen receptor was expressed in all primary and metastatic prostate cancer tissues and no mutations were identified . | 0.996 | 0.881 |
| urinary bladder cancer | PIK3CA | conclusions: increased levels of fgfr3 and pik3ca mutated dna in urine and plasma are indicative of later progression and metastasis in bladder cancer. | 0.995 | 0.866 |
| ovarian cancer | EPAS1 | the log-rank test showed that nuclear positive immunostaining for hif-1alpha ($p = .002$) and cytoplasmic positive immunostaining for hif-2alpha ($p = .0112$) in tumor cells are associated with poor prognosis of patients with ovarian carcinoma . | 0.994 | 0.86 |

Table 4: Contains the bottom ten Disease-associates-Gene confidence scores before and after model calbration. Disease mentions are highlighted in brown and Gene mentions are highlighted in blue.

| Disease Name | Gene Symbol | Text | Before Calibration | After Calibration |
|-----------------------|----------------|---|-----------------------|----------------------|
| endogenous depression | EP300 | from a clinical point of view , p300 amplitude should be considered as a psychophysiological index of suicidal risk in major depressive disorder . | 0.202 | 0.379 |
| Alzheimer's disease | PDK1 | from prion diseases to alzheimer 's disease : a common therapeutic target , [pdk1]. | 0.2 | 0.378 |

| Disease Name | Gene Symbol | Text | Before Calibration | After Calibration |
|--------------------------------|----------------|---|-----------------------|----------------------|
| endogenous depression | HTR1A | gepirone , a selective serotonin (5ht1a) partial agonist in the treatment of major depression . | 0.199 | 0.378 |
| Gilles de la Tourette syndrome | FGF9 | there were no differences in gender distribution , age at tic onset or td diagnosis , tic severity , proportion with current diagnoses of ocd/oc behavior or attention deficit hyperactivity disorder (adhd) , cbcl internalizing , externalizing , or total problems scores , ygtss scores , or gaf scores . | 0.185 | 0.37 |
| hematologic cancer | MLANA | methods: the sln sections (n = 214) were assessed by qrt assay for 4 established messenger rna biomarkers: mart-1, mage-a3, galnac-t, and pax3. | 0.18 | 0.368 |
| endogenous depression | MAOA | alpha 2-adrenoceptor responsivity in depression : effect of chronic treatment with moclobemide , a selective mao- a-inhibitor , versus maprotiline . | 0.179 | 0.367 |
| chronic kidney failure | B2M | to evaluate comparative beta 2-m removal we studied six stable end-stage renal failure patients during high-flux 3-h haemodialysis, haemodiafiltration, and haemofiltration, using acrylonitrile, cellulose triacetate, polyamide and polysulphone capillary devices. | 0.178 | 0.366 |
| hematologic cancer | C7 | serum antibody responses to four haemophilus influenzae type b capsular polysaccharide-protein conjugate vaccines (prp-d, hboc, c7p, and prp-t) were studied and compared in 175 infants, 85 adults and 140 2-year-old children. | 0.174 | 0.364 |
| hypertension | AVP | portohepatic pressures , hepatic function , and blood gases in the combination of nitroglycerin and vasopressin : search for additive effects in cirrhotic portal hypertension . | 0.168 | 0.361 |
| endogenous depression | GAD1 | within-individual deflections in gad , physical , and social symptoms predicted later deflections in depressive symptoms , and deflections in depressive symptoms predicted later deflections in gad and separation anxiety symptoms . | 0.149 | 0.349 |

Table 5: Contains the top ten Compound-treats-Disease confidence scores after model calbration. Disease mentions are highlighted in brown and Compound mentions are highlighted in red.

| Compound Name | Disease Name | Text | Before Calibration | After Calibration |
|------------------------|-----------------------|---|-----------------------|----------------------|
| Prazosin | hypertension | experience with prazosin in the treatment of hypertension . | 0.997 | 0.961 |
| Methyldopa | hypertension | oxprenolol plus cyclopenthiazide-kcl versus methyldopa in the treatment of hypertension . | 0.997 | 0.961 |
| Methyldopa | hypertension | atenolol and methyldopa in the treatment of hypertension . | 0.996 | 0.957 |
| Prednisone | asthma | prednisone and beclomethasone for treatment of asthma . | 0.995 | 0.953 |
| Sulfasalazine | ulcerative colitis | sulphasalazine, used in the treatment of ulcerative colitis, is cleaved in the colon by the metabolic action of colonic bacteria on the diazo bond to release 5-aminosalicylic acid (5-asa) and sulpharidine. | 0.994 | 0.949 |
| Prazosin | hypertension | letter : prazosin in treatment of hypertension . | 0.994 | 0.949 |
| Methylprednis olone | asthma | use of tao without methylprednisolone in the treatment of severe asthma . | 0.994 | 0.948 |
| Budesonide | asthma | thus , a regimen of budesonide treatment that consistently attenuates bronchial responsiveness in asthmatic subjects had no effect in these men; larger and longer trials will be required to establish whether a subgroup of smokers shows a favorable response . | 0.994 | 0.946 |
| Methyldopa | hypertension | pressor and chronotropic responses to bilateral carotid occlusion (bco) and tyramine were also markedly reduced following treatment with methyldopa , which is consistent with the clinical findings that chronic methyldopa treatment in hypertensive patients impairs cardiovascular reflexes . | 0.994 | 0.946 |
| Fluphenazine | schizophrenia | low dose fluphenazine decanoate in maintenance treatment of schizophrenia . | 0.994 | 0.946 |

Table 6: Contains the bottom ten Compound-treats-Disease confidence scores before and after model calbration. Disease mentions are highlighted in brown and Compound mentions are highlighted in red.

| Compound Name | Disease Name | Text | Before Calibration | After Calibration |
|------------------|----------------|--|-----------------------|----------------------|
| Indomethaci n | hypertension | effects of indomethacin in rabbit renovascular hypertension . | 0.033 | 0.13 |
| Alprazolam | panic disorder | according to logistic regression analysis, the relationships between plasma alprazolam concentration and response, as reflected by number of panic attacks reported, phobia ratings, physicians, and patients, ratings of global improvement, and the emergence of side effects, were significant. | 0.03 | 0.124 |

| Compound Name | Disease Name | Text | Before Calibration | After Calibration |
|--------------------|------------------------------|---|-----------------------|----------------------|
| Mestranol | polycystic ovary syndrome | the binding capacity of plasma testosterone-estradiol-binding globulin (tebg) and testosterone (t) levels were measured in four women with proved polycystic ovaries and three women with a clinical diagnosis of polycystic ovarian disease before , during , and after administration of norethindrone , 2 mg. , and mestranol , 0.1 mg . | 0.03 | 0.123 |
| Creatine | coronary artery disease | during successful and uncomplicated angioplasty (ptca), we studied the effect of a short lasting myocardial ischemia on plasma creatine kinase, creatine kinase mb-activity, and creatine kinase mmisoforms (mm1, mm2, mm3) in 23 patients. | 0.028 | 0.12 |
| Creatine | coronary artery disease | in 141 patients with acute myocardial infarction, creatine phosphokinase isoenzyme (cpk-mb) was determined by the activation method with dithiothreitol (rao et al.: clin. | 0.027 | 0.117 |
| Morphine | brain cancer | the tissue to serum ratio of morphine in the hypothalamus , hippocampus , striatum , midbrain and cortex were also smaller in morphine tolerant than in non-tolerant rats . | 0.026 | 0.115 |
| Glutathione | anemia | our results suggest that an association between gsh px deficiency and hemolytic anemia need not represent a cause-and-effect relationship. | 0.026 | 0.114 |
| Dinoproston e | stomach cancer | prostaglandin e2 (pge2) - and 6-keto-pgf1 alpha-like immunoactivity was measured in incubates of forestomach and gastric corpus mucosa in (a) unoperated rats , (b) rats with shamoperation of the kidneys and (c) rats with bilateral nephrectomy . | 0.023 | 0.107 |
| Creatine | coronary artery disease | the value of the electrocardiogram in assessing infarct size was studied using serial estimates of the mb isomer of creatine kinase (ck mb) in plasma , serial 35 lead praecordial maps in 28 patients with anterior myocardial infarction , and serial 12 lead electrocardiograms in 17 patients with inferior myocardial infarction . | 0.022 | 0.105 |
| Sulfamethaz ine | multiple sclerosis | quantitation and confirmation of sulfamethazine residues in swine muscle and liver by lc and gc/ms . | 0.017 | 0.093 |

Table 7: Contains the top ten Compound-binds-Gene confidence scores before and after model calbration. Gene mentions are highlighted in blue and Compound mentions are highlighted in red.

| Compound Name | Gene Symbol | Text | Before Calibration | After Calibration |
|---------------|----------------|------|--------------------|-------------------|
|---------------|----------------|------|--------------------|-------------------|

| Compound Name | Gene Symbol | Text | Before Calibration | After Calibration |
|-----------------------------------|----------------|--|--------------------|-------------------|
| Cyclic Adenosine Monophosphate | B3GNT2 | in sk-n-mc human neuroblastoma cells , the camp response to 10 nm isoproterenol (iso) is mediated primarily by beta 1-adrenergic receptors . | 0.903 | 0.93 |
| Indomethacin | AGT | indomethacin, a potent inhibitor of prostaglandin synthesis, is known to increase the maternal blood pressure response to angiotensin ii infusion. | 0.894 | 0.922 |
| Tretinoin | RXRA | the vitamin a derivative retinoic acid exerts its effects on transcription through two distinct classes of nuclear receptors , the retinoic acid receptor (rar) and the retinoid x receptor (rxr) . | 0.882 | 0.912 |
| Tretinoin | RXRA | the vitamin a derivative retinoic acid exerts its effects on transcription through two distinct classes of nuclear receptors , the retinoic acid receptor (rar) and the retinoid x receptor (rxr) . | 0.872 | 0.903 |
| D-Tyrosine | CSF1 | however, the extent of gap tyrosine phosphorylation induced by csf-1 was approximately 10 % of that induced by pdgf-bb in the nih3t3 fibroblasts. | 0.851 | 0.883 |
| D-Glutamic Acid | GLB1 | thus, the negatively charged side chain of glu-461 is important for divalent cation binding to beta-galactosidase. | 0.849 | 0.882 |
| D-Tyrosine | CD4 | second , we use the same system to provide evidence that the physical association of cd4 with the tcr is required for effective tyrosine phosphorylation of the tcr zeta-chain subunit , presumably reflecting delivery of p56lck (lck) to the tcr . | 0.825 | 0.859 |

| Compound Name | Gene Symbol | Text | Before Calibration | After Calibration |
|------------------|----------------|---|--------------------|-------------------|
| Calcium Chloride | TNC | the possibility that the enhanced length dependence of ca2 + sensitivity after cardiac tnc reconstitution was attributable to reduced tnc binding was excluded when the length dependence of partially extracted fast fibres was reduced to one-half the normal value after a 50 % deletion of the native tnc . | 0.821 | 0.855 |
| Metoprolol | KCNMB2 | studies in difi cells of the displacement of specific 125i-cyp binding by nonselective (propranolol), beta 1-selective (metoprolol and atenolol), and beta 2-selective (ici 118-551) antagonists revealed only a single class of beta 2-adrenergic receptors. | 0.82 | 0.854 |
| D-Tyrosine | PLCG1 | epidermal growth factor (egf) or platelet-derived growth factor binding to their receptor on fibroblasts induces tyrosine phosphorylation of plc gamma 1 and stable association of plc gamma 1 with the receptor protein tyrosine kinase. | 0.818 | 0.851 |

Table 8: Contains the bottom ten Compound-binds-Gene confidence scores before and after model calbration. Gene mentions are highlighted in blue and Compound mentions are highlighted in red.

| Compound Name | · Iext | | Before Calibration | After Calibration |
|--------------------|--------|---|-----------------------|----------------------|
| Deferoxamin e | TF | the mechanisms of fe uptake have been characterised using 59fe complexes of citrate , nitrilotriacetate , desferrioxamine , and 59fe added to eagle 's minimum essential medium (mem) and compared with human transferrin (tf) labelled with 59fe and iodine-125 . | 0.02 | 0.011 |
| Hydrocortiso ne | GH1 | group iv patients had normal basal levels of lh and normal lh , gh and cortisol responses . | 0.02 | 0.011 |
| Carbachol | INS | at the same concentration , however , iapp significantly (p less than 0.05) inhibited carbachol-stimulated (10 (-7) m) release of insulin by 30 %, and cgrp significantly inhibited carbachol-stimulated release of insulin by 33 % when compared with the control group. | 0.02 | 0.011 |

| Compound Name | Gene Symbol | Text | Before Calibration | After Calibration |
|-------------------|--|---|-----------------------|----------------------|
| Adenosine | ME2 | at physiological concentrations , atp , adp , and amp all inhibit the enzyme from atriplex spongiosa and panicum miliaceum (nad-me-type plants) , with atp the most inhibitory species . | 0.019 | 0.01 |
| Naloxone | POMC specifically, opioids, including 2-n-pentyloxy-2-phenyl-4-methyl-morpholine, naloxone, and beta-endorphin, have been shown to interact with il-2 receptors (134) and regulate production of il-1 and il-2 (48-50, 135). | | 0.018 | 0.01 |
| Cortisone acetate | POMC | sarcoidosis therapy with cortisone and acth – the role of acth therapy . | 0.017 | 0.009 |
| Epinephrine | INS | thermogenic effect of thyroid hormones : interactions with epinephrine and insulin . | 0.017 | 0.009 |
| Aldosterone | KNG1 | important vasoconstrictor , fluid - and sodium-retaining factors are the renin-angiotensin-aldosterone system , sympathetic nerve activity , and vasopressin ; vasodilator , volume , and sodium-eliminating factors are atrial natriuretic peptide , vasodilator prostaglandins like prostacyclin and prostaglandin e2 , dopamine , bradykinin , and possibly , endothelial derived relaxing factor (edrf) . | 0.016 | 0.008 |
| D-Leucine | POMC | cross-reactivities of leucine-enkephalin and beta- endorphin with the eia were less than 0.1 %, while that with gly-gly-phe-met and oxidized gly-gly-phe-met were 2.5 % and 10.2 %, respectively. | 0.011 | 0.005 |
| Estriol | LGALS1 | [diagnostic value of serial determination of estriol and hpl in plasma and of total estrogens in 24-h-urine compared to single values for diagnosis of fetal danger] . | 0.01 | 0.005 |

Table 9: Contains the top ten Gene-interacts-Gene confidence scores before and after model calbration. Both gene mentions highlighted in blue.

| Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration |
|--------------|--------------|--|--------------------|-------------------|
| ESR1 | HSP90AA1 | previous studies have suggested that the 90- kda heat shock protein (hsp90) interacts with the er, thus stabilizing the receptor in an inactive state . | 0.812 | 0.864 |

| Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration |
|--------------|--------------|--|--------------------|-------------------|
| TP53 | TP73 | cyclin g interacts with p53 as well as p73, and its binding to p53 or p73 presumably mediates downregulat ion of p53 and p73. | 0.785 | 0.837 |
| TP53 | AKT1 | treatment of c81 cells with ly294002 resulted in an increase in the p53-responsive gene mdm2, suggesting a role for akt in the tax-mediated regulation of p53 transcriptio nal activity. | 0.773 | 0.825 |
| ABCB1 | NR1I3 | valproic acid induces cyp3a4 and mdr1 gene expression by activation of constitutive androstane receptor and pregnane x receptor pathways. | 0.762 | 0.813 |
| PTH2R | PTH2 | thus, the juxtamembr ane receptor domain specifies the signaling and binding selectivity of tip39 for the pth2 receptor over the pth1 receptor. | 0.761 | 0.812 |

| Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration |
|--------------|--------------|--|--------------------|-------------------|
| CCND1 | ABL1 | synergy with v-abl depended on a motif in cyclin d1 that mediates its binding to the retinoblasto ma protein , suggesting that abl oncogenes in part mediate their mitogenic effects via a retinoblasto ma protein-dependent pathway . | 0.757 | 0.808 |
| CTNND1 | CDH1 | these complexes are formed independen tly of ddr1 activation and of betacatenin and p120-catenin binding to ecadherin; they are ubiquitous in epithelial cells. | 0.748 | 0.798 |
| CSF1 | CSF1R | this is in agreement with current thought that the c-fms proto-oncogene product functions as the csf-1 receptor specific to this pathway . | 0.745 | 0.795 |

| Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration |
|--------------|--------------|--|--------------------|-------------------|
| EZR | CFTR | without ezrin binding, the cytoplasmic tail of cftr only interacts strongly with the first amino- terminal pdz domain to form a 1:1 c- cftr. | 0.732 | 0.78 |
| SRC | PIK3CG | we have demonstrat ed that the sh2 (src homology 2) domains of the 85 kda subunit of pi-3k are sufficient to mediate binding of the pi-3k complex to tyrosine phosphoryla ted , but not nonphosphoryla ted il-2r beta , suggesting that tyrosine phosphoryla tion is an integral component of the activation of pi-3k by the il-2r . | 0.731 | 0.78 |

Table 10: Contains the bottom ten Gene-interacts-Gene confidence scores before and after model calbration. Both gene mentions highlighted in blue.

| | Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration | |
|--|--------------|--------------|------|--------------------|-------------------|--|
|--|--------------|--------------|------|--------------------|-------------------|--|

| Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration |
|--------------|--------------|---|--------------------|-------------------|
| AGTR1 | ACE | result (s): the luteal tissue is the major site of ang ii, ace, at1r, and vegf, with highest staining intensity found during the midluteal phase and at pregnancy. | 0.009 | 0.003 |
| ABCE1 | ABCF2 | in relation to normal melanocytes , abcb3 , abcb6 , abcc2 , abcc4 , abce1 and abcf2 were significantly increased in melanoma cell lines , whereas abca7 , abca12 , abcb4 , abcb5 and abcd1 showed lower expression levels . | 0.008 | 0.002 |
| IL4 | IFNG | in contrast , il-13ralpha2 mrna expression was up- regulated by ifn-gamma plus il-4 . | 0.007 | 0.002 |
| FCAR | CD79A | we report here the presence of circulating soluble fcalphar (cd89) - iga complexes in patients with igan . | 0.007 | 0.002 |

| Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration |
|--------------|--------------|--|--------------------|-------------------|
| IL4 | VCAM1 | similarly , il-4 induced vcam-1 expression and augmented tnf-alpha-induced expression on huvec but did not affect vcam-1 expression on hdmec . | 0.007 | 0.002 |
| IL2 | IFNG | prostaglandi n e2 at priming of naive cd4 + t cells inhibits acquisition of ability to produce ifn- gamma and il-2, but not il-4 and il-5. | 0.006 | 0.002 |
| IL2 | FOXP3 | il-1b promotes tgf-b1 and il- 2 dependent foxp3 expression in regulatory t cells. | 0.006 | 0.002 |
| IL2 | IFNG | the detailed distribution of lymphokine-producing cells showed that il-2 and ifn-gamma-producing cells were located mainly in the follicular areas . | 0.005 | 0.001 |

| Gene1 Symbol | Gene2 Symbol | Text | Before Calibration | After Calibration |
|--------------|--------------|--|--------------------|-------------------|
| IFNG | IL10 | results: we found weak mrna expression of interleukin-4 (il-4) and il-5, and strong expression of il-6, il-10 and ifngamma before therapy. | 0.005 | 0.001 |
| PIK3R1 | PTEN | both pten (pi3k antagonist) and pp2 (unspecific phosphatas e) were down- regulated . | 0.005 | 0.001 |

Top Ten Sentences for Each Edge Type

Table 11: Contains the top ten predictions for each edge type. Highlighted words represent entities mentioned within the given sentence.

| Edge Type | Source Target Node Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text | |
|--------------|----------------------------|--------------------------------|------------------------------------|-------------------------------|-------------|------|--|
|--------------|----------------------------|--------------------------------|------------------------------------|-------------------------------|-------------|------|--|

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|------------------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| DaG | urinary bladder cancer | TP53 | 1 | 0.945 | 2112 | Existing | conclusion: our findings indicate that the dsp5 3-285 can upre gulate wild-type p53 expression in hum an bladder cancer cells through rna activation, and suppresses cells proliferation and metastasis in vitro and in vivo. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| DaG | ovarian | EGFR | 1 | 0.937 | 1330 | Existing | conclusion: our data show ed that incre ased expression of egfris associated with poor prognosis of patients with eoc and dacomitinib may act as a novel, usefulchem other apy drug. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|-----------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| DaG | stomac h cancer | TP53 | 1 | 0.937 | 2679 | Existing | conclusion: this meta-analy sis sugg ests that p53 arg72 pro poly morp hism is associated with increased risk of gastric cancer in asian s. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| DaG | lung | TP53 | 1 | 0.936 | 6813 | Existing | conclusion: these results suggest that high expression of the p53 onco prote in is a favor able prognostic factor in a subset of patients with nsclc. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| DaG | breast cancer | TCF7L2 | 1 | 0.936 | 56 | Existing | this meta - analy sis dem onstr ated that tcf7l2 gene poly morp hism s (rs122 5537 2 and rs790 3146) are associated with an incre ased susce ptibili ty to breas t canc er . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|-------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| DaG | skin cancer | COX2 | 1 | 0.935 | 73 | Novel | eleva ted expr essio n of cox-2 has been assoc iated with tumo r progr essio n in skin canc er throu gh multi ple mech anis ms. |
| DaG | thyroid cancer | VEGFA | 1 | 0.933 | 592 | Novel | as a conclusion, we sugg est that vegf g +405 c poly morp hism is associated with incre ased risk of ptc. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|-----------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| DaG | stomac h cancer | EGFR | 1 | 0.933 | 1237 | Existing | recently, high lymp h node ratio is closel y associated with egfrexpression in advanced gastric c cancer. |
| DaG | liver cancer | GPC3 | 1 | 0.933 | 1944 | Novel | concl usion s seru m gpc3 was overe xpres sed in hcc patie nts. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|-----------------------|-----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| DaG | stomac h cancer | CCR6 | 1 | 0.931 | 24 | Novel | the cox regre ssion analy sis show ed that high expr ession of ccr6 was an indep ende nt prog nosti c facto r for gc patie nts. |
| CtD | Sorafen ib | liver cancer | 1 | 0.99 | 6672 | Existing | tace plus soraf enib for the treat ment of hepa tocell ular carci nom a: final resul ts of the multi cente r socra tes trial. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|------------------|-----------------------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CtD | Methot rexate | rheuma toid arthritis | 1 | 0.989 | 14546 | Existing | comp ariso n of low- dose oral pulse meth otrex ate and place bo in the treat ment of rheu mato id arthri tis. |
| CtD | Aurano fin | rheuma toid arthritis | 1 | 0.988 | 419 | Existing | aura nofin versu s place bo in the treat ment of rheu mato id arthri tis. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|-----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| CtD | Lamivu dine | hepatiti s B | 1 | 0.988 | 6709 | Existing | rand omiz ed contr olled trials (rcts) comp aring etv with lam for the treat ment of hepa titis b deco mpe nsate d cirrh osis were inclu ded . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|-----------------|------------------------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CtD | Doxoru bicin | urinary bladder cancer | 1 | 0.988 | 930 | Existing | 17- year follo w-up of a rand omiz ed pros pecti ve contr olled trial of adjuv ant intra vesic al doxo rubici n in the treat ment of supe rficial bladd er canc er. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|------------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CtD | Doceta xel | breast cancer | 1 | 0.987 | 5206 | Existing | curre ntly, rand omiz ed phas e iii trials have dem onstr ated that docet axel is an effect ive strat egy in the adjuv ant treat ment of breas t canc er. |
| CtD | Cimetid ine | psoriasi s | 0.999 | 0.987 | 12 | Novel | cimet idine versu s place bo in the treat ment of psori asis . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|-------------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CtD | Olanza | schizop hrenia | 1 | 0.987 | 3324 | Novel | a doub le-blind , rand omis ed comp arativ e trial of amis ulpri de versu s olanz apine in the treat ment of schiz ophr enia: short -term resul ts at two mont hs . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|------------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| CtD | Fulvestr | breast cancer | 1 | 0.987 | 826 | Existing | phas e iii clinic al trials have dem onstr ated the clinic al bene fit of fulve stran t in the endo crine treat ment of breas t canc er. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|------------------|--------------------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| CtD | Pimecr olimus | atopic dermati tis | 1 | 0.987 | 531 | Existing | intro ducti on: altho ugh sever al controlled clinic al trials have dem onstrated the effica cy and good toler abilit y of 1 % pime croli mus crea m for the treat ment of atopi c derm atitis, the resul ts of these trials may not apply to reallife usag e. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CbG | Gefitini b | EGFR | 1 | 0.99 | 8746 | Existing | morp holog ic featu res of aden ocarc inom a of the lung predictive of resp onse to the epide rmal grow th facto r recep tor kinas e inhibi tors erloti nib and gefiti nib. |
| CbG | Adenosi | EGFR | 1 | 0.987 | 644 | Novel | it is well estab lishe d that inhibiting atp binding within the egfr kinas e domain regulates its function. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CbG | Rosiglit | PPARG | 1 | 0.987 | 1498 | Existing | rosiglitazo ne is a pote nt pero xiso me prolif erato r- activ ated recep tor gam ma agoni st that decre ases hype rglyc emia by redu cing insuli n resist ance in patie nts with type 2 diabe tes mellit us. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|--------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| CbG | D- Tyrosin e | INSR | 0.998 | 0.987 | 1713 | Novel | this resul t sugg ests that tyrosi ne phos phor ylatio n of phos phati dylin ositol 3-kinas e by the insuli n recep tor kinas e may incre ase the speci fic activi ty of the form er enzy me in vivo . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|--------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CbG | D- Tyrosin e | IGF1 | 0.998 | 0.983 | 819 | Novel | affini ty- purifi ed insuli n-like grow th facto r i recep tor kinas e is activ ated by tyrosi ne phos phor ylatio n of its beta subu nit . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| CbG | Pindolo I | HTR1A | 1 | 0.983 | 175 | Existing | pind olol, a beta block er with weak parti al 5-ht1a recep tor agoni st activi ty has been show n to prod uce a more rapid onset of antid epres sant actio n of ssris. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| CbG | Progest | SHBG | 1 | 0.981 | 492 | Existing | howe ver, dng also elicits properties of progester one deriv ative s like neutrality in meta bolic and cardi ovasc ular syste m and consi dera ble antia ndro genic activi ty, the latter incre ased by lack of binding to shbg as specific property of dng. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|------------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| CbG | Mifepri stone | AR | 1 | 0.98 | 78 | Existing | ru48 6 boun d to the andr ogen recep tor. |
| CbG | Alfenta nil | OPRM1 | 1 | 0.979 | 10 | Existing | purp ose: alfen tanil is a high pote ncy mu opiat e recep tor agoni st com monl y used durin g pres urgic al induc tion of anest hesia . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|-----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| CbG | Candes artan | AGTR1 | 1 | 0.979 | 36 | Existing | tcv- 116 is a new, nonp eptid e, angio tensi n ii type- 1 recep tor antag onist that acts as a speci fic inhibi tor of the renin - angio tensi n syste m. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| GiG | BRCA2 | BRCA1 | 0.972 | 0.984 | 12257 | Novel | a total of 9 famili es (16 %) show ed muta tions in the brca1 gene , inclu ding the one new muta tion identi fied in this study (5382i nsc) , and 12 famili es (21 %) prese nted muta tions in the brca2 gene . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| GiG | MDM2 | TP53 | 0.938 | 0.978 | 17128 | Existing | no muta tions in the tp53 gene have been foun d in samp les with ampli ficati on of mdm 2. |
| GiG | BRCA1 | BRCA2 | 1 | 0.978 | 12257 | Existing | path ogeni c trunc ating muta tions in the brca1 gene were foun d in two tumo r samp les with allelic losse s, wher eas no muta tions were identi fied in the brca2 gene . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| GiG | KRAS | TP53 | 0.992 | 0.971 | 4106 | Novel | muta tions in the p53 gene did not corre late with muta tions in the c-k-ras gene , indic ating that color ectal canc er can devel op throu gh path ways indep ende nt not only of the prese nce of muta tions in any of these gene s but also of their coop erati on . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| GiG | TP53 | HRAS | 0.992 | 0.969 | 451 | Novel | path ologic exam ination of the uc specimens from aaexpo sed patients identified heter ozygous hras chan ges in 3 cases, and deletion or replacement mutations in the tp53 gene in 4. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|--|
| GiG | REN | NR1H3 | 0.998 | 0.966 | 8 | Novel | nucle ar recep tor lxralp ha is invol ved in camp - medi ated hum an renin gene expr essio n . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| GiG | ESR2 | CYP19A 1 | 0.999 | 0.96 | 159 | Novel | dna meth ylatio n, histo ne modi ficati ons, and bindi ng of estro gen recep tor, erb to regul atory dna sequ ence s of cyp1 9a1 gene were evalu ated by chro mati n imm unop recipi tatio n (chip) assay . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| GiG | RET | EDNRB | 0.816 | 0.96 | 136 | Novel | muta tions in the ret gene, which code s for a recep tor tyrosi ne kinas e, and in ednr b which code s for the endo thelin-b recep tor, have been shown to be associated with hscrin hum ans. |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| GiG | PKD1 | PKD2 | 1 | 0.959 | 1614 | Existing | appr oxim ately 85 % of adpk d cases are caus ed by muta tions in the pkd1 gene , while muta tions in the pkd2 gene acco unt for the rema ining 15 % of cases . |

| Edge Type | Source Node | Target Node | Generative Model Prediction | Discriminative Model Prediction | Number of Sentenc es | In Hetionet | Text |
|--------------|----------------|----------------|--------------------------------|------------------------------------|-------------------------------|-------------|---|
| GiG | LYZ | CTCF | 0.999 | 0.959 | 2 | Novel | in conju nctio n with the thyro id recep tor (tr), ctcf bindi ng to the lysoz yme gene trans cripti onal silenc er medi ates the thyro id horm one resp onse elem ent (tre) depe nden t trans cripti onal repre ssion . |