

ProPheno: An online dataset for completely characterizing the human protein-phenotype landscape in biomedical literature

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ProPheno: An Online Dataset for Completely Characterizing the Human Protein-phenotype Landscape in Biomedical Literature

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ABSTRACT

Identifying protein-phenotype relations is of paramount importance for applications such as uncovering rare and complex diseases. One of the best resources that captures the protein-phenotype relationships is the biomedical literature. In this work, we introduce ProPheno, a comprehensive online dataset composed of human protein/phenotype mentions extracted from the complete corpora of Medline and PubMed. Moreover, it includes co-occurrences of protein-phenotype pairs within different spans of text such as sentences and paragraphs. We use ProPheno for completely characterizing the human protein-phenotype landscape in biomedical literature. ProPheno, the reported findings and the gained insight has implications for (1) biocurators for expediting their curation efforts, (2) researches for quickly finding relevant articles, and (3) text mining tool developers for training their predictive models. The RESTful API of ProPheno is freely available at http://propheno.cs.montana.edu.

INTRODUCTION

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Proteins are the workhorses of life, and they perform a wide range of operations in cells. Thousands of proteins work together to provide the functionality of cells. However, when a gene is mutated, a malfunction of protein may occur which can lead to a genetic disorder (NIH, 2018).

The observation of phenotypes is important in studying genetic disorders. In the medical context, a phenotype can be characterized as a deviation from normal morphology or behavior (Robinson, 2012). The study of phenotypes in medicine consists of detailed understanding of the phenotypic abnormalities associated with each disease (Robinson, 2012). Variations of genes and proteins cause functional changes and identifying the effects of their mutations is necessary for understanding the resulting phenotype, i.e. the observed disease state (Baker and Rebholz-Schuhmann, 2009). Furthermore, many patients suffer from rare diseases caused by genomic variants, i.e. diseases caused by disruptions in regular gene expression. However, many variants are quite rare, making genotype-phenotype correlations dubious and clinical interpretations difficult (Firth et al., 2009). One way to increase certainty would be identifying patients who share the same or overlapping gene variants and phenotypic characteristics (Firth et al., 2009). Therefore, finding relations between proteins and phenotypes can be considered vital for applications such as finding treatments and cures for rare diseases.

Since biologists, researchers, and scientists report their findings and observations from wet-lab experiments and clinical studies in biomedical literature, it can be considered one of the most valuable resources for extracting protein-phenotype relations. Therefore, exploring the feasibility of extracting relations between proteins and phenotypes mentioned in biomedical literature through text mining and machine learning has recently gained significant attention (Singhal et al., 2016; Korbel et al., 2005; Goh et al., 2006; Khordad and Mercer, 2017). However, the first main hurdle in developing text mining models is the absence of a gold-standard dataset of bio-entity mentions, i.e. proteins and phenotypes, which

encompasses the complete corpora of published biomedical articles. In this work, we attempt to resolve this issue by developing such a comprehensive dataset.

Human Phenotype Ontology (HPO) is a standardized vocabulary which covers a wide range of phenotype abnormalities observed in human diseases (Köhler et al., 2013). HPO is composed of five subontologies among which *Phenotypic abnormalities* is the main sub-ontology. *Phenotypic abnormalities* sub-ontology, which describes clinical abnormalities, is also called the *Organ abnormality*. Each subontology is organized in a hierarchical structure where more general terms are close to the top while more specific terms are closer to the bottom. Each pair of terms in the hierarchy are linked with a *is-a* relationship. In this paper, we use *phenotypes* and *HPO terms*, interchangeably.

HPO website¹ provides gold-standard annotations for a large collection of human proteins. However, currently, only a small portion of known human proteins have HPO annotations (Köhler et al., 2013). But, it is believed that there are many other human proteins which are associated with diseases and hence should be annotated with HPO terms (Peter Robinson, personal communication, 2015). But, biocuration, which is the process of manually extracting relevant information from biomedical literature, is well known to be a highly resource consuming task. Therefore, computational models for relation extraction could provide a solution in expediting and making it less tedious. However, first step in prediction of protein-phenotype relations is extracting protein and phenotype names from biomedical literature. There is no publicly available dataset of proteins and phenotype names on all the biomedical literature.

In this paper, we introduce ProPheno, which is an online and publicly accessible dataset composed of proteins, phenotypes (HPO terms), and their co-occurrences (co-mentions) in text which are extracted from Medline abstracts and PubMed full-text articles using a sophisticated in-house developed text mining pipeline. This dataset covers all terms in the *Organ abnormality* sub-ontology. We also conduct a comprehensive characterization of the protein-phenotype landscape in biomedical literature using this data. The findings from this characterization has implications for biocurators and researchers working in related fields as well as practitioners in the area of developing automated text mining pipelines for biocuration. The dataset is accessible through a RESTful API, which can be used in many programming language as well as a web interface (online demo version) for online access.

METHOD

We developed the text mining pipeline² shown in Figure 1, which is an extension of the pipeline described elsewhere (Pourreza Shahri and Kahanda, 2018). This pipeline extracts proteins/phenotypes and their co-mentions by consuming 27,590,898 Medline abstracts (downloaded on 07/01/2017) and 1,873,381 PubMed full-text articles (downloaded on 3/15/2018) as the input. PubMed full-text articles were downloaded from the PubMed website in XML format. A small portion of these full-text articles (140,370) did not contain any text, so we removed them from consideration. We employed *PubMed XML Parser* (Achakulvisut and Acuna, 2015) to extract the paragraphs from the remaining 1,733,012 full-text articles, and stored the corresponding paragraphs in separate files. It is worth mentioning that to avoid duplicating abstracts, we do not take the abstract section of full-text articles into account.

We employed NCBO Virtual Appliance (NCBO Annotator) from BioPortal (Jonquet et al., 2009; Noy et al., 2009) for extracting HPO terms from the literature. Protein mentions were retrieved from the literature using LingPipe (Carpenter, 2007). We used UniProt (Apweiler et al., 2004) synonyms³ of proteins to improve the coverage when extracting proteins from literature.

We also considered other alternatives to extract these entities such as OBO annotator (Taboada et al., 2014) and Bio-Lark CR (Groza et al., 2015) for extracting phenotype names, and GNormPlus (Wei et al., 2015) and ABNER (Settles, 2005) for extracting protein names. However, most of these systems either did not provide desirable results or were difficult to access or use.

The introduced dataset comprises UniProt identifiers and HPO identifiers for proteins and phenotypes extracted from biomedical literature, respectively, which are categorized by the abstracts and full-text articles. This dataset also provides co-mentions of proteins and phenotypes in different spans of text. We used three spans: (1) sentence-level co-mentions (SCoM) which occur in a single sentence (2) paragraph-level co-mentions (PCoM) which occur in a single paragraph (i.e. across multiple sentences), and (3) document-level co-mentions (DCoM) which occur in a single document (i.e. across multiple paragraphs).

¹https://hpo.jax.org/app

²Image created using https://www.draw.io

 $^{^3} ftp: //ftp.uniprot.org/pub/databases/uniprot/current_release/knowledgebase/idmapping/by_organism/HUMAN_9606_idmapping.dat.gz$

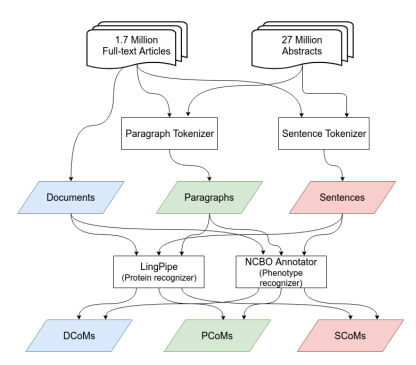


Figure 1. Overview of the text mining pipeline for extracting protein and phenotype names as well as their co-occurrences within different spans of text.

We have previously worked on extracting co-mentions in a limited setting (Pourreza Shahri and Kahanda, 2018) in which we showed that SCoMs and PCoMs can improve the performance of protein-phenotype prediction tools. In an effort to primarily make these co-mentions readily available for other researchers in the filed, ProPheno was born. Figure 2 shows an example of a co-mention of a protein and a phenotype in a sentence (i.e. a sentence-level co-mention). Figure 3 is a screenshot of ProPheno demo version, which shows a list of 10 random HPO terms extracted from abstracts. In this figure, each row depicts an occurrence of a phenotype mentioned in text, and "Start Location" and "End Location" indicated the actual location of occurrence.

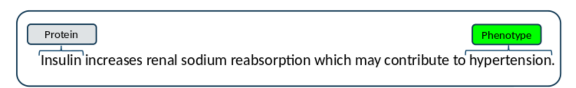


Figure 2. An example of a sentence-level protein-phenotype co-mention which is extracted from the article PMID: 10855734. The corresponding UniProt ID and HPO ID for the protein and the phenotype are P01308 and HP:0000822, respectively.

RESULTS

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In this section we present detailed statistics and analysis on the mentions of proteins and phenotypes in different spans of text.

Proteins and Phenotypes

Table 1 shows the total number of the unique and all proteins and phenotypes in abstracts and full-text articles. Since full-text articles include more details about proteins and phenotypes, intuitively, we expect more terms in full-text articles. The average number of proteins and phenotypes in abstracts and full-text articles supports this claim. By looking at the unique number of proteins and phenotypes in abstracts versus full-text articles, we observe that there are a few proteins and phenotypes which were extracted



ProPheno

PubMed ID	HPO ID	Start Location	End Location
304747	HP:0002665	694	702
304750	HP:0001945	98	103
304749	HP:0002045	440	451
304751	HP:0009778	11	22
304759	HP:0003002	57	70
304759	HP:0002664	64	70
304760	HP:0001882	155	165
304760	HP:0000716	189	199
304768	HP:0001627	283	307
304771	HP:0002088	1009	1021

Figure 3. The list of 10 random HPO terms mentioned in the abstracts shown in the online demo version. Each row shows an occurrence of an HPO term in an abstract. "Start Location" and "End Location" show the position of the matched phenotype in text, e.g. Start Location = 98 and End Location = 103 show that the phenotype has been mentioned in text starting from index 98 and ends at index 103.

only from either abstracts or full-text articles. In this study, we consider various combination of words which are detected by NCBO Annotator that are matched with a term in the HPO database. For instance, NCBO Annotator returns "prostate cancer", i.e. HP:0012125, and "cancer", i.e. HP:0002664, from "... prostate cancer ...", and both words can be matched with corresponding HPO terms in the HPO database. This can be attributed as the main reason for having a large number of extracted phenotypes.

Table 1. Stats of Protein/Phenotype Mentions in Biomedical Literature

	Abstracts	Full-texts	Total
Unique proteins	2,178	2,389	2,512
All proteins	1,807,246	2,173,695	3,980,941
Avg. number of proteins	2.11	4.63	-
Unique phenotypes	2,224	2,227	2,277
All phenotypes	30,954,930	32,639,095	63,594,025
Avg. number of phenotypes	3.58	24.8	-

Figures 4 and 5 show the distribution of unique proteins and phenotypes in abstracts and full-text articles, respectively⁴. As mentioned before, there are a few proteins and phenotypes which were extracted from either abstracts or full-text articles. We also observe that our pipeline can detect 74% of the proteins and 92% of the phenotypes curated in the HPO database.

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⁴Diagrams generated by https://github.com/tctianchi/pyvenn

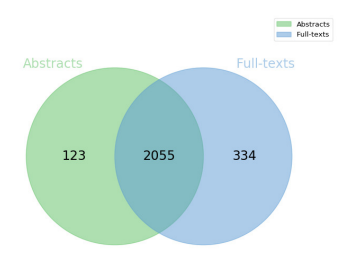


Figure 4. The Distribution of Unique Proteins in Abstracts and Full-texts

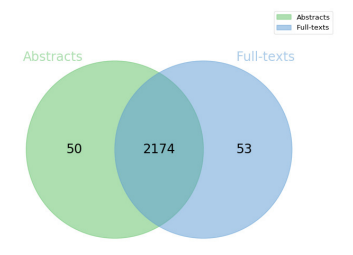


Figure 5. The Distribution of Unique Phenotypes in Abstracts and Full-texts

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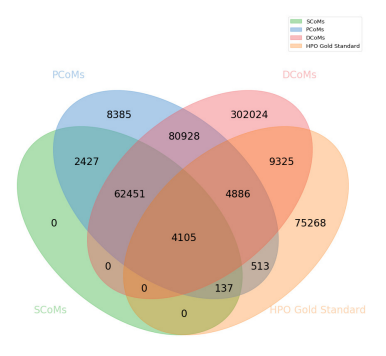


Figure 6. The Distribution of Unique Pairs of Co-mentions in SCoMs, PCoMs, DCoMs, and protein-HPO annotations curated in the HPO database

21 Co-mention of Proteins and Phenotypes in Text

As mentioned in the previous section, co-mentions are defined as the co-occurrences of proteins and phenotypes within a certain span of text extracted from our text mining pipeline. Statistics on these co-mentions in various spans of text, i.e. sentences, paragraphs, and full-text documents, are given in Table 2. The distribution of unique pairs of proteins and phenotypes in SCoMs, PCoMs, DCoMs, and the HPO gold standard are shown in Figure 6. We observe that 75,268 unique pairs in the HPO gold standard were not extracted by our pipeline. There are two possible reasons that can prevent the pipeline to extract these pairs. First, either LingPipe or NCBO Annotator may have failed to extract the corresponding proteins or phenotypes, respectively. Second, they may have been either not co-occurred in at least a document or one entity is mentioned in the abstract while the other entity is in the body of the article. Besides, there are 10,812 unique pairs in PCoMs (2,427 in common with SCoMs) which are not recognized by the DCoMs. This is because we treat the abstracts as paragraphs, and since we only have one copy of the abstracts, DCoMs do not contain those unique pairs.

Analysis of Protein and Phenotype Named Entities

Figure 7 shows the distribution of protein names in SCoMs, PCoMs, and DCoMs. The number of less frequent proteins in SCoMs is more than proteins with high frequency, and the number of proteins in for each frequency in SCoMs is higher than corresponding number in PCoMs and DCoMs. This observation suggests that the larger spans of text are able to identify more proteins.

Table 2. Statistics of Co-mentions Extracted from both Medline and PubMed. We consider abstracts as paragraphs, so we do not have Document-level information for the abstracts.

Full-text articles						
Span	Unique proteins	Unique HPO terms	Unique co-mentions	Total co-mentions		
Sentence-level	1,845	1,498	49,686	693,005		
Paragraph-level	2,185	1,896	122,522	4,323,395		
Document-level	2,362	2,126	463,719	99,818,140		
Abstracts						
Span	Unique proteins	Unique HPO terms	Unique co-mentions	Total co-mentions		
Sentence-level	1,684	1,461	42,774	496,969		
Paragraph-level	1,975	1,827	102,881	4,116,999		
Document-level	-	-	-	-		
All						
Span	Unique proteins	Unique HPO terms	Unique co-mentions	Total co-mentions		
Sentence-level	1,998	1,623	69,120	1,189,974		
Paragraph-level	2,313	1,976	225,403	8,440,394		
Document-level	2,362	2,126	436,719	99,818,140		

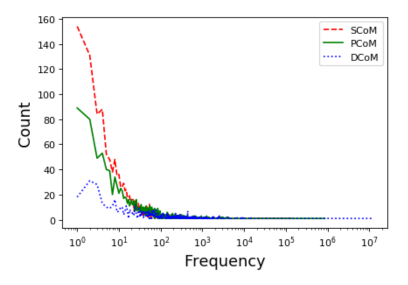


Figure 7. The Distribution of Protein Mentions

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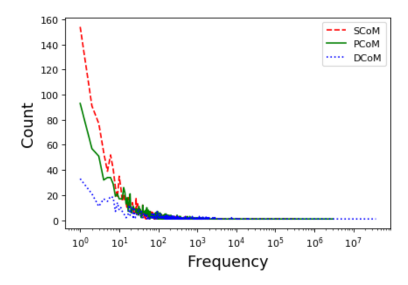


Figure 8. The Distribution of HPO-term Mentions

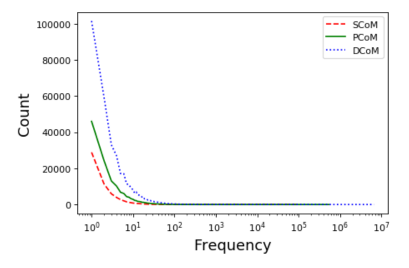


Figure 9. The Distribution of the Unique pair mentions

The same can be observed in Figure 8 which shows distribution of HPO terms in SCoMs, PCoMs, and DCoMs. However, in Figure 9 which shows the distribution of unique protein and HPO-term pairs, occurrences of unique pairs for each frequency in DCoMs is higher than corresponding values in SCoMs and PCoMs. One of the reasons of this observation can be that many of the unique pairs occur in sentences and do not occur more in larger spans of text. Therefore, larger spans of text have more occurrences of unique pairs for every frequency.

Figure 10 shows the distribution of the depths (with respect to the hierarchy) of all HPO terms recognized in the literature. This plot conveys that more specific HPO terms are less frequent in the literature. One of the reasons of this observation can be that there are less number of mentions of more specific terms in the literature. Figure 11 demonstrates the distribution of unique HPO terms detected in the literature and HPO terms from the HPO gold standard. We observe that our pipeline is able to detect the majority of HPO terms from the HPO gold standard.

In addition, the evolution of the protein-phenotype landscape in biomedical literature is shown in Figure 12. We observe a slight increase in the number of unique proteins, and a relatively higher increase in the number of unique HPO terms between 2009 and 2018.

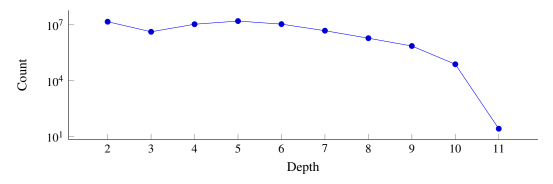


Figure 10. Distribution of the depths of terms detected in literature

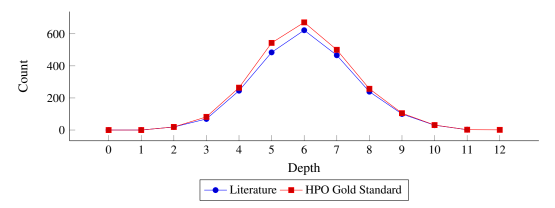


Figure 11. Distribution of the Depths of Unique Terms Detected in Literature and HPO Gold Standard

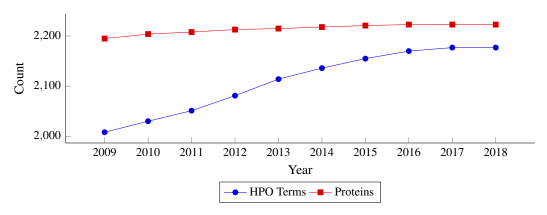


Figure 12. The evolution of the protein-phenotype landscape in Biomedical Literature

CONCLUSIONS AND FUTURE WORK

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In this paper we presented a dataset of proteins and phenotypes (HPO labels) in the entire biomedical corpora derived from Medline abstracts and PubMed full-text articles. This dataset was generated using an expanded text mining pipeline from one of our previous work (Pourreza Shahri and Kahanda, 2018).

We also reported detailed analysis and statistics on the mentions of proteins and phenotypes in the entire corpora along with co-mentions of these entities in various spans of biomedical text. Additionally, we presented the evolution of biomedical literature over the period of 2009-2018.

In this study, we used bio-entity recognizers which had shown good performance in identifying bioentities in biomedical text. However, bio-entity recognition is still a challenging problem. Consequently, advances in the performance of bio-entity recognizers would enhance the ability of our pipeline for correctly detecting more proteins and phenotypes in text.



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Despite detecting millions of proteins and phenotypes in text, it is understood that a portion of the comentions are false positives (the mere occurrence of two entities within a certain span does not constitute a valid relationship). Therefore, the next step would be developing a context-sensitive co-mention classifier or a filter to remove these false positives. This will provide the capability of listing the co-mentions in a ranked order according to the confidence scores predicted by the classifier.

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REFERENCES

- Achakulvisut, T. and Acuna, D. E. (2015). Pubmed Parser.
- Apweiler, R., Bairoch, A., Wu, C. H., Barker, W. C., Boeckmann, B., Ferro, S., Gasteiger, E., Huang, H., Lopez, R., Magrane, M., et al. (2004). UniProt: the universal protein knowledgebase. *Nucleic acids research*, 32(suppl_1):D115–D119.
- Baker, C. J. and Rebholz-Schuhmann, D. (2009). Between proteins and phenotypes: annotation and interpretation of mutations.
- Carpenter, B. (2007). LingPipe for 99.99% recall of gene mentions. In *Proceedings of the Second BioCreative Challenge Evaluation Workshop*, volume 23, pages 307–309.
- Firth, H. V., Richards, S. M., Bevan, A. P., Clayton, S., Corpas, M., Rajan, D., Van Vooren, S., Moreau, Y.,
 Pettett, R. M., and Carter, N. P. (2009). DECIPHER: database of chromosomal imbalance and phenotype in humans using ensembl resources. *The American Journal of Human Genetics*, 84(4):524–533.
- Goh, C.-S., Gianoulis, T. A., Liu, Y., Li, J., Paccanaro, A., Lussier, Y. A., and Gerstein, M. (2006). Integration of curated databases to identify genotype-phenotype associations. *BMC genomics*, 7(1):257.
- Groza, T. et al. (2015). Automatic concept recognition using the Human Phenotype Ontology reference and test suite corpora. *Database*, 2015:bav005.
- Jonquet, C., Shah, N. H., and Musen, M. A. (2009). The open biomedical annotator. *Summit on translational bioinformatics*, 2009:56.
- Khordad, M. and Mercer, R. E. (2017). Identifying genotype-phenotype relationships in biomedical text. *Journal of biomedical semantics*, 8(1):57.
- Köhler, S., Doelken, S. C., et al. (2013). The Human Phenotype Ontology project: linking molecular biology and disease through phenotype data. *Nucleic acids research*, 42(D1):D966–D974.
- Korbel, J. O., Doerks, T., Jensen, L. J., Perez-Iratxeta, C., Kaczanowski, S., Hooper, S. D., Andrade,
 M. A., and Bork, P. (2005). Systematic association of genes to phenotypes by genome and literature
 mining. *PLoS biology*, 3(5):e134.
- NIH (2018). How can gene mutations affect health and development?
- Noy, N. F. et al. (2009). BioPortal: ontologies and integrated data resources at the click of a mouse.

 Nucleic acids research, 37(suppl_2):W170–W173.
- Pourreza Shahri, M. and Kahanda, I. (2018). Extracting co-mention features from biomedical literature
 for automated protein phenotype prediction using PHENOstruct. In 10th International Conference on
 Bioinformatics and Computational Biology, BICOB 2018, pages 123–128.
- Robinson, P. N. (2012). Deep phenotyping for precision medicine. *Human mutation*, 33(5):777–780.
- Settles, B. (2005). ABNER: an open source tool for automatically tagging genes, proteins and other entity names in text. *Bioinformatics*, 21(14):3191–3192.
- Singhal, A., Simmons, M., and Lu, Z. (2016). Text mining genotype-phenotype relationships from biomedical literature for database curation and precision medicine. *PLoS computational biology*, 12(11):e1005017.
- Taboada, M., Rodríguez, H., Martínez, D., Pardo, M., and Sobrido, M. J. (2014). Automated semantic annotation of rare disease cases: a case study. *Database*, 2014.
- Wei, C.-H., Kao, H.-Y., and Lu, Z. (2015). GNormPlus: an integrative approach for tagging genes, gene families, and protein domains. *BioMed research international*, 2015.