PHENOstruct: Prediction of human phenotype ontology terms using heterogeneous data sources [version 1; referees: 2 approved]

Indika Kahanda¹, Christopher Funk², Karin Verspoor³,⁴, Asa Ben-Hur¹

¹Department of Computer Science, Colorado State University, Fort Collins, CO, 80523, USA
²Computational Bioscience Program, University of Colorado School of Medicine, Aurora, CO, 80045, USA
³Department of Computing and Information Systems, University of Melbourne, Parkville, Victoria, 3010, Australia
⁴Health and Biomedical Informatics Centre, University of Melbourne, Parkville, Victoria, 3010, Australia

Abstract

The human phenotype ontology (HPO) was recently developed as a standardized vocabulary for describing the phenotype abnormalities associated with human diseases. At present, only a small fraction of human protein coding genes have HPO annotations. But, researchers believe that a large portion of currently unannotated genes are related to disease phenotypes. Therefore, it is important to predict gene-HPO term associations using accurate computational methods. In this work we demonstrate the performance advantage of the structured SVM approach which was shown to be highly effective for Gene Ontology term prediction in comparison to several baseline methods. Furthermore, we highlight a collection of informative data sources suitable for the problem of predicting gene-HPO associations, including large scale literature mining data.

Corresponding author: Asa Ben-Hur (asa@cs.colostate.edu)

How to cite this article: Kahanda I, Funk C, Verspoor K and Ben-Hur A. PHENOstruct: Prediction of human phenotype ontology terms using heterogeneous data sources [version 1; referees: 2 approved] F1000Research 2015, 4:259 (doi: 10.12688/f1000research.6670.1)

Copyright: © 2015 Kahanda I et al. This is an open access article distributed under the terms of the Creative Commons Attribution Licence, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Grant information: This work was supported by the NSF Advances in Biological Informatics program through grants number 0965768 (awarded to Dr. Ben-Hur) and 0965616 (originally awarded to Dr. Verspoor).

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing interests: The authors declare that they have no competing interests.

First published: 16 Jul 2015, 4:259 (doi: 10.12688/f1000research.6670.1)
Introduction
In the medical context a phenotype is defined as a deviation from normal morphology, physiology, or behavior. The human phenotype ontology (HPO) is a standardized vocabulary that describes the phenotype abnormalities encountered in human diseases. It was initially populated using databases of human genes and genetic disorders such as OMIM, Orphanet and DECIPHER, and was later expanded using literature curation. The hierarchical structure of the HPO is very similar to that of the Gene Ontology (GO), and it too has the structure of a directed acyclic graph (DAG); like GO, more general terms are found at the top, and term specificity increases from the root to the leaves. This implies the “true-path rule”: whenever a gene is annotated with a given term, that implies all its ancestor terms.

HPO is composed of three subontologies: organ abnormality, mode of inheritance, and onset and clinical course. Organ abnormality is the main subontology which describes clinical abnormalities (Figure 1). The mode of inheritance subontology describes the inheritance patterns of the phenotypes. The onset and clinical course subontology describes the typical time of onset of clinical symptoms and their speed of progression. The organ abnormality, mode of inheritance and onset and clinical course subontologies are composed of ~10000, 25 and 30 terms respectively. Throughout this paper, the organ abnormality, the mode of inheritance, and the onset and clinical course subontologies will be referred to as the Organ subontology, Inheritance subontology and Onset subontology, respectively.

The HPO web site (http://www.human-phenotype-ontology.org) provides gene-disease-HPO annotations that can be used for research involving human diseases. Over 50,000 annotations of hereditary diseases are available at the moment. Specifically, the genes are annotated with a set of phenotype terms based on their known relationships with diseases (Figure 2).

Currently, only a small fraction (~3000) of human protein coding genes are known to be associated with hereditary diseases, and only those genes have HPO annotations at the moment. But researchers believe that there are many other disease-causing genes in the human genome and estimate that another 5000 genes can be associated with phenotypes (Peter Robinson, personal communication, 2014). However, experimentally finding disease-causing genes is a highly resource consuming and difficult task. Therefore, it is important to explore the feasibility of developing computational methods for predicting gene-HPO associations. While there is a plethora of computational approaches for the related task of prediction of gene-disease associations, no computational method that directly predicts gene-HPO term associations exists at this time.

Approach
We define the HPO prediction problem as directly predicting the complete set of HPO terms for a given gene. This problem is a hierarchical multilabel classification (HMC) problem, as a given gene can be annotated with multiple labels, and the set of labels have a hierarchy associated with them.

The traditional approach for solving HMC problems is to decompose the problem into multiple single label problems and apply independent binary classifiers for each label separately; however, this approach has several disadvantages. First, independent classifiers are not able to learn from the inter-relationships between the labels. Second, the leaf terms typically have a low number of annotated examples making it difficult to learn an effective classifier. Furthermore, the predicted labels are typically hierarchically inconsistent, i.e. a child term (e.g. Hearing abnormality) is predicted while its parent term (e.g. Abnormality of ear) is not—making it difficult to interpret the predictions. To remedy this problem, an additional reconciliation step of combining independent predictions to obtain a set of predictions that are consistent with the topology of the ontology is required (see e.g. 11 for a discussion of several reconciliation methods that are effective for GO term prediction).

![Figure 1. A portion of the Organ abnormality subontology. All HPO parent-child relationships represent “is-a” relationships.](image)

![Figure 2. HPO annotations. a) general format of annotations: genes are annotated with a set of phenotype terms based on their known relationships with diseases b) an example annotation: the amyloid precursor protein (APP) gene is associated with Alzheimer’s disease and cerebroarterial amyloidosis. Therefore, the APP gene is annotated with the set of HPO terms (Organ in orange, Inheritance in green) associated with these diseases.](image)
An alternative approach is to use a single classifier that learns a direct mapping from inputs to the space of hierarchically consistent labels; this can be achieved using structured prediction, which is a framework for learning a mapping from inputs to label spaces that have a structure associated with them. This framework can capture information from the inter-relationships between labels and allows the prediction of a set of labels that are hierarchically consistent, eliminating the need for multiple classifiers, and the need for establishing hierarchical consistency between the predictions. Previously we have shown the effectiveness of modeling the GO term prediction problem using a structured prediction framework in a method called GOstruct. In this work we demonstrate the effectiveness of this strategy for HPO term prediction using the same methodology, and explore a variety of data sources that are useful for this task, including large scale data extracted from the biomedical literature.

Methods

Data

Our models are provided with feature vectors and HPO annotations. Each gene/protein was characterized by several sets of features generated using four data sources: Network, GO, literature and variants, which are described below. We used the UniProt ID mapping service (http://www.uniprot.org/mapping/) for mapping genes to proteins.

HPO annotations

Gene-HPO annotations were downloaded from the HPO website (http://www.human-phenotype-ontology.org). We ignored the global root term (“ALL”) and root terms of the three subontologies. We also removed terms that were not annotated to 10 or more genes. Then we mapped the genes to proteins and generated corresponding protein-HPO annotations (see Table 1).

Network

We extracted protein-protein interactions and other functional association network data (i.e. co-expression, co-occurrence, etc.) from BioGRID 3.2.106, STRING 9.1 and GeneMANIA 3.1.2 (http://pages.genemania.org/data/) databases.

The BioGRID database provides protein-protein interaction networks acquired from physical and genetic interaction experiments. STRING provides networks based on several different evidence channels (co-expression, co-occurrence, fusion, neighborhood, genetic interactions, physical interactions, etc.). We combined edges from the two databases by taking the union of interactions from BioGRID and STRING and represented each gene by a vector of variables, where component \( i \) indicates if the corresponding protein interacts with protein \( i \) in the combined network.

The GeneMANIA website (http://pages.genemania.org/data/) provides a large number of protein-protein interaction/association networks generated using several types of evidence: co-expression, co-localization, genetic interactions, physical interactions and predicted interactions. A gene is represented by a vector of variables for each network, where component \( i \) indicates if the corresponding protein interacts with protein \( i \) with respect to that particular network.

Gene Ontology

We extracted GO annotations from the GO website (http://www.geneontology.org/) and UniProt-goa (http://www.ebi.ac.uk/GOA). We excluded all annotations that were obtained by computational methods. A gene is represented as a vector of indicator variables in which variable \( i \) is 1 if it is annotated with GO term \( i \).

Literature

We used two different sources for generating literature features: abstracts extracted from Medline on 10-23-13 and full-text articles extracted from PubMed Open Access Collection (PMCOA) on 11-06-13. A natural language processing pipeline was utilized to characterize genes/proteins by same-sentence word occurrences extracted from these sources, forming a bag-of-words (BoW) representation for each gene. First, all words were lower-cased and stop words were removed. Then they were further filtered to keep only the low frequency words (i.e. words that are present only in less than 1% of the proteins in the data). A gene is represented by a vector in which the element \( i \) gives the number of times the word \( i \) occurred in the same sentence with that gene/protein.

Variants

We extracted all the disease variants in the human genome and their associated diseases from UniProt (http://www.uniprot.org/docs/humsvavar). This data provides variants that have been found in patients and the disease-association is reported in literature. We also extracted gene-disease associations from the HPO website. This data associates a protein with diseases that are known to occur when the associated gene is mutated. To generate features from this data, we first extracted for each protein \( i \) its set of associated diseases \( D_i \) from the UniProt disease variants data. Finally, each gene was represented by a vector in which element \( j \) indicates if the variant \( j \) is in \( V_i \).

Models

In this work we compare a structured support vector machine approach against several baseline methods: a) binary support vector machines (SVMs) and b) a state-of-the-art HMC method based on decision tree ensembles (Clus-HMC-Ens). In this section we describe PHENOSTruct and the two baseline methods. In addition, we assessed the performance of: c) an indirect method that first predicts disease terms for a gene using a structured model and then
PHENOstruct

In earlier work we developed the GOstruct method which uses structured SVMs (SSVM) for GO term prediction\(^{13}\). In this work we apply the same methodology to HPO term prediction and refer to it as PHENOstruct to emphasize the different problem domain. Unlike collections of binary classifiers applied independently at each node of the hierarchy, PHENOstruct predicts a set of hierarchically consistent HPO terms for a given gene (Figure 3). More specifically, PHENOstruct learns a compatibility function that models the association between a given input and a structured output\(^{12}\), in this case the collection of all hierarchically consistent sets of HPO terms. Let \( \mathcal{X} \) be the input space where genes are represented and let \( \mathcal{Y} \) be the space of labels. The set of HPO terms associated with a given gene is collectively referred to as its (structured) label. \( \mathcal{Y} \) represents each HPO subontology in a vector space where component \( i \) represents term \( i \). Given a training set \( \{(x_i, y_i)\}_{i=1}^n \) where \( x_i \in \mathcal{X} \) and \( y_i \in \mathcal{Y} \), the compatibility function \( f: \mathcal{X} \times \mathcal{Y} \rightarrow \mathbb{R} \) maps input-output pairs to a score that indicates how likely is a gene \( x \) to be associated with a collection of terms represented by \( y \). The predicted label \( \hat{y} \) for an unseen input \( x \) can then be obtained by using the argmax operator as \( \hat{y} = \arg\max_{y \in \mathcal{Y}} f(x, y) \) where \( \mathcal{Y}_c \subseteq \mathcal{Y} \) is the set of all candidate labels. In this work we use the combinations of all terms in the training set as the set of candidate labels \( \mathcal{Y}_c \).

In order to obtain correct classification, the compatibility value of the true label (correct set of HPO annotations) of an input needs to be higher than that of any other candidate label (Figure 4). PHENOstruct uses structured SVM (SSVM) training where this is
used as a (soft) constraint; it tries to maximize the margin, or the difference between the compatibility value for the actual label and the compatibility for the next best candidate\[^1\]. In the structured-output setting, kernels correspond to dot products in the joint input-output feature space, and they are functions of both inputs and outputs. PHENOstruct uses a joint kernel that is the product of the input-space and the output-space kernels:

$$K((x_1, y_1), (x_2, y_2)) = K_1(x_1, x_2)K_2(y_1, y_2).$$

The motivation for this form is that two input/output pairs are considered similar if they are similar in both their input space features and their labels; the output space kernel, for which we use a linear kernel between label vectors, captures similarity of the annotations associated with two genes; the input space kernel combines several sources of data by the addition of multiple input-space kernels, one for each data source. Each kernel is normalized according to

$$K_{\text{norm}}(z_1, z_2) = K(z_1, z_2) / \sqrt{K(z_1, z_1)K(z_2, z_2)}$$

before being used with the joint input-output kernel. The Strut library (http://sourceforge.net/projects/strut/) with default parameter settings was used for the implementation of PHENOstruct.

**Binary SVMs**

As a baseline method we trained a collection of binary SVMs, each trained on a single HPO term. Binary SVMs were trained using the PyML (http://pyml.sourceforge.net) machine learning library with default parameter settings. We used linear kernels for each set of input space features.

**Clus-HMC-Ens**

Clus-HMC-Ens is a state-of-the-art HMC method based on decision tree ensembles which has been shown to be very effective for GO term prediction\[^1\]. In our study, we provide exactly the same set of features used with PHENOstruct as input to Clus-HMC-Ens and use parameter settings that provided the best performance for GO term prediction (https://dtai.cs.kuleuven.be/clus/hmc-ens/). The number of bags used was 50 for the Inheritance and Onset subontologies; 10 bags were used for the Organ subontology because of the large running times for this subontology.

**Evaluation**

Classifier performance was estimated using five-fold cross-validation. Since typically scientists/biologists are interested in knowing the set of genes/proteins associated with a certain HPO term, we primarily use a *term-centric* measure for presenting results. Term-centric measures average performance across terms as opposed to *protein-centric* measures which average performance across proteins as described elsewhere\[^1\]. More specifically, we use the macro AUC (area under the receiver operating curve), which is computed by averaging the AUCs across HPO terms. For comparing performance across classifiers, p-values were computed using paired t-tests. Additionally, we report performance in terms of several protein-centric measures (precision, recall, F-max) in the Supplementary material (Table S3 and Table S4). Definitions of all performance measures are given in the Supplementary material. PHENOstruct assigns a confidence score to each predicted HPO term, which is computed using the compatibility function as described elsewhere\[^4\]. The onset and clinical course subontology includes terms such as *pace of progression*, *age of onset* and *onset* which are only used for grouping terms. We ignore these grouping terms when computing performance.

**Results and discussion**

**PHENOstruct performance**

As illustrated in Table 2, PHENOstruct significantly outperforms Clus-HMC-Ens and the binary SVMs in the Organ and Onset subontologies. This suggests that modeling the HPO prediction problem as a structured prediction problem is highly effective. It is interesting to note that the biggest improvement of PHENOstruct over binary SVMs is seen in the Organ subontology. Given its very large number of terms, as well as the deep hierarchy, this further confirms the value of the structured approach. PHENOstruct outperforms binary SVMs in the Inheritance and Onset subontologies but to a lesser extent than in the Organ subontology because they are far less complex than the Organ subontology. We note that the two methods that first predict OMIM terms, which are then mapped

| Subont. | Terms | Method          | AUC  | P-value   |
|---------|-------|----------------|------|-----------|
| Organ   | 1,796 | Binary SVMs    | 0.66 | 1.7E-262  |
|         |       | Clus-HMC-Ens   | 0.65 | 0.0E+00   |
|         |       | PHENOstruct    | 0.73 | —         |
| Inherit.| 12    | Binary SVMs    | 0.72 | 2.2E-01   |
|         |       | Clus-HMC-Ens   | 0.73 | 7.3E-01   |
|         |       | PHENOstruct    | 0.74 | —         |
| Onset   | 23    | Binary SVMs    | 0.62 | 4.4E-03   |
|         |       | Clus-HMC-Ens   | 0.58 | 3.3E-05   |
|         |       | PHENOstruct    | 0.64 | —         |
to HPO terms performed poorly (see details in the Supplementary material). It is also interesting to see that Clus-HMC-Ens performs worse than binary SVMs with respect to macro AUC (Table 2) but performs slightly better than binary SVMs according to protein-centric F-max (Table S3).

PHENOstruct’s average AUC for the Organ and Inheritance subontologies are 0.73 and 0.74, respectively. Even though the Organ subontology is a far more complex subontology than the Inheritance subontology (with thousands of terms and 13 levels as opposed to tens of terms and only 3 levels) they show similar performance. The Onset subontology is the hardest to predict accurately, with an average AUC of 0.64. Only six Onset subontology terms have individual AUCs above 0.7 (Table 4).

Even though PHENOstruct outperforms the baseline methods, there is much room for improvement, especially in the Onset subontology. The small number of annotated genes in this subontology (Table 1) makes it difficult to train an effective model while the incomplete nature of the current gold standard used for evaluation tends to underestimate performance of classifiers\(^1\). See section for a detailed analysis of false positives.

In general, Organ subontology terms with few annotations show a mix of both high and low performance as illustrated in Figure 5. This suggests that PHENOstruct is not necessarily affected by the frequency of the terms. But, terms with more annotations tend to show moderate performance. See Figure 6 for an example of experimental and predicted annotations (Organ subontology) for a protein.

It is interesting to note that “polygenic inheritance” and its parent term “multifactorial inheritance” have the lowest number of annotations as well as the lowest individual AUCs in the Inheritance subontology (see Table 3). These are the two terms with the lowest AUC with binary SVMs as well (see Table S6). It is not surprising that these two terms have lower accuracy because each describes inheritance patterns that depend on a mixture of determinants. Moreover, the diseases inherited in this manner – termed complex diseases – are not as well characterized and annotated compared to Mendelian/single gene diseases. On the other hand, the mitochondrial inheritance term has an exceptional AUC of 0.98. It is also the term with the highest AUC with the binary SVMs as well (see Table S6). The human mitochondrial DNA was the first significant part of the human genome to be fully sequenced, two decades before the completion of the human genome project\(^5\). Due to this, and the relative ease of sequencing the mitochondrial genome\(^2\), diseases caused by mutations in human mitochondrial DNA have been reported very early\(^2\). It is likely that this well-studied nature of mitochondrial DNA leads to the high performance of the mitochondrial inheritance term.

| Name                        | Freq. | Depth | AUC |
|------------------------------|-------|-------|-----|
| Multifactorial inheritance   | 15    | 1     | 0.54|
| Polygenic inheritance        | 15    | 2     | 0.54|
| Mitochondrial inheritance    | 41    | 1     | 0.98|
| Sporadic                     | 52    | 1     | 0.61|
| Somatic mutation             | 61    | 1     | 0.76|
| X-linked dominant inheritance| 62    | 3     | 0.83|
| X-linked recessive inheritance| 111  | 3     | 0.77|
| Heterogeneous                | 148   | 1     | 0.69|
| Gonosomal inheritance        | 198   | 1     | 0.80|
| X-linked inheritance         | 198   | 2     | 0.80|
| Autosomal dominant inherit.  | 1096  | 1     | 0.78|
| Autosomal recessive inher.   | 1665  | 1     | 0.73|

**Table 3. Performance of PHENOstruct in the Inheritance subontology.** The average macro AUC for the Inheritance subontology is 0.74. Terms are displayed in ascending order of frequency.
Figure 5. **Performance of PHENOstruct in the Organ subontology.** Performance for each term is displayed using AUC against its frequency. The average AUC for the Organ subontology is 0.73.

Figure 6. **Example of experimental and predicted annotations.** a) experimental annotation of protein P43681  b) PHENOstruct's prediction for P43681 (protein-centric precision and recall for this individual protein is 1.0 and 0.62, respectively).
As a potential improvement to PHENOstruct we explored an approximate inference algorithm that replaces computation of the most compatible label by looping overall combinations of labels that occur in the training data with a dynamic programming algorithm that performs approximate evaluation of all possible combinations of hierarchically consistent labels. However, this led to a slight decrease in performance, showing the advantage of considering only the biologically relevant combinations. Further research should consider other alternatives.

All experiments were performed on Linux running machines with 8 cores (64-bit, 3.3GHz) and 8GB memory. Combined running times for performing five-fold cross-validation for all three subontologies are: binary SVMs: 55 hours, Clus-HMC-Ens: 825 hours and PHENOstruct: 90 hours.

**Effectiveness of individual data sources**

We performed the following set of experiments in order to identify the most effective data sources for HPO prediction using PHENOstruct. First, to identify the individual effectiveness of each source, we performed a series of experiments in which we provided features generated from a single source of data at a time as input to PHENOstruct. Then to understand how much each data source is contributing to the overall performance we conducted leave-one-source-out experiments.

In all three subontologies, network data is the most informative individual data source as illustrated in Figure 7. Moreover, it is by far the main contributor to the overall performance both in the Organ and Inheritance subontologies (Figure 8). This is intuitive because if two genes/proteins are known to be interacting and/or active in the same pathways it leads to association with the same/similar diseases/phenotypes.

Although the genetic variant features provide the lowest performance in the Organ and Onset subontologies, leaving out variant data hurts the overall performance noticeably in all three subontologies as can be seen in Figure 8. This suggests that variant data are very

![Figure 7. Performance of PHENOstruct with individual data sources. Results are shown for each source of data: network (functional association data); Gene Ontology annotations; literature mining data; genetic variants; and the model that combines all features together.](image_url)
useful especially as a complementary dataset to the others. Moreover, we found that variant data are very effective for predicting cancer-related terms in the Organ subontology (see Table S1).

It is very encouraging to see that the literature data with a simple BoW representation by itself is very informative (Figure 7) and leaving out literature features shows considerable performance drop in the other two subontologies (Figure 8). In an analysis of the SSVM weight vector, we found that the majority of the most important tokens extracted from literature consist of names of proteins, genes and diseases (see Table S2).

We also considered an alternative representation of the literature data where a gene is represented by a vector in which the element $i$ gives the number of times the word $i$ occurred in the same sentence with that particular gene/protein divided by the total number of unique genes/proteins that word co-occurred with. This representation is analogous to the TFIDF (term frequency * inverse document frequency) representation typically used in information retrieval and text mining. However, these features led to slight deterioration of performance in all three subontologies (macro AUCs 0.60, 0.58 and 0.56 for Organ, Inheritance and Onset subontologies, respectively). Although GO features provide the second best individual performance both in the Organ and Onset subontologies (Figure 7), their contribution to the overall performance is very minimal (Figure 8). In fact leaving out GO features increases the overall performance in the Inheritance and Onset subontologies. The incompleteness of GO annotations may have contributed towards this.

Finally, the combination of all the features provides higher performance than individual feature sets in all three subontologies as can be
seen in Figure 7. However, leaving out GO features in the Inheritance and Onset subontologies, led to improved performance, suggesting that not all sources contribute to the overall performance. This shows that the selection of data sources must be performed carefully in order to find the optimal combination of sources for each subontology.

Validating false positives

Like other biological ontologies, the HPO is incomplete due to various factors such as slowness of the curation process\(^27\). In other words, the set of HPO annotations we considered as the gold standard does not fully represent all the phenotypes that should be associated with the currently annotated genes; this leads to performance estimates that underestimate the true performance of a classifier\(^21\). To explore this issue, we selected 25 predictions made by PHENOstruct which were considered false positives according to the current gold standard and looked for evidence in the current biomedical literature that can be used as evidence for those predictions. For 14 of those predictions we were able to find supporting evidence. The details of the complete validation process are given in the Supplementary material.

Conclusions and future work

This is the first study of directly predicting gene-HPO term associations. We modeled this problem as a hierarchical multi-label problem and used the SSVM framework for developing PHENOstruct. Our results demonstrate that using the SSVM is more effective than the traditional approach of decomposing the problem into a collection of binary classification problems. In our experiments we evaluated several types of data which were found to be informative for HPO term prediction: networks of functional association, large scale data mined from the biomedical literature and genetic variant data.

There are several ways in which this work can be extended. For the literature data we used a simple BoW representation. An alternative is to try and extract gene-HPO term co-mentions directly; in the context of GO term prediction we have found that both approaches lead to similar overall performance\(^17\). However, co-mentions have the added value that they are easy to verify by a human curator. Another source of information that can be utilized is semantic similarity of HPO terms to other phenotypic ontologies such the mammalian phenotype ontology, which is currently used for annotating the rat genome\(^28\). Finally, exploring the effectiveness of combining all three subontologies, as opposed to treating them as three independent subontologies as we have done here, is also worth exploring.

Although PHENOstruct outperformed the baseline methods, there is considerable room for improvement in all three subontologies. While some improvement can likely be obtained as described above, its performance will also improve as the number of HPO annotations increases. HPO is a relatively new ontology that will likely see substantial growth in the coming years, which will help in improving the accuracy of computational methods that contribute to its expansion.

Data and software availability

Zenodo: Data and software associated with PHENOstruct: Prediction of human phenotype ontology terms using heterogeneous data sources, 10.5281/zenodo.18764\(^29\)

Author contributions

IK and AB conceived and designed the method and experiments. CF and KV developed a NLP pipeline and generated literature features. IK performed all experiments with PHENOstruct. All authors read and approved the manuscript.

Competing interests

The authors declare that they have no competing interests.

Grant information

This work was supported by the NSF Advances in Biological Informatics program through grants number 0965768 (awarded to Dr. Ben-Hur) and 0965616 (originally awarded to Dr. Verspoor). I confirm that the funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.
Supplementary material

Analysis of variant features
In this section we analyze the performance of the features generated from genetic variant data in detail. The macro AUC for the variants data is only 0.56 in the Organ subontology. However, 37 terms have an AUC equal to or above 0.9. As listed in the Table S1, 21 out of those 37 terms well-predicted by the variant data are terms related to cancer. But interestingly, only 53 out of 1796 of all the Organ subontology terms are related to cancer. This shows strong evidence that the genetic variant data are highly effective for predicting cancer related phenotype terms. Terms predicted with high accuracy by the literature features do not show a similar tendency (data not shown).

| HPO ID   | HPO term                        | Freq | Depth | AUC  | Cancer-related |
|----------|---------------------------------|------|-------|------|----------------|
| HP:0006846 | Acute encephalopathy            | 15   | 5     | 1.00 | No             |
| HP:0006965 | Acute necrotizing encephalopathy | 15   | 6     | 1.00 | No             |
| HP:0003287 | Abnormality of mitochondrial metabolism | 16   | 4     | 1.00 | No             |
| HP:0008316 | Abnormal mitochondria in muscle tissue | 16   | 5     | 1.00 | No             |
| HP:0012103 | Abnormality of the mitochondrion | 16   | 3     | 1.00 | No             |
| HP:0002141 | Gait imbalance                   | 14   | 5     | 1.00 | No             |
| HP:000148  | Vaginal atresia                 | 19   | 7     | 1.00 | No             |
| HP:0001827 | Genital tract atresia           | 19   | 4     | 1.00 | No             |
| HP:0002862 | Bladder carcinoma               | 22   | 5     | 1.00 | Yes            |
| HP:0006740 | Transitional cell carcinoma of the bladder | 22   | 6     | 1.00 | Yes            |
| HP:0009725 | Bladder neoplasm                | 22   | 4     | 1.00 | Yes            |
| HP:0010784 | Uterine neoplasm                | 29   | 7     | 0.98 | Yes            |
| HP:002672 | Gastrointestinal carcinoma       | 23   | 6     | 0.98 | Yes            |
| HP:0006716 | Hereditary nonpolyposis colorectal carcinoma | 23   | 7     | 0.98 | Yes            |
| HP:0006749 | Malignant gastrointestinal tract tumors | 23   | 5     | 0.98 | Yes            |
| HP:0010747 | Medial flaring of the eyebrow    | 14   | 6     | 0.98 | No             |
| HP:0002891 | Uterine leiomyosarcoma           | 20   | 8     | 0.98 | Yes            |
| HP:0100243 | Leiomyosarcoma                   | 20   | 4     | 0.98 | Yes            |
| HP:0004481 | Progressive macrocephaly         | 18   | 6     | 0.97 | No             |
| HP:0007707 | Congenital primary aphaikia      | 14   | 7     | 0.97 | No             |
| HP:0100834 | Neoplasm of the large intestine  | 33   | 6     | 0.97 | Yes            |
| HP:0006519 | Alveolar cell carcinoma          | 13   | 6     | 0.97 | Yes            |
| HP:0100552 | Neoplasm of the tracheobronchial system | 13   | 5     | 0.97 | Yes            |
| HP:0009806 | Nephrogenic diabetes insipidus   | 15   | 3     | 0.95 | No             |
| HP:0100273 | Neoplasm of the colon            | 15   | 7     | 0.95 | Yes            |
| HP:0005584 | Renal cell carcinoma             | 28   | 6     | 0.94 | Yes            |
| HP:0003002 | Breast carcinoma                 | 20   | 3     | 0.94 | Yes            |
| HP:0006753 | Neoplasm of the stomach          | 39   | 5     | 0.94 | Yes            |
| HP:0100013 | Neoplasm of the breast           | 31   | 2     | 0.92 | Yes            |
| HP:0004808 | Acute myeloid leukemia           | 22   | 5     | 0.92 | No             |
| HP:0003006 | Neuroblastoma                    | 19   | 7     | 0.91 | Yes            |
| HP:0004376 | Neuroblastic tumors              | 19   | 6     | 0.91 | Yes            |
| HP:0002370 | Poor coordination                | 18   | 6     | 0.90 | No             |
| HP:0010786 | Urinary tract neoplasm           | 49   | 3     | 0.90 | Yes            |
| HP:0000142 | Abnormality of the vagina        | 27   | 6     | 0.90 | No             |
| HP:0001413 | Micronodular cirrhosis           | 14   | 5     | 0.90 | No             |
| HP:0009726 | Renal neoplasm                   | 47   | 5     | 0.90 | Yes            |

Terms are listed in the ascending order of their individual AUCs. 21 out of the 37 (57%) terms well-predicted Organ subontology terms by the variant data are terms related to cancer.
In the Inheritance subontology it was noticeable that the variant data are more effective for the categories with fewer annotations compared to the literature features (data not shown). The average number of annotations of the Inheritance subontology HPO categories with relatively higher AUC by variant features (compared to literature features) is only 46. This trend is also visible, albeit to a lesser extent, in the Organ and Onset subontologies as well; the corresponding numbers for the Organ and the Onset subontologies are 26 and 91, respectively. Furthermore, only mitochondrial inheritance term achieves an AUC above 0.9 with all data sources. However, with variant data alone, both mitochondrial inheritance and somatic mutation terms achieve AUCs above 0.9.

**Analysis of literature features**

In order to identify the most important literature features, we looked at the weight vectors of the structured SVM model underlying PHENOstruct that was trained only on the literature features. Typically, the input space features with higher weight in the weight vector correspond to the features that are considered most important by the model for the given predictive task.

In the dual formulation of the Structured SVM, \( \alpha \) values are defined for each pair of example \( i \) and structured output \( j \). In order to calculate the weight vector for a specific HPO term \( j \) (\( W_j \)), we first identified the subset of input examples (i.e. proteins) that are annotated with the given term (referred to as \( S_j \)). Then \( W_j \) is the summation of \( \alpha_{ij} \times x_i \) where \( x_i \) is the feature vector of example \( i \) and \( x_i \in S_j \). Features with higher weights in the weight vector \( W_j \) correspond to the features that were considered most informative by the model for the task of predicting the term \( j \).

We trained PHENOstruct on literature features and computed the weight vectors as described above. Then we ranked the literature features by their weights and examined the top-100 literature features. In the Organ subontology we analysed the top-100 literature features with respect to the 8 HPO terms that have individual AUCs above 0.9. For those 8 terms the union set of top-100 features is composed of 107 unique tokens. By far, the majority (>70%) of these tokens are genes/proteins/protein complexes/pathways names. Another 12 tokens are disease/phenotype names (Table S2).

| Category                        | Tokens                                                                                                                                                                                                 |
|---------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| proteins/protein complexes      | cx32, kissepeptin, -308, t308, smn2, ns5, trap-positive, mpp+-induced, 1-methyl-4-phenylpyridinium, tfn-alpha-mediated, tfn-alpha-stimulated, tfn-alpha-treated, frataxin, ki-ras, connexin32, tdcb, recq4, =-galcer, tyrosinase-related, hpms2, her4, cd40-cd40l, imp2a, rys, mg2+-atpase, ews-fl1, abeta42, fancc, p40phox, her1, bdnf-induced, trap+, gfap-ir, daf-16/foxo, hdl3, -238, [tfn-alpha], cd40/cd40l, anti-ngf, tep1, recq, rt-4, pfemamp1, zo-2, nphp1, tfn-alpha-dependent, pomt1, igm-positive, apoai, p110alpha, fancf, tbx4, anti-cd40l, igg |
| genes                           | hmsn2, cx26, fkrp, smn1, cin3, nphp4, mn1, nnt, apex2, akt-2                                                                                                                                              |
| pathways                        | ras/raf/mek/erk, pi3k-akt-mtor                                                                                                                                                                           |
| diseases/phenotypes             | cmt1a, hnp, hdl2, cin2, hpp, fmf, rtt, hnpcc, charcot-marie-tooth, amenorrhea, rett, antiiardioiipin                                                                                                          |
| misc.                           | sheldrick, shelx97, bruker, farrugia, ortep-3, platon, shelxs97, spek, sgdid, wlds, caii, aoa, tdf, crysalis, wingx, amf                                                                                                                                 |

The union set of the top-100 literature features with respect to the 8 HPO terms that have individual AUCs equal to or above 0.9 is composed of 107 unique tokens. The token “-308” and “t308” in the “proteins/protein complexes” category are due to mis-tokenization of “miR-308”. Similarly, “-238” in the same category is due to mis-tokenization of “BG-23”. Also “=-galcer” in the same category originated from \( \alpha \)-galcer and \( \beta \)-galcer due to mishandling of UTF characters \( \alpha \) and \( \beta \).
Additional methods
We describe here the results of experiments that we conducted with two additional methods.

**SSVM → disease → HPO method** This is an indirect method that first predicts gene-disease associations and then maps them to HPO terms using associations available on the HPO website. This method uses the same input space data as PHENOstruct and learns a structured SVM using the same methodology. Using this model it predicts diseases along with confidence scores for unseen genes. Subsequently, the predicted scores for disease terms are directly transferred to all the HPO terms associated with those diseases (Figure S1). When multiple diseases are associated with a single HPO term, scores are accumulated. It is surprising that this method shows mediocre performance (Table S3). One of the main reasons for this is the low performance of the underlying SSVM for predicting disease terms (average AUC of 0.64), which consequently affects the accuracy of predicted HPO terms.

**PhenoPPIOrth** We also evaluated the performance of PhenoPPIOrth (Wang et al., 2013). PhenoPPIOrth is a computational tool that can predict a set of diseases for a given human gene. Specifically, it predicts OMIM disease terms for human genes using protein-protein interaction and orthology data. Then it also maps the predicted OMIM terms to HPO terms using the disease-HPO mapping available in the HPO website11. We downloaded the pre-computed predictions from the PhenoPPIOrth website. Compared to PHENOstruct, PhenoPPIOrth’s performance was quite low (see Table S3). It is important to note that PhenoPPIOrth makes predictions for only a subset of proteins with respect to all three ontologies (1487, 175 and 155 in Organ, Inheritance and Onset subontologies, respectively). One of the main reasons is that HPO annotations are generated using three sources: OMIM, Orphanet and DECHIPER but PhenoPPIOrth predicts only OMIM terms.

---

**Figure S1. SSVM → disease → HPO method.** This method takes feature vectors and disease annotations associated with each gene as the input for training a SSVM model. Then, it predicts diseases for unseen genes using the learned model. Subsequently, the predicted scores for disease terms are directly transferred to all the HPO terms associated with those diseases.

---

**Table S3. Comparison between PHENOstruct and other methods.**

| Subontology | Method            | F-max | Precision | Recall | mac-AUC |
|-------------|-------------------|-------|-----------|--------|---------|
| Organ       | PhenoPPIOrth      | 0.20  | 0.27      | 0.15   | 0.52    |
|             | Struct->Dis->HPO  | 0.23  | 0.16      | 0.41   | 0.49    |
|             | Binary SVMs       | 0.35  | 0.32      | 0.40   | 0.66    |
|             | Clus-HMC-Ens      | 0.41  | 0.39      | 0.43   | 0.65    |
|             | PHENOstruct       | 0.42  | 0.35      | 0.56   | 0.73    |
| Inheritance | PhenoPPIOrth      | 0.12  | 0.16      | 0.10   | 0.55    |
|             | Struct->Dis->HPO  | 0.11  | 0.07      | 0.25   | 0.46    |
|             | Binary SVMs       | 0.69  | 0.62      | 0.78   | 0.72    |
|             | Clus-HMC-Ens      | 0.73  | 0.64      | 0.84   | 0.73    |
|             | PHENOstruct       | 0.74  | 0.68      | 0.81   | 0.74    |
| Onset       | PhenoPPIOrth      | 0.25  | 0.25      | 0.24   | 0.53    |
|             | Struct->Dis->HPO  | 0.07  | 0.06      | 0.10   | 0.49    |
|             | Binary SVMs       | 0.33  | 0.24      | 0.51   | 0.62    |
|             | Clus-HMC-Ens      | 0.35  | 0.27      | 0.48   | 0.58    |
|             | PHENOstruct       | 0.39  | 0.31      | 0.52   | 0.64    |

The performance is evaluated using macro AUC and protein-centric F-max, Precision and Recall (as defined above) on the complete HPO graph (i.e. true-path rule is applied to annotations and predictions).
Table S4. Comparison between PHENOstruct vs. other methods only leaf terms.

| Subontology | Method          | F-max | Precision | Recall | mac-AUC |
|-------------|-----------------|-------|-----------|--------|---------|
| Organ       | PhenoPPIOrth    | 0.03  | 0.05      | 0.03   | 0.51    |
|             | Struct->Dis->HPO| 0.01  | 0.01      | 0.02   | 0.50    |
|             | Binary SVMs     | 0.20  | 0.19      | 0.22   | 0.66    |
|             | Clus-HMC-Ens    | 0.08  | 0.04      | 0.31   | 0.64    |
|             | PHENOstruct     | 0.30  | 0.21      | 0.50   | 0.77    |
| Inheritance | PhenoPPIOrth    | 0.11  | 0.15      | 0.09   | 0.55    |
|             | Struct->Dis->HPO| 0.01  | 0.01      | 0.02   | 0.46    |
|             | Binary SVMs     | 0.69  | 0.62      | 0.74   | 0.71    |
|             | Clus-HMC-Ens    | 0.72  | 0.63      | 0.84   | 0.73    |
|             | PHENOstruct     | 0.74  | 0.68      | 0.82   | 0.74    |
| Onset       | PhenoPPIOrth    | 0.19  | 0.20      | 0.17   | 0.52    |
|             | Struct->Dis->HPO| 0.03  | 0.02      | 0.14   | 0.49    |
|             | Binary SVMs     | 0.29  | 0.21      | 0.47   | 0.62    |
|             | Clus-HMC-Ens    | 0.28  | 0.21      | 0.43   | 0.58    |
|             | PHENOstruct     | 0.35  | 0.28      | 0.48   | 0.68    |

The performance is evaluated by using the exact annotations (i.e. only leaf terms) as ground truth. In other words, true-path rule is not applied. Performance is presented using macro AUC and protein-centric F-max, Precision and Recall as defined above.

**Performance measures**

We use term-centric AUC or macro AUC as our primary evaluation measure for reporting results. In addition, we use several protein-centric measures. Protein-centric precision and recall at a given threshold \( t \) are defined as

\[
Pr_{pc}(t) = \frac{1}{N} \sum_{i=0}^{N} \frac{TP(t)_i}{TP(t)_i + FP(t)_i},
\]

\[
Rec_{pc}(t) = \frac{1}{N} \sum_{i=0}^{N} \frac{TP(t)_i}{TP(t)_i + FN(t)_i},
\]

where \( TP(t)_i \), \( FP(t)_i \) and \( FN(t)_i \) are the number of true positives, number of false positives and number of false negatives w.r.t. protein \( i \) at threshold \( t \). Now we can define protein-centric F-max as

\[
F_{pc} - max = \max_t \frac{2Pr_{pc}(t)Rec_{pc}(t)}{Pr_{pc}(t) + Rec_{pc}(t)}.
\]

**Complete results**

In this section we present performance of all five methods using several performance measures.

**Validating false positives**

First we ranked the test proteins in descending order of the protein-centric precision of their Organ subontology predictions made by PHENOstruct. Then we retrieved the 25 false positive predictions for the top 17 proteins in that list. Next, we performed online searches using the pair of protein name and phenotype name as the query for the search engine. This resulted in a list of publications for each false positive prediction. Then we manually extracted the excerpts from those papers that contained supporting evidence that suggests the particular false positive is in fact correct. Using this manual process we found evidence for 14 of the 25 false predictions considered for this study (see Table S5). For two of the cases the evidence comes from studies involving mice (indicated within parentheses with the PubMed ID). Overall success of this study strongly suggests that the performance of PHENOstruct is under-estimated due to the incompleteness of the current gold standard.
| Gene     | HPO term                    | PubMed ID               | Evidence                                                                                   |
|----------|-----------------------------|-------------------------|-------------------------------------------------------------------------------------------|
| DKC1     | Postnatal growth retardation| PMID: 1085321            | "...Hoyeraal-Hreidarsson (HH) syndrome is a multisystem disorder affecting boys characterized by aplastic anaemia (AA), immunodeficiency, microcephaly, cerebellar hypoplasia and growth retardation... We therefore analysed the DKC1 gene in two HH families. One family has a nucleotide change at position 146G > T in exon 3 of the DKC1 gene..." |
| PEX13    | Retinal dystrophy           | PMID: 12897163          | "...Autosomal-dominant Retinitis pigmentosa... These results strongly suggest that causal mutations in PEX13 are responsible for retinal dystrophy in these affected individuals..." |
| PEX5     | Retinal dystrophy           | PMID: 19878916          | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in PEX5 causes impairment of pre-mRNA splicing and autosomal-dominant retinitis pigmentosa..." |
| BEST1    | Retinitis pigmentosa        | PMID: 19878916          | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in BEST1 causes retinitis pigmentosa..." |
| RPE65    | Retinitis pigmentosa        | PMID: 21549338          | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in RPE65 causes impairment of pre-mRNA splicing and autosomal-dominant retinitis pigmentosa..." |
| SNRNP200 | Emphysema                   | PMID: 21549338          | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in SNRNP200 causes impairment of pre-mRNA splicing and autosomal-dominant retinitis pigmentosa..." |
| ORC1     | Emphysema                   | PMID: 21549338          | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in ORC1 causes impairment of pre-mRNA splicing and autosomal-dominant retinitis pigmentosa..." |
| CUL7     | Postnatal microcephaly      | PMID: 21549338          | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in CUL7 causes impairment of pre-mRNA splicing and autosomal-dominant retinitis pigmentosa..." |
| ACTB     | Progressive hearing impairment | PMID: 21549338   | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in ACTB causes impairment of pre-mRNA splicing and autosomal-dominant retinitis pigmentosa..." |
| MSH2     | Maligant gastrointestinal tract tumours | PMID: 21549338 | "...Autosomal-dominant retinitis pigmentosa... A missense mutation in MSH2 causes impairment of pre-mRNA splicing and autosomal-dominant retinitis pigmentosa..." |

*The columns “HPO term”, “PubMed ID” and “Evidence” provides the false positive prediction made by PHENOStruct for the given gene, PubMed ID of the literature that contains evidence which actually suggests that the prediction should be considered true and the excerpt from that literature which contains the evidence, respectively. We used the 25 false positive predictions for the 17 proteins that had the highest individual protein-centric precision and found evidence for 14 predictions. Two of the evidence comes from studies involving mice (indicated within parentheses with the PubMed ID)."
Table S6. Performance of Binary SVMs in the Inheritance subontology.

| Name                                      | Freq | Depth | AUC  |
|-------------------------------------------|------|-------|------|
| Multifactorial inheritance                | 15   | 1     | 0.62 |
| Polygenic inheritance                     | 15   | 2     | 0.62 |
| Mitochondrial inheritance                 | 41   | 1     | 0.96 |
| Sporadic                                  | 52   | 1     | 0.66 |
| Somatic mutation                          | 61   | 1     | 0.71 |
| X-linked dominant inheritance             | 62   | 3     | 0.79 |
| X-linked recessive inheritance            | 111  | 3     | 0.70 |
| Heterogeneous                             | 148  | 1     | 0.65 |
| X-linked inheritance                      | 198  | 2     | 0.78 |
| Gonosomal inheritance                     | 198  | 1     | 0.78 |
| Autosomal dominant inheritance            | 1096 | 1    | 0.69 |
| Autosomal recessive inheritance           | 1665 | 1    | 0.68 |

The macro AUC for the Inheritance subontology is 0.72. Terms are displayed in ascending order of frequency.

References

1. Robinson PN: Deep phenotyping for precision medicine. Hum Mutat. 2012; 33(5): 777–780. PubMed Abstract | Publisher Full Text
2. Kihara S, Doekken SC, Mungall CJ, et al.: The human phenotype ontology project: linking molecular biology and disease through phenotype data. Nucleic Acids Res. 2014; 42(Database issue): D966–D974. PubMed Abstract | Publisher Full Text | Free Full Text
3. Hamosh A, Scott AF, Amberger JS, et al.: Online Mendelian Inheritance in Man (OMIM), a knowledgebase of human genes and genetic disorders. Nucleic Acids Res. 2005; 33(Database issue): D514–D517. PubMed Abstract | Publisher Full Text | Free Full Text
4. Ayme S, Schmitz-Essen D, Schumacher H, et al.: Networking for rare diseases: a necessity for Europe. Bundesgesundheitsblatt Gesundheitsforschung Gesundheitsschutz. 2007; 50(12): 1477–1483. PubMed Abstract | Publisher Full Text
5. Bragin E, Chatzimichali EA, Wright CF, et al.: DECIPHER: database for the interpretation of phenotype-linked pathogenic sequence and copy-number variation. Nucleic Acids Res. 2014; 42(Database issue): D993-D1000. PubMed Abstract | Publisher Full Text | Free Full Text
6. Ashburner M, Ball CA, Blake JA, et al.: Gene ontology: tool for the unification of biology. The Gene Ontology Consortium. Nat Genet. 2000; 25(1): 25–29. PubMed Abstract | Publisher Full Text | Free Full Text
7. Robinson PN, Köhler S, Gelrich A, et al.: Improved exome prioritization of disease genes through cross-species phenotype comparison. Genome Res. 2014; 24(2): 340–348. PubMed Abstract | Publisher Full Text | Free Full Text
8. Moreau T, Tranchevent LC: Computational tools for prioritizing candidate genes: boosting disease gene discovery. Nat Rev Genet. 2012; 13(8): 523–536. PubMed Abstract | Publisher Full Text
9. Bi W, Kwok JT: Multi-label classification on tree- and dag-structured hierarchies. In Lise Getoor and Tobias Scheffer, editors, New York, NY, USA, ACM. Proceedings of the 28th International Conference on Machine Learning (ICML-11). 2011: 17–24. Reference Source
10. Silla CN Jr, Freitas AA: A survey of hierarchical classification across different application domains. Data Min Knowl Discov. 2011; 22(1-2): 31–72. PubMed Abstract | Publisher Full Text
11. Obozinski G, Lankriet G, Grant C, et al.: Consistent probabilistic outputs for protein function prediction. Genome Biol. 2008; 9(Suppl 1): S6. PubMed Abstract | Publisher Full Text | Free Full Text
12. Tschantz tantalidis I, Joachims T, Hofmann T, et al.: Large margin methods for structured and interdependent output variables. J Mach Learn Res. 2005; 6: 1453–1484. Reference Source
13. Sokolov A, Ben-Hur A: Hierarchical classification of gene ontology terms using the GOstruct method. J Bioinform Comput Biol. 2010; 8(2): 357–376. PubMed Abstract | Publisher Full Text
14. Sokolov A, Funk C, Gramm K, et al.: Combining heterogeneous data sources for accurate functional annotation of proteins. BMC Bioinformatics. 2013; 14(Suppl 3): S10. PubMed Abstract | Free Full Text
15. Chatr-aryamontri A, Breitkreutz BJ, Heinicke S, et al.: The BioGRID Interaction database: 2013 update. Nucleic Acids Res. 2013; 41(Database issue): D816–D823. PubMed Abstract | Publisher Full Text | Free Full Text
16. Szklarczyk D, Franceschini A, Kuhn M, et al.: The STRING database in 2011: functional interaction networks of proteins, globally integrated and scored. Nucleic Acids Res. 2011; 39(Database issue): D561–8. PubMed Abstract | Publisher Full Text | Free Full Text
17. Funk C, Kahanda I, Ben-Hur A, et al.: Evaluating a variety of text-mined features for automatic protein function prediction with GOstruct. J Biomed Semantics. 2015; 6: 9. PubMed Abstract | Publisher Full Text | Free Full Text
18. Wang P, Lai WF, Li MJ, et al.: Inference of gene-phenotype associations via protein-protein interaction and orthology. PLoS One. 2013; 8(10): e77478. PubMed Abstract | Publisher Full Text | Free Full Text
19. Schegt L, Vens C, Stroy J, et al.: Predicting gene function using hierarchical multi-label decision tree ensembles. BMC Bioinformatics. 2010; 11: 2. PubMed Abstract | Publisher Full Text | Free Full Text
20. Radijovic P, Clark WT, Orton TR, et al.: A large-scale evaluation of computational protein function prediction. Nat Methods. 2013; 10(3): 221–227. PubMed Abstract | Publisher Full Text | Free Full Text
21. Huntenhower C, Hibbs MA, Myers CL, et al.: The impact of incomplete knowledge on evaluation: an experimental benchmark for protein function prediction. Bioinformatics. 2009; 25(18): 2404–2410. PubMed Abstract | Publisher Full Text | Free Full Text
22. Anderson S, Bankier AT, Barrett BG, et al.: Sequence and organization of the human mitochondrial genome. Nature. 1981; 290(5806): 457–465. PubMed Abstract | Publisher Full Text | Free Full Text
23. Taylor RW, Turnbull DM: Mitochondrial DNA mutations in human disease. Nat Rev Genet. 2005; 6(5): 389–402. PubMed Abstract | Publisher Full Text | Free Full Text
24. Wallace DC, Singh G, Lott MT, et al.: Mitochondrial DNA mutation associated with Leber’s hereditary optic neuropathy. Science. 1988; 242(4884):1427–1430. PubMed Abstract | Publisher Full Text

25. Holt U, Harding AE, Morgan-Hughes JA: Deletions of muscle mitochondrial DNA in patients with mitochondrial myopathies. Nature. 1988; 331(6158):717–719. PubMed Abstract | Publisher Full Text

26. Jones KS: A statistical interpretation of term specificity and its application in retrieval. J Doc 1972; 28(1): 11–21. Publisher Full Text

27. Baumgartner WA Jr, Cohen KB, Fox LM, et al.: Manual curation is not sufficient for annotation of genomic databases. Bioinformatics. 2007; 23(13): i41–i48. PubMed Abstract | Publisher Full Text | Free Full Text

28. Smith CL, Eppig JT: The mammalian phenotype ontology: enabling robust annotation and comparative analysis. Wiley Interdiscip Rev Syst Biol Med. 2009; 1(3): 390–399. PubMed Abstract | Publisher Full Text | Free Full Text

29. Kahanda I, Funk C, Verspoor K, et al.: Data and software associated with PHENOstruct: Prediction of human phenotype ontology terms using heterogeneous data sources. Zenodo. 2005. Data Source
Open Peer Review

Current Referee Status: ✔️ ✔️

Version 1

Referee Report 28 August 2015
doi:10.5256/f1000research.7166.r9995

Shaillay Dogra
Vishuo BioMedical Pte Ltd, Singapore, Singapore

A well written article with detailed methodology towards mapping genes to diseases. The method proposes to overcome the limitations of traditional approaches, which take single-label at a time. Author's approach uses structured prediction that takes into account related set of labels.

Some general points for authors consideration:
1. What is a possible, direct application of this method? How can this be integrated with other software tools or called as a service, for example?

2. Different data sources have been used of which protein-protein interactions, for example, is a noisy one with false-interactions reported via experimental methods like co-precipitation and yeast-two-hybrid systems. Can the author's comment on how the quality of data affects their approach? How to deal with noisy data sources or reduce the weight of contribution from that particular data source?

3. How easy is to update this, for example, integrating latest PubMed abstracts? What's the pipeline or process to do so?

Some specific points for authors to consider:
1. Why do we expect more genes to be disease-causing? Is it just a general line of reasoning that functional genes should cause an aberrant phenotype if they do not work properly?

2. Perhaps the authors can expand a bit more on the problem formulation - "therefore, it is important to explore the feasibility..."

3. Under "Approach", perhaps the authors can explain a bit on HMC for the benefit of the readers. Also, a bit more on "structured prediction/learning" in layman terms or illustrated with an example can help the reader grasp the concept.

4. Authors can consider simplifying this into shorter sentences for easy grasp - "An alternate approach is to use a single classifier..."

5. Under "HPO annotations" - for general understanding, could the authors tell more about why they 'removed terms that were not annotated to 10 or more genes.'
6. Under "Literature" - abstracts extracted are from 2013 and not up to date with 2015.

7. Under "Literature" - for general understanding, could the authors tell more about why they 'filtered to keep only the low frequency words'.

8. Under "variants" - does this data from Uniprot covers data sources like clinvar, dbGaP, GWAS studies etc?

9. Under "Models", for the general understanding of the reader could the authors expand on what is meant by a structured model?

10. Figure 3 - last part of panel is barely legible.

11. Under "Evaluation" could the authors expand on what's implied by term-centric and protein-centric, F-max?

12. "The human mitochondrial DNA... Due to this, and the relative..." - I am not sure if this is the reason.

13. Authors can consider simplifying this into shorter sentences for easy grasp - "As a potential improvement to PHENOstruct..."

14. I really liked the way authors dealt with "validating false positives" (text and table S5).

15. Figure S1 - last part of panel is almost illegible.

16. Supplementary material, "Performance Measures" - for the benefit of the readers, what does F-max mean in a literal, intuitive sense?

17. Table S4 -- for the benefit of the readers, what is 'true-path rule', 'ground truth', 'macro AUC'...?

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.
term/Gene association predictions one at a time. If a prediction is YES for "ventricular septal defect" but NO for "abnormality of the ventricular septum", then the result is mutually inconsistent. A number of machine learning algorithms have emerged to tackle this problem, including one that the authors previously learned for an analogous project with Gene Ontology predictions. In general, the paper is very well done and it is likely to be accessible to a wide audience because it is well written. The topic of HPO annotation prediction is very new, and there is no other published work on the topic that I am aware of at present, although a number of groups, including the authors, participated in a CAFA competition at the 2014 ISMB.

Suggestions:
1. The authors should cite GeneMania (The GeneMANIA prediction server: biological network integration for gene prioritization and predicting gene function Warde-Farley et al, 2010, NAR).

2. The authors should state the version of the HPO and the HPO annotation data they used. In the meantime, the number of annotations has increased substantially, and a number of improvements to the HPO structure have been made (for instance, the Organ abnormality term has been renamed to Phenotypic abnormality).

3. It would be nice to have a little more self-contained explanation about some of the methods employed, such as Clu-HMC-Ens.

4. The authors observed an excellent AUROC score for mitochondrial inheritance, and speculate that the reason is that the mitochondrial genome is well studied. I suspect that the true reason might be that mitochondrial genes have a very specialized functional profile (energy etc) that is much more homogeneous than say "autosomal recessive".

5. The Mammalian phenotype ontology is not only used to annotate the rat genome, but is the major tool used to annotate the mouse genome, and is an extremely useful resource used now by the International Mouse Phenotyping Consortium the Mouse Genome Informatics group, and many others. This should be added to the text.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.
Author/s: Kahanda, I; Funk, C; Verspoor, K; Ben-Hur, A

Title: PHENOstruct: Prediction of human phenotype ontology terms using heterogeneous data sources.

Date: 2015

Citation: Kahanda, I., Funk, C., Verspoor, K. & Ben-Hur, A. (2015). PHENOstruct: Prediction of human phenotype ontology terms using heterogeneous data sources. F1000Res, 4, pp.259-. https://doi.org/10.12688/f1000research.6670.1.

Persistent Link: http://hdl.handle.net/11343/127454

File Description: Accepted version

License: CC BY