Development and Application of an Interaction Network Ontology (INO) for Literature Mining of Vaccine-associated Gene-Gene Interactions

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Abstract — Literature mining of gene-gene interactions has been enhanced by ontology-based name classifications. However, in biomedical literature mining, interaction keywords have not been carefully studied and used beyond a collection of keywords. In this study, we report the development of a new Interaction Network Ontology (INO) that classifies >800 interaction keywords and incorporates interaction terms from the PSI Molecular Interaction (PSI-MI) and Gene Ontology (GO). Using INO-based literature mining results, a modified Fisher's exact test was established to analyze significantly over- and under-represented enriched gene-gene interaction types within a specific area. Such a strategy was applied to study the vaccine-mediated gene-gene interactions using all PubMed abstracts. The Vaccine Ontology (VO) and INO were used to support the retrieval of vaccine terms and interaction keywords from the literature. In total 14 over-represented and 17 under-represented interaction types were identified. The analysis of these interaction types and their associated gene-gene pairs uncovered many scientific insights.

Keywords—ontology; vaccine; Interaction Network Ontology; INO; literature mining; enrichment analysis

I. INTRODUCTION

Two common strategies of literature retrieval of reported gene-gene interactions include gene-gene co-occurrence and interaction keywords-based literature mining. In this paper, the gene-gene interaction represents a broad interactive relation between two genes or gene products [1]. Such a relation does not have to be direct physical interaction. The co-occurrence strategy identifies two related genes both listed in the same literature, or more specifically in the same title, abstract, or sentence. An example of such a strategy is the PubGene system that extracts gene relationships based on co-occurrence of gene symbols in MEDLINE titles and abstracts [2]. The other strategy relies on the identification of two genes together with an interaction keyword in the same sentence. Such a method may still generate many false-positive results. To improve the interaction keyword-based approach machine learning algorithms (e.g., support vector machine (SVM) [3]) with features extracted from syntactic analysis of sentences (e.g., dependency parse trees) can be used [4].

Ontologies can be applied to enhance literature mining performance. For example, in our previous work, a vaccine-specific sub-network was built by considering only the interactions that were extracted from sentences that contain the “vaccine” term (or its variants like “vaccines”, “vaccination”, and “vaccinated”). This strategy does not retrieve the sentences where more specific vaccine names such as BCG (a commercial tuberculosis vaccine) are mentioned. Such vaccine names and their hierarchical relations are represented in Vaccine Ontology (VO) [5]. We found that the application of VO has significantly improved the analysis of the vaccine-specific sub-network [6].

An ontology that logically represents various interaction keywords/types and their semantic relations would help address the challenge of retrieving and classifying the types of gene-gene interactions in the interaction keyword-based literature mining. The GENIA ontology provides a semantically annotated corpus for biological literature mining [7]. However, this ontology does not specify various types of interactions between genes or proteins. Initiated from the classification of >800 interaction keywords [6], we have developed the Interaction Network Ontology (INO) that ontologically represents various interaction types and their relations, and collects and assigns interaction keywords to these different interaction types. The detail about the ontology will for the first time be provided in this manuscript.

In addition to supporting the literature mining of gene-gene interactions, INO can be used for interaction type enrichment analysis. Gene Ontology (GO)-based gene set enrichment analyses have been widely used to determine over- or under-represented biological functions in a set of genes obtained from high-throughput Omics studies. GO provides controlled vocabulary of standard terms for describing gene product characteristics in a hierarchical structure. The input for the GO term enrichment analysis is a list of genes. Such a method does not classify enriched gene-gene interactions. Since INO classifies different interaction types into a structured ontology, it becomes possible to perform gene-gene interaction enrichment study by comparing the INO-based literature mined data of gene-gene interactions in some specific domain over the data from the broad background literature mining.

In this manuscript, we will first introduce the INO development with a focus on its representation of interaction types and keywords for literature mining. An INO-based gene interaction enrichment method based on a modified Fisher’s exact test will then be introduced. Such a method has been applied to analyze the vaccine-mediated gene-gene interactions.
interactions. The results of over- and under-represented gene-gene interaction types and gene-gene interactions will also be described in detail.

II. METHODS

A. INO development

INO was developed by following the Open Biological Ontology (OBO) Foundry ontology development principles, including openness and collaboration [8]. Its development is aligned and integrated with existing OBO Foundry library ontologies. INO imports existing terms by using OntoFox [9]. New terms generated in INO use the “INO_” prefix. INO uses the format of W3C standard Web Ontology Language (OWL2) (http://www.w3.org/TR/owl-guide/). For efficient editing of INO, the Protégé 4.3 OWL ontology editor (http://protege.stanford.edu/) was used.

The INO source is open freely under a Creative Commons (CC) license for public and commercial usage. INO has been deposited at the INO SourceForge project page (http://sourceforge.net/projects/ino/). It is available in the ontology repositories of NCBO BioPortal (http://purl.bioontology.org/ontology/INO) and Ontobee [10] (http://www.ontobee.org/browser/index.php?o=INO).

B. INO-based literature mining of gene-gene interaction pairs and interaction types

The sentences from the complete PubMed abstracts (published up to 12/31/2013) were obtained from the BioNLP database in the National Center for Integrative Biomedical Informatics (NCIBI; http://ncibi.org/). VO-SciMiner [11] was used to identify gene names/symbols, VO and INO terms from these sentences. Sentences with two gene names and INO terms (e.g., interacts, binds, activates) were selected. We obtained the dependency parse trees of the sentences using the Stanford Parser [12] and extracted the shortest dependency path between each pair of genes in a sentence. We defined an edit distance-based kernel function among these dependency paths and used SVM [3] to classify if a path describes an interaction between a gene pair [6]. A confidence score calculated based on SVM was used to measure the confidence of association between two genes in a sentence in the literature. Positively-scored sentences were kept, and the gene pairs together with the interaction keywords from these sentences were extracted. The extracted interaction keywords were mapped to INO to define the interaction types.

C. Development of INO-based statistical enrichment analysis of literature mined gene-gene interaction data

A modified Fisher’s exact test has gained popularity over the last decade in high-throughput gene expression studies as a preferred method for identifying enriched biological functions among given gene sets [13, 14]. We implemented the modified Fisher’s exact test in Perl using the Ngram Statistics Package [15] to identify enriched gene-gene interaction types, in terms of INO terms, within a concept-specific sub-network. For each INO term, a 2x2 contingency table is obtained on which the Fisher’s test runs, as shown in Table 1. Both under-represented and over-represented terms are selected as a significantly enriched INO term with a p-value < 0.05 after Benjamini-Hochberg (BH) multiple testing corrections. In the current study, vaccine-associated gene-gene interaction network was defined based on the gene-gene interactions obtained from the PubMed abstracts, including those retrieved by a PubMed search of ’vaccine’ and those identified by SciMiner-VO using 186 specific vaccine terms extracted from the VO ‘vaccine’ branch. These 186 vaccine terms (e.g., tuberculosis vaccine BCG) are easily identified by natural language processing programs. This vaccine-associated network was compared against the complete gene-gene interaction network.

Table 1. 2x2 contingency table.

| # of gene-gene Interaction pairs | Concept-specific sub-network | Whole Network |
|----------------------------------|-------------------------------|---------------|
| With the INO term                | 30 – 1                        | 500           |
| Without the INO term             | 150                           | 30000         |

The sub-network has 30 gene pairs associated with this INO term out of a total of 180 gene pairs. A modified Fisher's exact test, with the “-1” modification made to the typical Fisher’s exact test to make the statistical test more conservative, was employed to identify significantly over-represented terms (p-value of 6.9E-20).

III. RESULTS

A. The Interaction Network Ontology (INO)

1) INO overall design and hierarchy

INO is a biomedical ontology in the domain of molecular interactions and interaction networks. INO is aligned with the upper level Basic Formal Ontology (BFO) [16] (Figure 1). BFO contains two branches, continuant and occurrent. The continuant branch represents time-independent entity such as material entity, and the occurrent branch represents time-related entity such as process. BFO has currently been used by over 100 domain ontologies, including many (e.g., GO) within the framework of the OBO Foundry [8]. By aligning different domain ontologies under the two branches of BFO, INO is able to efficiently use the terms from other ontologies in representing signaling pathway elements.

Three key INO terms are interaction, network, and pathway. In INO, an interaction is defined as a processual entity that has two or more participants (i.e., interactors) that have an effect upon one another under a particular condition. An interactor (or called interactant) is defined as a material entity that plays the role of “interactor role”. With different roles, an interactor can be an ‘input interactor’, ‘output interactor’, ‘catalyst’, ‘positive regulator’, or ‘negative regulator’. An interaction consumes its input interactors (but not the catalysts or regulators) and generates its output interactors. A network is a process that includes at least two connected interactions. A network does not have to include a predefined start or end entity. A pathway is a type of network that has specified distinct start(s) and end(s). Each of the three INO key terms includes many subclasses.
INO imports terms from other ontologies, particularly from the Proteomics Standard Initiative-Molecular Interaction (PSI-MI), which is a standard molecular interaction data exchange format established by the HUPO Proteomics Standard Initiative (http://www.psidev.info). Their PSI-MI format has been widely used in the proteomics community and PSI-MI is also an OBO Foundry library ontology. To be compatible with PSI-MI, we have imported the branch of the ‘interaction type’ (MI_0190) to INO (Figures 1 and 2).

Compared to PSI-MI, GO Biological Processes (BP) branch often has more detailed subclasses (or subtypes) to specific interaction types. Using more general PSI-MI terms (e.g., PSI-MI ‘lipid addition’) as parent terms, INO has imported many specific GO subtypes of interactions (e.g., GO ‘protein myristoylation’) to INO as subclasses of the MI-based interaction terms (Figure 1). As a specific example, we have imported GO ‘protein myristoylation’ and all of its GO subclasses to INO (Figure 2). The GO term ‘protein myristoylation’ has been used to replace the PSI-MI term ‘myristoylation reaction’. It is noted that the top level GO Biological Processes hierarchy is not used because many biological processes (e.g., ‘metabolic process’) in GO are not ‘interaction’ per se and thus cannot be imported to INO for interaction representation.

While PSI-MI focuses on direct protein-protein interactions, it does not include many other interaction types such as regulation types. Therefore, INO also includes interaction types that are out of current PSI-MI scope, especially different regulation types (Figure 1). Many of these interaction types were generated by classifying the over 800 interaction keywords used in our previous literature mining studies [1, 6].

### (2) Literature mining support in INO

The over 800 interaction keywords used in our previous literature mining studies [1, 6] do not correspond to the same number of interaction types. While an interaction type or term in INO has its ontology ID, such a term may be associated with different synonyms or related keywords that can be used for literature mining. To support identification of genetic interactions in literature, synonyms and related keywords are needed. To meet this need, we have generated an annotation property called ‘has literature mining keywords’ (Figure 2). Such an annotation type allows the listing of different keywords mapping to the interaction type.

For example, the term ‘protein myristoylation’ in INO has five related literature mining terms including ‘myristoylate’, ‘myristoylates’, ‘myristoylated’, ‘myristoylating’, and ‘myristoylation’. These term variations are listed as an annotation of the interaction type using the annotation property ‘has literature mining keywords’ (Figure 2). The list of keywords can be easily extracted from the ontology by SPARQL or other methods and used for literature mining.

### (3) Statistics of INO terms and interaction keywords

As of May 31, 2014, INO contains 259 terms including 189 new INO terms and 70 terms from existing ontologies (Table 2). In addition to the ontologies introduced above, INO also imports terms from other authoritative domain ontologies such as the Chemical Entities of Biological Interest (ChEBI)
[17] and the Ontology of Genes and Genomes (OGG) [18]. Provenance and source ontology IDs are kept in our term importing [9].

In total, INO includes 355 terms under the branch of INO interaction. These INO interaction terms and their associated literature mining keywords can be used for efficient literature text tagging and retrieval of sentences containing these keywords. The usage of these terms and keywords in our literature mining study is described below.

B. INO-based literature mining of gene-gene interactions

(1) Workflow and system design

The workflow of the ontology-based gene pair enrichment analysis is illustrated in Figure 3.

![Figure 3. The workflow of INO-based gene-gene interaction enrichment analysis. The detail about the workflow is described in the text.](image)

Specifically, all publications from PubMed were first downloaded. The sentences of article titles and abstracts were parsed and pre-processed. Human gene names and interaction keywords were tagged. To tag human gene names, the HUGO human gene nomenclature assignments (http://www.genenames.org/) were used. These human gene names are also available in the OGG [18]. The INO interaction types and associated keywords were used for tagging interaction keywords. As detailed in the Methods section, an INO-based modified Fisher’s exact test was developed to identify enriched statistically significantly gene-gene interaction types and associated gene-gene pairs (Figure 3).

The INO-based workflow for literature mining of gene-gene interactions is applicable for different use case studies. Below we introduce the application of such a strategy for studying the gene-gene interactions in the vaccine domain.

(2) INO-based literature enrichment analysis of vaccine-associated gene-gene interaction data

Our literature mining analysis used all PubMed documents published as of 12/31/2013. A total of 23,481,042 PubMed documents were used as the background data set for the analysis. Using this data set, SciMiner identified 314,152 gene pairs, each of which was associated with at least one INO term.

We applied our study to the vaccine domain. A PubMed search for vaccine-related documents resulted in 237,061 hits (as of 12/31/2013). VO-SciMiner additionally identified 28,908 documents using VO terms, resulting in a total of 265,969 documents to define the vaccine-associated document sets. The gene-gene interactions (i.e., gene pairs) with positive SVM scores and at least one INO term at the same sentence level were generated from these 265,969 PubMed abstracts. A total of 6,116 gene pairs were associated with at least one INO term.

Out of 78 INO interaction terms associated with at least five gene-pairs of the vaccine-associated sub-network, 14 terms were significantly over-represented (BH p-value < 0.05 and a minimal enrichment fold of 2) (Table 2). The results indicate that these 14 interaction types are more extensively studied in the vaccine context among the research of all the gene-gene interaction types published in PubMed.

Table 2. Significantly over-represented INO terms among the gene-gene interaction pairs of vaccine-associated sub-network.

| INO_ID   | Reference Term     | Enrichment Fold | BH * P-value | Most Frequent Gene-pair (#) |
|----------|--------------------|-----------------|--------------|----------------------------|
| 0000140  | neutralization     | 6.6             | 0            | IFNG IL12A (5)**           |
| 0000096  | induction of production | 6.2           | 0            | TNF IFNG (2)               |
| 0001006  | gene fusion        | 5.6             | 0            | CD40LG CD40 (3)            |
| 00010103 | accessory regulation | 3.9            | 0            | CDRA CD4 (55)              |
| 0000062  | costimulation      | 3.7             | 0            | CD40 CD8A (4)              |
| 0000169  | synergization      | 3.0             | 0            | CD8A CD40 (5)              |
| 0000089  | co-regulation      | 2.9             | 0            | CD8A CD40 (5)              |
| 0559     | denitrification    | 2.9             | 0            | IL17A MUC8 (1)             |
| 0195     | covalent binding  | 2.5             | 0            | CSF2 AC0P (2)              |
| 0208     | genetic interaction | 4.9            | 1.82E-10     | CD40LG CD40 (3)            |
| 0571     | mRNA cleavage      | 23.2            | 2.58E-07     | CFI SUPT2H (1)             |
| 0982     | RNA cleavage       | 16.2            | 2.21E-06     | CFI SUPT3H (1)             |
| 0910     | nucleic acid cleavage | 6.4           | 6.31E-04     | CFI SUPT3H (1)             |
| 0018377  | trefoil myristoylation | 2.3          | 2.68E-03     | CD4 S100B (2)              |

Notes: * BH: Benjamini-Hochberg; ** IFNG_IL12A (5): represents the IFNG and IL12A gene pair shown in five papers.

Furthermore, our gene-gene interaction enrichment analysis was able to retrieve all the gene pairs associated with each interaction type (last column in Table 2). For example, as indicated in five publications (PubMed IDs: 19915058, 8557339, 15557182, 17517055, and 7525727), the cytokines interferon-gamma (IFNG) and interleukin-12A (IL12A) have been found to be closely related, and the neutralization of one cytokine often leads to decreased production of another one [19, 20]. Such neutralization-related research has been typically found in the field of vaccinology. In another example, associated with the interaction type “induction of production”, the production of one cytokine TNF (or IFNG) was found to be induced by another cytokine IFNG (or TNF) [21]. A close examination of all the gene pairs recorded in Table 2 shows that they are all related to the vaccine and immunity research. These results also confirm the specificity of our INO-based enrichment analysis.

In addition, our study found 17 significantly under-represented INO terms with a maximum enrichment fold of 0.5 (equivalent to 2 fold in over-representation) and BH P-value < 0.05 (Table 3). Compared to the general gene-gene interaction research, these interaction types are likely less
studied in the vaccinology research field. The reasons of these under-represented interaction types may vary. It is likely that some of these under-represented interactions represent new research opportunities in the vaccinology.

Table 3. Significantly under-represented INO terms among the gene-gene interaction pairs of vaccine-associated sub-network.

| INO_ID     | Reference Term                  | Enrichment Fold | BH P-value |
|------------|---------------------------------|-----------------|------------|
| MI 0203    | dephosphorylation reaction       | 0.06            | 0          |
| INO 0000178| tyrosine-phosphorylation         | 0.09            | 0          |
| INO 0000044| gene expression regulation       | 0.25            | 0          |
| INO 0000172| transactivation                  | 0.26            | 0          |
| INO 0000960| coprecipitation                 | 0.28            | 0          |
| GO 0016310 | phosphorylation                 | 0.36            | 0          |
| MI 0403    | colocalization                   | 0.36            | 0          |
| MI 0414    | enzymatic reaction              | 0.42            | 0          |
| MI 0194    | cleavage reaction               | 0.49            | 0          |
| MI 0213    | methylation reaction            | 0.37            | 6.84E-16   |
| INO 0000992| dissociation                     | 0.28            | 6.27E-15   |
| INO 0000484| coimmunoprecipitation           | 0.35            | 1.0E-13    |
| INO 0001115| hyperphosphorylation            | 0.27            | 2.54E-08   |
| INO 0000884| Destabilization                 | 0.28            | 1.49E-05   |
| GO 0000641 | protein complex assembly        | 0.24            | 1.97E-05   |
| INO 0000888| protein dimerization            | 0.26            | 6.41E-05   |
| INO 000171 | Termination                     | 0.42            | 3.98E-03   |

One advantage of INO based study is that we can rely on the INO hierarchy to identify the relations among enriched interaction types. Such a strategy is used to generate the hierarchies of enriched 14 over-represented and 17 under-represented INO interaction types (Figure 4). This study clearly shows the relations between many different interaction terms. For example, among the three over-represented terms ‘mRNA cleavage’, ‘RNA cleavage’, and ‘nucleic acid cleavage’, there are two parent-child relations as clearly shown in Figure 4. Interestingly, the term ‘cleavage reaction’ is one of the 17 under-represented terms (Table 3).

Both sets of over-represented and under-represented interaction terms share some common top-level terms including ‘regulation’, ‘direction interaction’, ‘association’, and ‘interaction’. Otherwise, specific profiles of the two sets are in general distinct at the bottom levels (Figure 4).

Figure 4. The hierarchies of over- and under-represented INO interaction terms. (A) The hierarchy of 14 over-represented INO interaction terms. (B) The hierarchy of 17 under-represented INO interaction terms. The results were generated using OntoFox [9] with the OntoFox setting “includeComputedIntermediates”, and visualized using the Protege-OWL editor (http://protege.stanford.edu/). The box-enclosed terms are over- or under-represented interaction types directly identified in our program (see Tables 2 and 3). Other terms not enclosed in boxes are terms retrieved by OntoFox to ensure the completeness of the hierarchies.

IV. DISCUSSION

This paper introduces two major contributions in the area of ontology-based literature mining research. First, we have for the first time systematically introduced the development of the INO ontology targeting for robust literature mining of gene-gene interaction types. It is noted that in addition to literature mining, INO is also being developed to model various interactions and networks among different molecules [22]. However, the INO development was initiated from meeting our literature mining need [6]. Second, we have proposed and implemented a novel INO-based gene-gene interaction enrichment strategy. The INO-based gene pair enrichment analysis is novel in that the input of such analysis is the literature mined gene-gene interaction types and gene pairs. It differs from a typical GO enrichment analysis where a list of genes is the input. Such a strategy was further used to study the enriched gene-gene interaction types and gene pairs in the domain of vaccinology. Our results demonstrate that the INO offers a repository of hierarchical interaction keywords and platform for allowing systematical retrieval of interaction types from the literature. The INO-based gene-gene interaction enrichment method further provides a strategy for analyzing the retrieved gene-gene interaction literature results.

Our INO-based literature mining study found that while it is relatively easy to describe the relation between two genes when only one interaction keyword exists in the sentence containing these two genes, it is much difficult to describe the relation between the two genes if multiple keywords exist. For example, in our IFNG-IL12A neutralization-related interaction type, we can know two of these genes participate in a neutralization-related interaction(s). However, it does not mean that IFNG neutralizes IL-12A, or vice versa. We can only say that these two genes interact somehow in a neutralization-related pattern. It is likely that multiple interaction-related keywords co-exist in one sentence. For example, an IFNG-IL12A neutralization-related interaction sentence is “In vitro IL-12 neutralization dramatically impairs the IFN-gamma response to S. typhimurium but not to ConA” [23]. This sentence contains two interaction-related keywords “neutralization” and “impaired”. This is a complex relation where a neutralization of one gene impairs another gene expression. It hints that one gene positively regulates another. In this case, the neutralization is really an experimental condition. Our literature mining program retrieved both keywords independently without considering them together. It would be more advanced if we could process these two keywords simultaneously and assign a unique interaction type, such as ‘impairment after neutralization’, which would be a subclass (or child term) of the existing INO term ‘positively regulation’. While this example demonstrates a new direction of future research, such analysis does not undermine the contributions of
the new INO-based literature mining strategy first reported in this manuscript. Indeed, our strategy provides a new start point and platform for further addressing these challenges.

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Authors’ contributions: JH developed the INO-based gene interaction enrichment assay and generated data with the vaccine domain use case. AO developed the SVM-based literature mining pipeline. ZX generated the script to execute the literature mining pipeline. YH developed the INO and was the primary writer of the manuscript. YH, JH, and AO all participated in the project design, result interpretation, and manuscript writing. All authors agreed on the manuscript publication.

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REFERENCES

[1] A. Ozgur, Z. Xiang, D. Radev, and Y. He, "Literature-based discovery of IFN-γ and vaccine-mediated gene interaction networks," *Journal of Biomedicine and Biotechnology*, vol. 2010, p. Article ID 426479 (13 pages). 2010.

[2] T. K. Jenssen, A. Laegreid, J. Komorowski, and E. Hovig, "A literature network of human genes for high-throughput analysis of gene expression," *Nat Genet*, vol. 28, pp. 21-8, May 2001.

[3] T. Joachims, "Making large-scale support vector machine learning practical," in *Advances in Kernel Methods: Support Vector Learning*, C. J. B. B. Schölkopf, and A. J. Smola, Eds., ed Cambridge, MA.: MIT Press, 1999, pp. 169-184.

[4] G. Erkan, A. Ozgur, and D. R. Radev, "Semi-supervised classification for extracting protein interaction sentences using dependency parsing," presented at the Proceedings of the 2007 Joint Conference on Empirical Methods in Natural Language Processing and Computational Natural Language Learning (EMNLP-CoNLL), Prague, Czech Republic, 2007.

[5] Y. He, L. Cowell, A. D. Diehl, H. L. Mobley, B. Peters, A. Ruttenberg, et al., "VO: Vaccine Ontology," in *The 1st International Conference on Biomedical Ontologies (ICBO-2009)*, Buffalo, NY, USA, 2009, p. http://precedings.nature.com/documents/3552/version/1.

[6] A. Ozgur, Z. Xiang, D. R. Radev, and Y. He, "Mining of vaccine-associated IFN-gamma gene interaction networks using the Vaccine Ontology," *J Biomed Semantics*, vol. 2 Suppl 2, p. S8, 2011.

[7] J. D. Kim, T. Ohta, Y. Tateisi, and J. Tsujii, "GENIA corpus—semantically annotated corpus for bio-textmining," *Bioinformatics*, vol. 19 Suppl 1, pp. i180-2, 2003.

[8] B. Smith, M. Ashburner, C. Rosse, J. Bard, W. Bug, W. Ceusters, et al., "The OBO Foundry: coordinated evolution of ontologies to support biomedical data integration," *Nat Biotechnol*, vol. 25, pp. 1251-5, Nov 2007.

[9] Z. Xiang, M. Courtot, R. R. Brinkman, A. Ruttenberg, and Y. He, "OntoFox: web-based support for ontology reuse," *BMC Res Notes*, vol. 3, p. 175, 2010.

[10] Z. Xiang, C. Mungall, A. Ruttenberg, and Y. He, "Ontobee: A linked data server and browser for ontology terms," in *The 2nd International Conference on Biomedical Ontologies (ICBO)*, Buffalo, NY, USA, 2011, pp. Pages 279-281 [http://ceur-ws.org/Vol-833/paper48.pdf].

[11] J. Hur, Z. Xiang, E. L. Feldman, and Y. He, "Ontology-based Brucella vaccine literature indexing and systematic analysis of gene-vaccine association network," *BMC Immunol*, vol. 12, p. 49, 2011.

[12] M. C. de Marmeffe, B. Maccartney, and C. D. Manning, "Generating typed dependency parses from phrase structure parses," in *Proceedings of LREC-06*, 2006, pp. 449-454.

[13] D. A. Hosack, G. Dennis, Jr., B. T. Sherman, H. C. Lane, and R. A. Lempicki, "Identifying biological themes within lists of genes with EASE," *Genome Biol*, vol. 4, p. R70, 2003.

[14] M. A. Sartor, V. Mahavisino, V. G. Keshamouni, J. Cavalcioni, Z. Wright, A. Karnovsky, et al., "ConceptGen: a gene set enrichment and gene set relation mapping tool," *Bioinformatics*, vol. 26, pp. 456-63, Feb 15 2010.

[15] S. Banerjee and T. Pedersen, "The Design, Implementation, and Use of the Ngram Statistic Package," in *Proceedings of the Fourth International Conference on Intelligent Text Processing and Computational Linguistics*, Mexico City, Mexico, 2003, pp. Pages 370-381 [http://www.d.umn.edu/~tpederse/Pubs/cicling2003-2.pdf].

[16] P. Grenon and B. Smith, "SNAP and SPAN: Towards Dynamic Spatial Ontology," *Spatial Cognition and Computation*, vol. 4, pp. 69-103, 2004.

[17] J. Hastings, P. de Matos, A. Dekker, M. Ennis, B. Harsha, N. Kale, et al., "The ChEBI reference database and ontology for biologically relevant chemistry: enhancements for 2013," *Nucleic Acids Res*, vol. 41, pp. D456-63, Jan 2013.

[18] Y. He, Y. Liu, and B. Zhao, "OGG: a biological ontology for representing genes and genomes in specific organisms," in *The 2014 International Conference on Biomedical Ontologies (ICBO 2014)*, Houston, TX, USA, 2014, pp. 1-6.

[19] X. Chen, M. A. O'Donnell, and Y. Luo, "Dose-dependent synergy of Th1-stimulating cytokines on bacille Calmette-Guérin-induced interferon-gamma production by human mononuclear cells," *Clin Exp Immunol*, vol. 149, pp. 178-85, Jul 2007.

[20] T. A. Wynn, I. P. Oswald, I. A. Eltoum, P. Caspar, C. J. Lowenstein, F. A. Lewis, et al., "Elevated expression of Th1 cytokines and nitric oxide synthase in the lungs of vaccinated mice after challenge infection with Schistosoma mansoni," *J Immunol*, vol. 153, pp. 5200-9, Dec 1 1994.

[21] S. J. Green, L. F. Scheller, M. A. Marletta, M. C. Seguin, F. W. Klotz, M. Slayter, et al. "The OBO Foundry: coordinated evolution of gene-vaccine association networks,” in *Proceedings of the 5th International Conference on Intelligent Text Processing and Computational Linguistics*, Mexico City, Mexico, 2003, pp. Pages 279-281 [http://ceur-ws.org/Vol-833/paper48.pdf].