Overview of the Pathway Curation (PC) task of BioNLP Shared Task 2013

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Abstract

We present the Pathway Curation (PC) task, a main event extraction task of the BioNLP shared task (ST) 2013. The PC task concerns the automatic extraction of biomolecular reactions from text. The task setting, representation and semantics are defined with respect to pathway model standards and ontologies (SBML, BioPAX, SBO) and documents selected by relevance to specific model reactions. Two BioNLP ST 2013 participants successfully completed the PC task. The highest achieved F-score, 52.8%, indicates that event extraction is a promising approach to supporting pathway curation efforts. The PC task continues as an open challenge with data, resources and tools available from http://2013.bionlp-st.org/

1 Introduction

Following developments in molecular biology, biological phenomena are increasingly understood on the molecular level, as the products of complex systems of molecular reactions. Pathway models formalizing biomolecules and their reactions in machine readable representations are a key way of sharing and communicating human understanding of these phenomena and of developing computational models of biological systems (Kitano, 2002). Many pathway models integrate knowledge from hundreds or thousands of scientific publications, and their curation requires substantial manual effort. To support this effort, we have developed PathText (Kemper et al., 2010) which provides a seamless environment integrating a pathway visualizer, text mining systems and annotation tools. Furthermore, automatic processing of the domain literature could thus potentially play an important role in the support of pathway curation.

Information extraction targeting biomolecular reactions has been a major focus of efforts in biomedical natural language processing, with several tasks, resources, and tools addressing in particular protein-protein interactions (Krallinger et al., 2007; Pyysalo et al., 2008; Tikk et al., 2010). However, most such efforts have employed simple representations, such as entity pairs, that are not sufficient for capturing molecular reactions to the level of detail required to support the curation of pathway models. Additionally, previous efforts have not directly involved the semantics (e.g. reaction type definitions) of such models. Perhaps in part due to these reasons, natural language processing and information extraction methods have not been widely embraced by biomedical pathway curation communities (Ohta et al., 2011c; Ohta et al., 2011a).

We believe that the extraction of structured event representations (Figure 1) pursued in the BioNLP Shared Tasks offers many opportunities to make significant contributions to support the development, evaluation and maintenance of biomolecular pathways. The Pathway Curation (PC) task, a main task of the BioNLP Shared Task 2013, is proposed as a step toward realizing these opportunities. The PC task aims to evaluate the applicability of event extraction systems to pathway curation and to encourage the further development of methods for related tasks. The design of the task aims to address current issues in information extraction for pathway curation by explicitly basing its representation and extraction targets on ma-
Figure 2: Illustration of pathway reaction (left), matching representation as an idealized text-bound event structure (middle) and applied event representation for statements actually appearing in text (right).

2 Task setting

The PC task is formulated as an event extraction task (Ananiadou et al., 2010) following the general representation and task setting first introduced in the BioNLP ST 2009 (Kim et al., 2011). The primary aim is the extraction of event structures, or events, each of which can involve any number of physical entities or other events in specific roles.

The event representation is sufficiently expressive to allow the definition of event structures that closely parallel the definition of reactions in pathway representations such as SBML and BioPAX. These pathway representations differentiate between three primary groups of reaction participants: reactants (“inputs”), products (“outputs”), and modifiers, where the specific roles of modifiers can be further identified to differentiate e.g. reaction catalysts from inhibitors. Correspondingly, the PC task applies the Theme role defined in previous BioNLP ST tasks to capture reactants, introduces a new Product role for products, and applies the previously defined Cause role and regulatory events to capture modifiers (Figure 2; see also Section 2.3).

It is important to note that while the event representation allows a one-to-one mapping to reactions in principle, an annotation scheme cannot guarantee that actual statements in text map to fully specified reactions: in free-form text, authors frequently omit mention of some entities taking part in reactions, perhaps most typically to avoid redundancies such as in “p38γ is phosphorylated into phospho-p38γ” (Figure 2b). Representations extracted from explicit statements in text will thus in some cases omit aspects of the corresponding complete reactions in pathway models.

Systems addressing the PC task are expected to extract events of specific types given 1) free-form text and 2) gold standard annotation for mentions of physical entities in that text. The task annotations also include equivalence relations and event modifications, a secondary extraction target. The annotation types are detailed below.

2.1 Entities

The entity annotation marks mentions of physical entities using start and end offsets in text (contiguous span) and a type selected from a fixed set. The following four entity types are marked in the PC task: SIMPLE CHEMICAL, annotated with reference to the Chemical Entities of Biological Interest (ChEBI) resource (Degtyarenko et al., 2008);
Table 1: Entity types, definitions, and reference resources.

| Entity type            | Scope                                    | Reference | Ontology ID |
|------------------------|------------------------------------------|-----------|-------------|
| SIMPLE CHEMICAL        | simple, non-repetitive chemical entities  | ChEBI     | SBO:0000247 |
| GENE OR GENE PRODUCT   | genes, RNA and proteins                  | gene/protein DBs | SBO:0000246 |
| COMPLEX                | entities of non-covalently linked components | complex DBs | SBO:0000253 |
| CELLULAR COMPONENT     | parts of cell and extracellular environment | GO-CC     | SBO:0000290 |

Table 2: Event types and arguments. “Molecule” refers to an entity annotation of any of the types SIMPLE CHEMICAL, GENE OR GENE PRODUCT, or COMPLEX, and “ANY” refers to an annotation of any type, either entity or event. The indentation corresponds to ontological relationships between the event types: for example, PHOSPHORYLATION is a CONVERSION and TRANSCRIPTION part of GENE EXPRESSION.

GENE OR GENE PRODUCT, annotated with reference to gene and protein databases such as UniProt (Consortium, 2011), Entrez Gene (Maglott et al., 2005) and PFam (Finn et al., 2010); COMPLEX, annotated with reference to database resources covering complexes; and CELLULAR COMPONENT, annotated following the scope of the Gene Ontology cellular component subontology (Ashburner et al., 2000) (Table 1). For discussion of the relation between these types and the representations applied in pathway models, we refer to Ohta et al. (2011c).

In terms of mention types in text, the annotation for SIMPLE CHEMICAL, GENE OR GENE PRODUCT and COMPLEX covers entity name mentions only, while the annotation for CELLULAR COMPONENT covers entity name mentions, nominal mentions, and adjectival references (e.g. “mitochondrial”).

### 2.2 Relations

The PC task defines one relation type, **Equiv** (equivalence), which can hold between entity mentions of the same type and specifies that they refer to the same real-world entity (Figure 3). These relations are only applied to determine if two events match during evaluation, where entities connected by an **Equiv** relation are considered interchangeable. Gold standard **Equiv** relations are applied also for test data, and systems participating in the task are not expected to extract these relations.

**2.3 Events**

The event annotation marks references to reactions, processes and comparable associations in scope of the annotation using the event representation. For the definition and scope of the event annotation, we rely primarily on the Systems Biology Ontology (SBO), drawing some general types not in scope of this ontology from the Gene Ontology (GO). Table 2 presents the event types anno-
Pathway Repository ID Publication
mTOR BioModels MODEL1012220002 (Caron et al., 2010)
mTORC1 upstream regulators BioModels MODEL1012220003 (Caron et al., 2010)
TLR BioModels MODEL2463683119 (Oda and Kitano, 2006)
Yeast Cell Cycle BioModels MODEL1011020000 (Kaizu et al., 2010)
Rb BioModels MODEL4132046015 (Calzone et al., 2008)
EGFR BioModels MODEL2463576061 (Oda et al., 2005)
Human Metabolic Network BioModels MODEL6399676120 (Duarte et al., 2007)
NF-kappaB pathway - (Oda et al., 2008)
p38 MAPK PANTHER DB P05918 -
p53 PANTHER DB P00059 -
p53 feedback loop pathway PANTHER DB P04392 -
Wnt signaling pathway PANTHER DB P00057 -

Table 3: Pathway models used to select documents for the task, with pathway repository model identifiers and publications presenting each model (when applicable).

The role in which each event argument (entity or other event) participates in an event is specified as one of the following:

**Theme** entity/event that undergoes the effects of the event. For example, the entity that is transcribed in a TRANSCRIPTION event or transported in a TRANSPORT event.

**Cause** entity/event that is causally active in the event. Marks, for example, “P₁” in “P₁ inhibits P₂ expression”.

**AtLoc,FromLoc,ToLoc** : location in which the Theme entity of a LOCALIZATION event is localized (At) in LOCALIZATION events not involving movement or is transported (or moves) from/to (From/To) in LOCALIZATION and TRANSPORT events involving movement.

**Site** site on the Theme entity that is modified in the event. Can be specified for modification events such as PHOSPHORYLATION.

**Participant** general role type identifying an entity that participates in some underspecified way in a high-level process. Only applied for the PATHWAY type.

### 2.4 Event modifications

In addition to events, the PC task defines a secondary extraction target, event modifications. Two modification types are defined: NEGATION and SPECULATION. Both are binary flags that modify events, the former marking an event as being explicitly stated as not occurring (e.g. “P is not phosphorylated”) and the latter as being stated in a speculative context (“P may be phosphorylated.”). Both are defined in terms of annotation scope and semantics identically as in the BioNLP ST’09 (Kim et al., 2009).

### 2.5 Evaluation

The PC task evaluation applies the standard evaluation criteria established in the BioNLP ST 2009. These criteria relax exact matching between gold and predicted events in two aspects: approximate trigger boundary matching, and approximate recursive event matching. The former allows predicted event triggers to differ from gold triggers by one word, and the latter requires recursively referred events to only match in their core arguments (see Table 2). We refer to Kim et al. (2011) for a detailed definition of these criteria.

### 3 Corpus

This section presents the PC task corpus and its annotation process.

#### 3.1 Document selection

To assure that the documents annotated for the PC task corpus are relevant to pathway reactions, we applied two complementary approaches, both selecting documents on the basis of relevance to a specific pathway reaction. First, we selected from the BioModels repository those pathway models with the largest numbers of manually created annotations referencing a specific PubMed document identifier. For each of these models, we extracted literature references, selected a random subset, downloaded the documents, and manually filtered to select abstracts that explicitly discuss relevant molecular reactions. Second, as only a small subset of models include explicit references to the
literature providing evidence for specific pathway reactions, we applied an alternative strategy where reactions from a selection of PANTHER DB models were entered into the PathText system (Kemper et al., 2010), which is capable of suggesting documents relevant to given reactions based on an SBML model. We then selected a random set of reactions to query the system, and manually evaluated the highest-ranking documents to identify those whose abstracts explicitly discuss the selected reaction. We refer to Miwa et al. (2013a) for a detailed description of this approach. Table 3 presents the pathway models on which the document selection was based.

3.2 Annotation process

The base entity annotation for the PC corpus was created automatically using state-of-the-art entity mention taggers for each of the targeted entity types. For SIMPLE CHEMICAL tagging, the OS-CAR4 system (Jessop et al., 2011) trained on the chemical named entity recognition corpus of Corbett and Copestake (2008) was applied. For GENE OR GENE PRODUCT mention detection, the NERSuite5 system trained on the BioCreative 2 Gene Mention task (Wilbur et al., 2007) corpus was used. NERSuite was also applied for CELLULAR COMPONENT mention detection, for this task trained on the Anatomical Entity Mention (AnEM) corpus (Ohta et al., 2012). Finally, COMPLEX annotations were created using a combination of a dictionary and heuristics making use of the GENE OR GENE PRODUCT annotation (for mentions such as “cyclin E/CDK2 complex”). To support the curation process, these tools were integrated into the NaCTeM text-analysis workflow system Argo (Rak et al., 2012).

Based on the evaluations of each of these tools in the studies presenting them, we expected initial automatic tagging performance to be in the range 80-90% in both precision and recall. Following initial automatic annotation, the entity mention annotation was manually revised to improve quality and consistency. As the entity annotation is not itself a target of extraction in the shared task, we did not separately evaluate the consistency of the revised entity mention annotation.

To assure that the quality and consistency of the event annotation are as high as possible, ini-

| Item     | Train | Devel | Test  | Total |
|----------|-------|-------|-------|-------|
| Documents| 260   | 90    | 175   | 525   |
| Words    | 53811 | 18579 | 35966 | 108356|
| Entities | 7855  | 2734  | 5312  | 15901 |
| Events   | 5992  | 2129  | 4004  | 12125 |
| Modifications | 317  | 80    | 174   | 571   |

Table 4: PC corpus statistics

4 Results

4.1 Participation

Two groups submitted final results to the PC task, one from the National Centre for Text Mining (NaCTeM) and one from the University of Turku BioNLP group (TEES-2.1) (Table 5). Both participants applied their well-established, state-of-the-art event extraction systems, EventMine7 (Miwa et al., 2012) (NaCTeM) and the Turku

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5http://lersuite.nlplab.org/

6This selection implies that the consistency of the event annotation of the final corpus is expected to exceed the 61% F-score of the IAA experiment. Consistency after selection was not separately evaluated.

7http://nactem.ac.uk/EventMine/
Event Extraction System\(^8\) (Björne et al., 2011) (TEES). The two systems share the same overall architecture, a one-best pipeline with SVM-based stages for event trigger detection, trigger-argument relation detection, argument grouping into event structures, and modification prediction. The feature representations of both systems draw on substructures of dependency-like representations of sentence syntax, derived from full parses of input sentences. TEES applies the Charniak and Johnson (2005) parser with the McClosky (2009) biomedical model, converting the phrase-structure parses into dependencies using the Stanford tools (de Marneffe et al., 2006). By contrast, EventMine uses a combination of the predicate-argument structure analyses created by the deep parser Enju (Miyao and Tsujii, 2008) and the output of the the GDep best-first shift-reduce dependency parser (Sagae and Tsujii, 2007). All three parsers have models trained in part on the biomedical domain GENIA treebank (Tateisi et al., 2005).

Interestingly, both systems make use of the GE task data, but the application of EventMine extends on this considerably by applying a stacked model (Miwa et al., 2013b) with predictions also from models trained on the BioNLP ST 2011 EPI and ID tasks (Pyysalo et al., 2012) as well as from four corpora introduced outside of the shared tasks by Thompson et al. (2011), Pyysalo et al. (2011), Ohta et al. (2011b) and Ohta et al. (2011c).

### 4.2 Evaluation results

Table 6 summarizes the primary evaluation results. The two systems demonstrate broadly similar performance in terms of F-scores, with NaCTeM achieving an 1.7% point higher overall result.

| Team     | recall | prec. | F-score |
|----------|--------|-------|---------|
| NaCTeM   | 52.23  | 53.48 | 52.84   |
| TEES-2.1 | 47.15  | 55.78 | 51.10   |

Table 6: Primary evaluation results

However, the systems show quite different performance in terms of the precision/recall balance: while the NaCTeM system has little difference between precision and recall, TEES-2.1 shows a clear preference for precision, with 8.6% lower recall than precision.

Results are shown separately for each event type in Table 7. The results largely mirror the overall performance, with the NaCTeM system showing better performance for 13 out of the 21 event types present in the test data and more balanced precision and recall than TEES-2.1, which emphasizes precision over recall for almost all event types. Although the results do not include evaluation of EventMine with a reduced set of stacked models in training, the modest difference in performance suggests that comprehensive use of previously released event resources in EventMine did not confer a decisive advantage, perhaps in part due to differences in the event definitions between the PC task and previous resources.

Overall, the two systems appear quite similar not only in architecture but also performance, with the clearest systematic difference observed being the different emphases on precision vs. recall. As both systems are based on machine learning methods with real-valued outputs, it would be relatively straightforward to use prediction confidences to analyse performance over the entire precision-recall curve instead of a single fixed point. Such analysis could provide further insight into the relative strengths and weaknesses of these two systems.

### 5 Discussion

Although participation in this initial run of the PC task was somewhat limited, the two participating systems have been applied to a large variety of event extraction tasks over the last years and have shown consistently competitive performance with the state of the art (Björne and Salakoski, 2011; Miwa et al., 2012). It is thus reasonable to assume that the higher performance achieved by the
Table 7: Primary evaluation results by event type.

| Event                  | NaCTeM recall | NaCTeM prec. | NaCTeM F-score | TEES-2.1 recall | TEES-2.1 prec. | TEES-2.1 F-score |
|------------------------|--------------|--------------|---------------|----------------|----------------|-----------------|
| CONVERSION             | 34.33        | 35.48        | 34.90         | 35.82          | 42.86          | 39.02           |
| PHOSPHORYLATION        | 62.46        | 55.94        | 59.02         | 53.40          | 66.00          | 59.03           |
| DEPHOSPHYLATION        | 45.00        | 56.25        | 50.00         | 35.00          | 77.78          | 48.28           |
| ACETYLATION            | 69.57        | 72.73        | 71.11         | 82.61          | 76.00          | 79.17           |
| DEACETYLATION          | 33.33        | 33.33        | 33.33         | 0.00           | 0.00           | 0.00            |
| METHYLATION            | 42.86        | 60.00        | 50.00         | 57.14          | 80.00          | 66.07           |
| DEMETHYLATION          | 100.00       | 100.00       | 100.00        | 100.00         | 100.00         | 100.00          |
| UBIQUITINATION         | 52.94        | 64.29        | 58.06         | 58.82          | 76.92          | 66.07           |
| DEUBIQUITINATION       | 100.00       | 100.00       | 100.00        | 100.00         | 100.00         | 100.00          |
| LOCALIZATION           | 42.25        | 61.22        | 50.00         | 43.66          | 54.39          | 48.44           |
| TRANSPORT              | 65.52        | 61.29        | 63.33         | 56.55          | 59.85          | 58.16           |
| GENE EXPRESSION        | 90.65        | 83.15        | 86.74         | 84.55          | 79.39          | 81.89           |
| TRANSCRIPTION          | 71.15        | 82.22        | 76.29         | 57.69          | 73.17          | 64.52           |
| TRANSLATION            | 0.00         | 0.00         | 0.00          | 50.00          | 100.00         | 66.07           |
| Simple-total           | 66.42        | 64.80        | 65.60         | 60.40          | 67.87          | 63.92           |
| DEGRADATION            | 78.57        | 89.19        | 83.54         | 78.57          | 78.57          | 78.57           |
| ACTIVATION             | 78.54        | 70.96        | 74.56         | 72.06          | 72.06          | 72.06           |
| INACTIVATION           | 44.62        | 55.77        | 49.57         | 38.46          | 45.45          | 41.67           |
| BINDING                | 64.96        | 47.30        | 54.74         | 53.96          | 53.96          | 53.96           |
| DISSOCIATION           | 38.46        | 46.88        | 42.25         | 35.90          | 45.16          | 40.00           |
| PATHWAY                | 84.91        | 75.50        | 79.93         | 70.94          | 75.50          | 73.15           |
| General-total          | 69.07        | 62.69        | 65.72         | 61.16          | 65.74          | 63.37           |
| REGULATION             | 33.33        | 33.97        | 33.65         | 29.73          | 39.51          | 33.93           |
| POSITIVE REGULATION    | 35.49        | 42.81        | 38.81         | 34.51          | 45.45          | 39.23           |
| NEGATIVE REGULATION    | 45.75        | 50.64        | 48.07         | 41.02          | 47.37          | 43.97           |
| Regulation-total       | 37.73        | 42.79        | 40.10         | 35.17          | 44.76          | 39.39           |
| Sub-total              | 53.47        | 53.96        | 53.72         | 48.23          | 56.22          | 51.92           |
| NEGATION               | 24.52        | 35.87        | 29.13         | 25.16          | 41.30          | 31.27           |
| SPECULATION            | 15.79        | 22.22        | 18.46         | 0.00           | 0.00           | 0.00            |
| Modification-total     | 23.56        | 34.65        | 28.05         | 22.41          | 40.00          | 28.73           |
| Total                  | 52.23        | 53.48        | 52.84         | 47.15          | 55.78          | 51.10           |

task participants, a balanced F-score of 52.8%, is a good estimate of the performance level that can be attained for this task by present event extraction technology.

The results achieved by the two systems are broadly comparable to the best results achieved by any system in similar previously introduced event extraction tasks (Kim et al., 2012; Pyysalo et al., 2012). Given the novelty of the task domain and reference resource and the broad selection of documents, we find the results highly encouraging regarding the applicability of event extraction technology to supporting the development, evaluation, and maintenance of pathway models.

6 Conclusions

This paper presented the Pathway Curation (PC) task, a main event extraction task of the BioNLP ST 2013. The task was organized in collaboration between groups with an interest in pathway curation with the aim of evaluating and advancing the state of the art in event extraction toward methods for developing, evaluating and maintaining formal pathway models in representations such as SBML and BioPAX. We introduced an event extraction task setting with reference to pathway model standards and the Systems Biology Ontology, selected a set of 525 publication abstracts relevant to specific model reactions, and created fully manual
event annotation marking over 12,000 event structures in the corpus.

Two participants in the BioNLP ST 2013 submitted final predictions to the PC task, applying established, state-of-the-art event extraction systems, EventMine and the Turku Event Extraction System. Both systems achieved F-scores over 50%, with the EventMine system achieving the best overall result of 52.8%. This level of performance is broadly comparable with results achieved in comparable previously proposed tasks, indicating that current event extraction technology is applicable to the projected pathway curation support tasks.

To allow the further development and evaluation of event extraction methods for the task, the PC task continues as an open challenge to all interested participants, with the annotated corpus data, supporting resources, and evaluation tools available under open licenses from the task homepage, http://2013.bionlp-st.org/

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