Tunable Domain-Independent Event Extraction in the MIRA Framework

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Abstract

We describe the system of the PIKB team for BioNLP'09 Shared Task 1, which targets tunable domain-independent event extraction. Our approach is based on a three-stage classification: (1) trigger word tagging, (2) simple event extraction, and (3) complex event extraction. We use the MIRA framework for all three stages, which allows us to trade precision for increased recall by appropriately changing the loss function during training. We report results for three systems focusing on recall (R = 28.88%), precision (P = 65.58%), and F1-measure (F1 = 33.57%), respectively.

1 Introduction

Molecular interactions have been the focus of intensive research in the development of in-silico biology. Recent developments like the Pathway and Interaction Knowledge Base (PIKB) aim to make available to the user the large semantics of the existing molecular interactions data using massive knowledge syndication. PIKB is part of LinkedLifeData1, a platform for semantic data integration based on RDF2 syndication and lightweight reasoning.

Our system is based on the MIRA framework where, by appropriately changing the loss function on training, we can achieve any desirable balance between precision and recall. For example, low precision with high recall would be appropriate in a search that aims to identify as many potential candidates as possible to be further examined by the user, while high precision might be essential when adding relations to a knowledge base. Such a tunable system is practical for a variety of important tasks, including but not limited to, populating extracted facts in PIKB and reasoning on top of new and old data.

Our system is based on a three-stage classification process: (1) trigger word tagging using a linear sequence model, (2) simple event extraction, and (3) complex event extraction. In stage (2), we generate relations between a trigger word and one or more proteins, while in stage (3), we look for complex interactions between simple events, trigger words and proteins. We use MIRA for all three stages with a loss function tuned for high recall.

2 One-best MIRA and Loss Functions

In what follows, $x_i$ will denote a generic input sentence, and $y_i$ will be the “gold” labeling of $x_i$. For each pair of a sentence $x_i$ and a labeling $y$, we compute a vector-valued feature representation $f(x_i, y)$. Given a weight vector $w$, the dot-product $w \cdot f(x_i, y)$ ranks the possible labelings $y$ of $x$; we will denote the top scoring labeling as $y_{w} (x)$. As with hidden Markov models (Rabiner, 1989), $y_{w} (x)$ can be computed efficiently for suitable feature functions using dynamic programming.

The learning portion of our method requires finding a weight vector $w$ that scores the correct labeling of the training data higher than any incorrect labeling. We used a one-best version of MIRA (Crammer, 2004; McDonald et al., 2005) to choose $w$. MIRA is an online learning algorithm that updates the weight vector $w$ for each training sentence $x_i$ according to the following rule:

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1http://www.linkedlifedata.com
2http://www.w3.org/RDF/
\[ w_{\text{new}} = \arg\min_w \|w - w_{\text{old}}\| \]

s.t. \[ w \cdot f(x_i, y_i) - w \cdot f(x, \hat{y}) \geq L(y_i, \hat{y}) \]

where \( L(y_i, y) \) is a measure of the loss of using \( y \) instead of the correct labeling \( y_i \), and \( \hat{y} \) is a shorthand for \( y_{\text{old}}(x_i) \). In case of a single constraint, this program has a closed-form solution. The most straightforward and the most commonly used loss function is the Hamming loss, which sets the loss of labeling \( y \) with respect to the gold labeling \( y_i \) as the number of training examples where the two labelings disagree. Since Hamming loss is not flexible enough for targeted training towards recall or precision, we use a number of task-specific loss functions (see Sections 3 and 5 for details). We implemented one-best MIRA and the corresponding loss functions in an in-house toolkit called Edlin. Edlin provides general machine learning architecture for linear models and a framework with implementations of popular learning algorithms including Naive Bayes, perceptron, maximum entropy, one-best MIRA, and conditional random fields (CRF) among others.

### 3 Trigger Word Tagging

The training and the development abstracts were first tokenized and split into sentences using maximum entropy models trained on the Genia\(^3\) corpora. Subsequently, we trained several sequence taggers in order to identify the trigger words in text. All our experiments used the standard BIO encoding (Ramshaw and Marcus, 1995) with different feature sets and learning procedures. We focused on recall since it determines the upper bound on the performance of our final system. In our experiments, we found that simultaneously identifying trigger words and the event types they trigger yielded low recall; thus, we settled on identifying trigger words in text as one kind of entity, regardless of event types.

In our initial experiments, we used a CRF-based sequence tagger (Lafferty et al., 2001), which yielded R=43.51\%. We further tried feature induction (McCallum, 2003) and second-order Markov assumptions for the CRF, achieving 44.72\% and 49.64\% recall, respectively.

\(^3\)http://www-tsujii.is.s.u-tokyo.ac.jp/GENIA/home/wiki.cgi

### 4 Event Extraction

The input to our event extraction algorithm is a list of trigger words and a list of genes or gene products.
ucts (e.g., proteins); the output is a set of relations as defined for Task 1. Our algorithm works in two stages. First, we generate events corresponding to relations between a trigger word and one or more proteins (simple events); then we generate events for relations between trigger words, proteins and simple events (complex events). The two stages differ only in the input data; thus, below we will describe our system for the first stage only.

For each sentence, we considered all pairs of entities (trigger words and proteins), and we used an unstructured classifier to determine the relationship for a given pair. These relationships encoded both the type of event (e.g., binding, regulation) and entities’ roles in that event (e.g., theme, cause); there was also a special relationship for unrelated entities. We constructed labeled examples to train a MIRA classifier using the training data provided by the task organizers; \(n\)-ary relations were then reconstructed from classifier’s predictions. The features we used are summarized in Table 2: they are over the words separating the two entities and their part-of-speech tags. We further used some simple features from syntactic phrases (OpenNLP\(^4\) parser) and dependency parse trees (McDonald et al., 2005), extracted using parsers trained on Genia corpora.

After some initial experiments, we found that our features were not sufficiently rich to allow us to learn the relationships between proteins that are part of the same event: we achieved a very low recall of about 20%. Consequently, we focused on the relationships between a trigger word and a protein. Since the competition stipulated that each trigger could be associated with only one type of event, we first chose the event type for each trigger by selecting the protein-label pair with the highest score. We then fixed the event type for this trigger word, and we discarded all proteins for which our classifier assigned a different event type to the target trigger-protein pair. Finally, we added to our output list all binary relations where the role of the protein was theme.

For some event classes — binding, regulation, positive regulation and negative regulation — the output of the binary classifier was further transformed so that \(n\)-ary relations can be formed. However, the way we did this was somewhat ad-hoc. For binding events, we added a 3-ary relation between the trigger, the highest scoring protein, and the second highest scoring protein. For regulation events, we added a 3-ary relation between the trigger and every pair of proteins where one was a theme and the other one was a cause. This aggressive addition of potential matches slightly reduced the overall precision, but helped improve the recall for the final system.

### Table 3: Our official results: for an erroneous submission.

| Event Class       | R   | P   | F₁  |
|-------------------|-----|-----|-----|
| Localization      | 10.92 | 82.61 | 19.29 |
| Binding           | 7.20  | 39.68 | 12.20 |
| Gene expression   | 30.47 | 74.58 | 43.26 |
| Transcription     | 10.95 | 39.47 | 17.14 |
| Protein catabolism| 28.57 | 57.14 | 38.10 |
| Phosphorylation   | 34.07 | 86.79 | 48.94 |
| Event Total       | 21.52 | 68.68 | 32.77 |
| Regulation        | 1.37  | 26.67 | 2.61  |
| Positive regulation | 1.12 | 25.58 | 2.14  |
| Negative regulation | 0.26 | 100.00 | 0.53 |
| Regulation Total  | 0.97  | 27.12 | 1.87  |
| Overall           | 10.84 | 64.13 | 18.55 |

5 Results and Discussion

Unfortunately, we made an error when making our official submission, which resulted in low scores; Table 3 shows the results for that submission.

The rest of this section describes the results and the implementation for the system we intended to submit. All reported results are for exact span matches and were obtained using the online tool provided by the task organizers.

As stated in Section 4, we used a linear model trained using one-best MIRA with ten runs over the data for the event extraction system. We oversampled the unstructured training instances that corresponded to a relation so that they become roughly equal in number to those that do not correspond to a relation. Finally, we performed parameter averaging as described in (Freund and Schapire, 1999). These details turned out to be very important for the system performance.

Table 4 shows the results for three different loss functions that gave the best results in our experiments. In describing the loss functions, we define three different types of errors: (1) if the system correctly predicted that a relation should be present,
but guessed the wrong type, we call this a cross-labeling; (2) a false positive occurs when the learner guessed some relation while there should have been none; (3) the reverse is a false negative. All loss functions we considered had a cross-labeling loss of 1. The 0-1 loss also has a loss of 1 for false positives and false negatives. The high-recall loss function penalizes false positives with 0.1 and false negatives with 5. The high-precision loss function penalizes false negatives with 0.1 and false positives with 5. The values 0.1 and 5 were chosen on the development data, but were not optimized aggressively.

In conclusion, we have built three domain-independent event extraction systems based on the MIRA framework, each using a different loss function. Overall, they perform quite well and would have been ranked second on precision\(^5\), and 6\(^{th}\) on recall, and 7\(^{th}\) on F\(_1\)-measure.

### 6 Future Work

After integrating domain knowledge, which should improve the recall for complex events and should boost the overall precision, we intend to transform the system output into RDF and add it to the PIKB repository. The required efforts discouraged us from building a middle ontology between the BioNLP and the PIKB data models, especially given the time limitations for the present task competition. However, we believe this is a promising direction, which we plan to pursue in future work.

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\(^5\)Our official submission is second on precision as well.

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