Extracting Protein-Protein interactions using simple contextual features

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1 Introduction

There has been much interest in recent years on the topic of extracting Protein-Protein Interaction (PPI) information automatically from scientific publications. This is due to the need that has emerged to organise the large body of literature that is generated through research, and collected at sites such as PubMed. Easy access to the information contained in published work is vital for facilitating new research, but the rate of publication makes manual collection of all such data unfeasible. Information Extraction approaches based on Natural Language Processing can be, and are already being used, to facilitate this process.

The dominant approach so far has been the use of hand-built, knowledge-based systems, working at levels ranging from surface syntax to full parses (Blaschke and Valencia, 2002; Huang et al., 2004; Plake et al., 2005; Rebholz-Schuhmann et al., 2005; Yakushiji et al., 2005). A similar work to the one presented here is by (Sugiyama et al., 2003), but it is not possible to compare results due to differing datasets and the limited information available about their methods.

2 Data

A gene-interaction corpus derived from the BioCreAtiVe task-1A data will be used for the experiments. This data was kindly made available by Jörg Hakenberg\(^1\) and is described in (Plake et al., 2005). The data consists of 1000 sentences marked up for POS tags, genes (both genes and proteins are marked as ‘gene’; the terms will be used interchangeably in this paper) and iWords. The corpus contains 255 relations, all of which are intra-sentential, and the “interaction word” (iWord)\(^2\) for each relation is also marked up.

I utilise the annotated entities, and focus only on relation extraction. The data contains directionality information for each relation, denoting which entity is the ‘agent’ and which the ‘target’, or denoting that this distinction cannot be made. This information will not be used for the current experiments, as my main aim is simply to identify relations between entities, and the derivation of this information will be left for future work.

I will be using the Naive Bayes, KStar, and JRip classifiers from the Weka toolkit, Zhang Le’s Maximum Entropy classifier (Maxent), TiMBL, and LibSVM to test performance. All experiments are done using 10-fold cross-validation. Performance will be measured using Recall, Precision and F1.

3 Experiments

Each possible combination of proteins and iWords in a sentence was generated as a possible relation ‘triple’, which combines the relation extraction task with the additional task of finding the iWord to describe each relation. 3400 such triples occur in the data. After each instance is given a probability by the classifiers, the highest scoring instance for each protein pairing is compared to a threshold to decide

\(^1\)See http://www.informatik.hu-berlin.de/~hakenberg/publ/suppl/sac05/

\(^2\)A limited set of words that have been determined to be informative of when a PPI occurs, such as interact, bind, inhibit, phosphorylation. See footnote 1 for complete list.
the outcome. Correct triples are those that match the iWord assigned to a PPI by the annotators.

For each instance, a list of features were used to construct a ‘generic’ model:

**interindices** The combination of the indices of the proteins of the interaction; “P1-position:P2-position”

**interwords** The combination of the lexical forms of the proteins of the interaction; “P1:P2”

**p1prevword, p1currword, p1nextword** The lexical form of P1, and the two words surrounding it

**p2prevword, p2currword, p2nextword** The lexical form of P2, and the two words surrounding it

**p2pdistance** The distance, in tokens, between the two proteins

**inbetween** The number of other identified proteins between the two proteins

**iWord** The lexical form of the iWord

**iWordPosTag** The POS tag of the iWord

**iWordPlacement** Whether the iWord is between, before or after the proteins

**iWord2ProteinDistance** The distance, in words, between the iWord and the protein nearest to it

A second model incorporates greater domain-specific features, in addition to those of the ‘generic’ model:

**patterns** The 22 syntactic patterns used in (Plake et al., 2005) are each used as boolean features.

**lemmas and stems** Lemma and stem information was used instead of surface forms, using a system developed for the biomedical domain.

4 Results

Tables 1 and 2 show the results for the two models described above. The system achieves a peak performance of 59.2% F1, which represents a noticeable improvement over previous results on the same dataset (52% F1 (Plake et al., 2005)), and demonstrates the feasibility of the approach adopted.

It is seen that simple contextual features are quite informative for the task, but that a significant gains can be made using more elaborate methods.

| Algorithm | Recall | Precision | F1  |
|-----------|--------|-----------|-----|
| Naive Bayes | 61.3   | 35.6      | 45.1 |
| KStar     | 65.2   | 41.6      | 50.8 |
| Jrip      | 66.0   | 45.4      | 53.8 |
| Maxent    | 58.5   | 48.2      | 52.9 |
| TiMBL     | 49.0   | 41.1      | 44.7 |
| LibSVM    | 49.4   | 56.8      | 52.9 |

Table 1: Results using ‘generic’ model

| Algorithm | Recall | Precision | F1  |
|-----------|--------|-----------|-----|
| Naive Bayes | 64.8   | 44.1      | 52.5 |
| KStar     | 60.9   | 45.0      | 51.8 |
| Jrip      | 44.3   | 45.7      | 45.0 |
| Maxent    | 57.7   | 56.6      | 57.1 |
| TiMBL     | 42.7   | 74.0      | 54.1 |
| LibSVM    | 54.5   | 64.8      | 59.2 |

Table 2: Results using extended model

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