Learning for Collective Information Extraction

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Abstract

An Information Extraction (IE) system analyses a set of documents with the aim of identifying certain types of entities and relations between them. Most IE systems treat separate potential extractions as independent. However, in many cases, considering influences between different candidate extractions could improve overall accuracy. For example, phrase repetitions inside a document are usually associated with the same entity type, the same being true for acronyms and their corresponding long form. One of our goals in this thesis is to show how these and potentially other types of correlations can be captured by a particular type of undirected probabilistic graphical model. Inference and learning using this graphical model allows for "collective information extraction" in a way that exploits the mutual influence between possible extractions. Preliminary experiments on learning to extract named entities from biomedical and newspaper text demonstrate the advantages of our approach.

The benefit of doing collective classification comes however at a cost: in the general case, exact inference in the resulting graphical model has an exponential time complexity. The standard solution, which is also the one that we used in our initial work, is to resort to approximate inference. In this proposal we show that by considering only a selected subset of mutual influences between candidate extractions, exact inference can be done in linear time. Consequently, a short term goal is to run comparative experiments that would help us choose between the two approaches: exact inference with a restricted subset of mutual influences or approximate inference with the full set of influences.

The set of issues that we intend to investigate in future work is two fold. One direction refers to applying the already developed framework to other natural language tasks that may benefit from the same types of influences, such as word sense disambiguation and part-of-speech tagging. Another direction concerns the design of a sufficiently general framework that would allow a seamless integration of cues from a variety of knowledge sources. We contemplate using generic sources such as external dictionaries, or web statistics on discriminative textual patterns. We also intend to alleviate the modeling problems due to the intrinsic local nature of entity features by exploiting syntactic information. All these generic features will be input to a feature selection algorithm, so that in the end we obtain a model which is both compact and accurate.

Contents

1	Intr	oduction	4
2	Bac	kground and Related Work	6
	2.1	Phrase Classification Approaches	7
		2.1.1 Rapier	7
		2.1.2 SRV	7
	2.2	Token Classification Approaches	8
		2.2.1 Hidden Markov Models	8
		2.2.2 Discriminative Models	10
		2.2.3 Maximum Entropy Models	10
		2.2.4 Conditional Random Fields	12
	2.3	Markov Random Fields	13
	2.4	Relational Markov Networks	14
3	Con	nnleted Research	16
U	31	Candidate Entities	16
	3.2	Entity Features	17
	33	The RMN Framework for Entity Recognition	17
	3.4	Local Clique Templates	19
	3.5	Global Clique Templates	$\frac{1}{20}$
	0.0	3.5.1 The Overlap Template	21
		3.5.2 The Repeat Template	21
		3.5.3 The Acronym Template	$\frac{1}{22}$
	3.6	Inference in Factor Graphs	23
	3.7	Learning Potentials in Factor Graphs	24
	3.8	Experimental Results	25
		1 A A A A A A A A A A A A A A A A A A A	
4	Cur	rent Research	28
	4.1	Local models	29
		4.1.1 Exact, linear time inference	29
		4.1.2 Learning Algorithm	31
		4.1.3 Experimental Results	33
5	Pro	posed Work	34
-	5.1	Restricted global models	34
		5.1.1 Exact. linear time inference	35
		5.1.2 Learning Algorithm	35
	5.2	Collective classification for WSD and POS tagging	36
	5.3	Using the web to improve information extraction	37
	5.4	Flexible use of external dictionaries	38
	5.5	Feature selection	39
	5.6	Relation extraction and syntactic information	40
	G		
6	Con	clusion	41

References

1 Introduction

Information Extraction (IE) is an important task in natural language processing, with many practical applications. It involves analyzing text documents, identifying particular types of entities, relations or events, and populating database slots with information about them. A basic component of an IE system is that of named entity recognition - the task of locating references to specific types of items in natural-language text. Since IE systems are difficult and time-consuming to construct, most recent research has focused on empirical techniques that automatically construct information extractors by training on supervised corpora (Cardie, 1997; Califf, 1999). Traditionally, IE systems have been trained to recognize names of people, organizations and locations (MUC (Grishman, 1995), CoNLL (Tjong Kim Sang & De Meulder, 2003)). Recently, substantial resources have been allocated for automatically extracting information from biomedical corpora, which has naturally led to the need of locating biologically relevant entity types, such as genes, proteins, or diseases. The wide variety of names used in the biomedical literature, coupled with their lack of formal structure, have made the IE problem especially difficult. This has further motivated the search for methods which are able to efficiently use any type of task-relevant knowledge. One particular type of knowledge which is especially useful for recognizing biological entities refers to correlations between the labels of repeated phrases inside a document, as well as between acronyms and their corresponding long form. In both cases, the mentioned phrases tend to have the same entity label. For example, Figure 1 shows part of an abstract from Medline, an online database of biomedical articles. In this abstract, the protein referenced by 'rpL22' is first introduced by its long name 'ribosomal protein L22', followed by the short name 'rpL22' between parentheses. The presence of the word 'protein' is a very good indicator that the entire phrase 'ribosomal protein L22' is a protein name. Also, 'rpL22' is an acronym of 'ribosomal protein L22' which increases the likelihood that it too is a protein name. The same name 'rpL22' occurs later in the abstract in contexts which do not indicate so clearly the entity type, however we can use the fact that repetitions of the same name tend to have the same type inside the same document.

The control of human **ribosomal protein L22** (rpL22) to enter into the nucleolus and its ability to be assembled into the ribosome is regulated by its sequence . The nuclear import of rpL22 depends on a classical nuclear localization signal of four lysines at positions 13 - 16 . RpL22 normally enters the nucleolus via a compulsory sequence of KKYLKK (I - domain , positions 88 - 93) ... Once it reaches the nucleolus , the question of whether rpL22 is assembled into the ribosome depends upon the presence of the N - domain .

Figure 1: Medline abstract with all protein names emphasized.

It is not always the case that repeated phrases have the same label. Figure 2 shows an example, where the first occurrence of 'eNOS' is a protein name, while its second occurrence is not a protein name by itself, because it is included in another protein name 'eNOS interaction protein'. Constraining repeated words like 'eNOS' to have the same label (i.e. either Inside or Outside a protein name) does not solve the problem either, as shown in Figure 2, where both tokens 'nitric' and 'oxide' are first tagged as Outside, and then Inside a protein name. In Section 3.5.2 we show how to capture the correlations between the labels of repeated phrases so that all the exceptions above are taken into account.

The capitalization pattern of the name itself is another useful indicator, nevertheless it is not sufficient by itself, as similar patterns are also used for other types of biological entities such as cell types or aminoacids (see Figure 3). Therefore, correlations between the labels of repeated phrases, or between acronyms and

Production of nitric oxide (NO) in endothelial cells is regulated by direct interactions of **endothelial nitric oxide synthase** (**eNOS**) with effector proteins such as Ca2+ - calmodulin . Here we have used the yeast two - hybrid system and identified a novel 34 kDa protein , termed NOSIP (**eNOS interaction protein**) , which avidly binds to the carboxyl terminal region of the **eNOS** oxygenase domain .

Figure 2: Medline abstract with all protein names emphasized.

their long form can provide additional useful information. Our intuition is that a method that could use this kind of information would show an increase in performance, especially when doing extraction from biomedical literature, where phenomena like repetitions and acronyms are pervasive.

The 5' upstream region (-448 / -443) of the human $dipeptidyl\,peptidase\,IV$ gene promoter containing a consensus E - box (CACGTG) was shown to bind upstream stimulatory factor using nuclear extracts from mouse (3T3) fibroblasts and the human intestinal and hepatic epithelial cell lines Caco - 2 and HepG2 .

Figure 3: Medline abstract with all protein names emphasized.

In this proposal, we describe how this type of document-level knowledge can be captured using Relational Markov Networks (RMNs) (Taskar, Abbeel, & Koller, 2002), a version of undirected graphical models which have already been successfully used to improve the classification of hyper-linked web pages. While other types of graphical models, such as Conditional Random Fields (CRFs) (Lafferty, McCallum, & Pereira, 2001), have modeled the entity recognition task as one of token classification, we take the different approach where candidate phrases in a document are classified according to the desired set of entity types. We then show how this phrase classification approach facilitates the modeling of correlations among labels of candidate entities, with the additional strength of phrase based features such as the actual text of the candidate entity, its capitalization pattern, or similarity with dictionary entries. Experimental results show that by factoring in global label correlations, the performance of the phrase classification approach is significantly improved.

In a typical application of CRFs, influences between the labels of consecutive tokens are the only correlations considered. This leads to a sequence labeling scenario, in which inference can be done efficiently using dynamic programming algorithms. Compared with CRFs, the increased representational power of RMNs comes at a cost in the time complexity of the inference algorithms. In our initial work, we resorted to approximate inference, based on an algorithm which has already exhibited competitive performance in other applications (Murphy, Weiss, & Jordan, 1999). However, in subsequent work we have discovered that exact inference for the phrase classification approach can be done efficiently, if no correlations between different candidate entities are to be considered. Moreover, by adding a carefully selected subset of document-level correlations, the same exact inference algorithm can be updated so that its running time remains linear in the number of candidate entities. We present how to select the correlations that are to be included in the model, and prove the linear complexity of the inference algorithm when run on the resulting structure.

As short-term goals, we intend to compare the exact inference / limited set of correlations approach with the original approach based on approximate inference / full set of correlations. Another direction that might lead to improved results is that of using a generalized version of the belief propagation algorithm, where

messages are passed between sets of nodes, at additional computational cost (Yedidia, Freeman, & Weiss, 2000), or using alternative approximate-inference methods.

Long-term goals include:

- Applying the same approach to other natural language tasks that may benefit from document-level correlations. Two examples are word sense disambiguation (WSD) and part-of-speech (POS) tagging. In WSD, the one sense per discourse hypothesis has been previously used by Yarowsky in (Yarowsky, 1995). The RMN framework however is able to incorporate this type of knowledge in a more probabilistic, sound manner. As for the task of POS tagging, while the benefit of using correlations between tags of repeated words is debatable if the tagger is trained and tested on documents from the same corpora, given its already competitive performance, it has nevertheless the potential of reducing the number of tagging errors on texts from different corpora.
- Using features based on similarities with existing dictionary entries. Such features can be incorporated in our phrase classification approach in a natural, straightforward manner.
- The "web as a corpus" is still an under-utilized idea. In this context, textual patterns that disambiguate the type of a candidate entity can be provided to a web search engine, so that statistics derived from the number of returned hits may be used in order to increase the IE system's performance.
- Syntactic information has already been proved to increase the accuracy on the task of relation extraction. Besides designing an IE system for extracting relations specific to biomedical entities, we intend to leverage entity recognition through the use of features derived from syntactic parses. Some of these features have the benefit of encoding long-range dependencies which cannot be captured from a flat representation of sentences.
- The previous three goals can be seen as part of the effort to design a general framework that would allow the use of information from various knowledge sources in order to increase the final IE system's performance. We intend to increase the robustness of this approach through the use of an efficient feature selection algorithm.

2 Background and Related Work

The task of automatically constructing information extractors has received a lot of attention in the past decade, and as such we observe a high diversity in the proposed approaches and the learning algorithms used therein. Nevertheless, a careful analysis reveals that most of these systems can be classified into two basic types of approaches:

- **Token Classification**: Word tokens in a document are sequentially classified as being inside or outside of a given named entity. Named entities are extracted by doing token classification and then assembling maximally contiguous sequences of inside tokens.
- **Phrase Classification**: Candidate phrases from a document are classified as to whether they are instances of some entity types or not. This can be done by either learning a multi-class classifier, in which case the number of classes is equal with the number of entity types plus one (for non-entity phrases), or by separately learning sets of extraction patterns, one set of patterns for each of the entity types.

2.1 Phrase Classification Approaches

Relational learning has been one of the learning paradigms used in some of the early IE systems, such as Rapier (Califf & Mooney, 1999) and SRV (Freitag, 1998). Both systems belong to the phrase classification approach.

2.1.1 Rapier

In RAPIER, the IE task is defined in terms of filling the slots contained in a template. A template specifies a particular type of event, such as joint ventures, corporate acquisitions, or job offerings. For example, a job offering template contains slots for title, salary, area of expertise, OS platform required, or job location. The training data consists of filled templates, one template per document. During testing, the IE system fills the template slots with data extracted from the document.

For each template slot, a set of rules is learned in a bottom-up fashion, with each rule composed of patterns that can make use of limited syntactic information. More exactly, the extraction rules consist of three parts:

- 1. A pre-filler pattern that matches text immediately preceding the slot filler,
- 2. A pattern that matches the actual field, and
- 3. A post-filler pattern that matches the text immediately following the slot filler.

Each pattern is a sequence of pattern elements of one of two types: *pattern items* and *pattern lists*. A pattern item matches exactly one word that satisfies its constraints. A pattern list has a maximum length N and matches 0 to N words, each satisfying a set of constraints. Besides constraining on words and their part-of-speech tags, Rapier can also incorporate semantic class information, such as that provided by WordNet (Miller, 1991). Consequently, each constraint is represented as a disjunctive list of one or more words, tags or WordNet synsets.

During testing, phrases are extracted by matching them against the set of rules learned for each slot. For the template filling task, extracted phrases which are duplicated are ignored, however the system can be easily modified to work in a named entity scenario, such that the output contains all extracted phrases, duplicates included. Because each pattern is designed to match phrases, we can view Rapier as belonging to the generic class of phrase classification approaches.

2.1.2 SRV

SRV (Freitag, 1998) too is based on a relational learning procedure. Like FOIL (Quinlan, 1990), it proceeds in a top-down fashion, starting with the entire set of examples - all negative examples and any positive examples not covered by already induced rules. At each step it greedily adds predicates, trying to cover as many positive, and as few negative examples as possible. There is a set of predefined predicate templates including tests on the length of the candidate entity or tests on features of tokens inside the candidate phrases. Token features are predefined too and come in two categories:

- *simple* features such as the word, its capitalization pattern, binary features testing whether the token is a punctuation sign, or a number.
- two *relational* features the previous and the next tokens.

As with any phrase classification approach, SRV needs to address the issue of searching through a typically huge negative examples set. The authors do this by handling negative examples implicitly, on a token-by-token basis – examples are indexed based on the tokens they contain. Because a token is generally shared by many candidate phrases, this leads to a more tractable search method.

2.2 Token Classification Approaches

2.2.1 Hidden Markov Models

Another class of approaches to learning IE systems is based on Hidden Markov Models (HMMs) (Rabiner, 1989). HMMs have been successfully used for speech recognition before becoming a model of choice for other natural language tasks such as POS tagging or named entity recognition. An HMM can be defined as the stochastic version of a finite state automaton. Thus, there is a set of states (hidden), with transitions between them. Given a state, there is a probability distribution over all possible transitions from that state. Symbols can be generated from any state, one symbol at a time, based on a symbol emission distribution. In a typical application of HMMs, a sequence of symbols is given, together with an HMM that is assumed to have produced it. The generative process by which the HMM produces a string of symbols starts by choosing a distinguished state (referred to as a starting state), then transitioning to another state according the the corresponding transition probability. This process of transitioning from one state to another continues until it reaches another distinguished state (referred to as the final state). Each time a transition is made from a state, a symbol is generated according to that state's symbol emission probability distribution. Graphically, an unrolled HMM can be represented as a directed graph, as in Figure 4. In this and all subsequent figures, the *X* symbols are used to denote observations, while *Y* symbols refer to hidden variables (states or labels).



Figure 4: Unrolling an HMM as a directed graphical model

One of the questions that an HMM inference algorithm is usually required to answer is what is the sequence of states that is most likely to have generated a given sequence of symbols. For example, in the case of POS tagging, each state corresponds to a POS tag, whereas symbols correspond to words. Given a particular sentence, the POS tagging is defined as the most likely sequence of states that generated the sentence.

HMMs are particularly attractive as they have a solid mathematical foundation, and the associated inference problem can be solved in time linear with the number of observed symbols using dynamic programming (the Viterbi algorithm). During learning, if the data is fully observable (e.g. labeled training data), the HMM parameters are simply set to their maximum likelihood estimates. If the data is only partially observable i.e. the states are hidden, the Baum-Welch algorithm, an instantiation of the more general Expectation Maximization (EM) algorithm (Dempster, Laird, & Rubin, 1977), can be used to find a set of parameters such that the likelihood function is locally optimized. IE systems based on HMMs belong naturally to the category of token classification approaches. The most likely path through the Markov model leads to a tagging of the input symbols, and consequently entities are extracted by assembling maximal contiguous sequences of words which are tagged with the same entity tag.

There exist numerous IE systems based on HMMs, and with them a whole diversity of augmentations to the basic model was introduced in order to better address various aspects of the task, such as the need for adequate representational power, or how to deal with sparsity due to insufficient training data.

Nymble. Nymble (Bikel, Schwartz, & Weischedel, 1999) is one of the earliest learning systems for named entity recognition based on HMMs. It consists of an ergodic model with one state for each entity type, together with an additional state for tokens outside any entity. Inside each name-class state, words are generated based on a statistical bigram language model. The generation of name-classes (states) and words proceeds in three steps, which are repeated until the entire word sequence is observed.

- 1. Select a name-class, conditioning on the previous name-class and the previous word.
- 2. Generate the first word from inside that name-class, by conditioning on the current and previous name-classes.
- Generate subsequent words inside the current name-class, where each word is conditioned on its immediate predecessor.

In this model, words are generally assimilated with ordered pairs of words and word features $\langle w, f \rangle$, where features belong to a predefined set of features, similar with those used in the SRV system. This further exacerbates the problem of insufficient training data for estimating the model parameters. Consequently, the authors rely on a multi-level back-off scheme, with weights for each level of back-off set based on an empirical formula.

HMMs and Shrinkage. A different approach is proposed in (Freitag & McCallum, 1999), where a separate HMM is created for each of the extraction fields. The states in each HMMs are either background or target states. Prefix and suffix states are distinguished from other background states in order to account for distributional peculiarities in the case of tokens occurring before or after the target field. Similarly, because certain tokens tend to occur at the beginning or end of the fragment, the target state is expanded into an array of parallel paths of varying length. The problem of data sparsity is alleviated through the use of "shrinkage", a statistical technique which combines parameter estimates from data-sparse states of a complex model with estimates from data-rich states of a simpler model. The method relies on a hierarchy that represents expected similarity between parameter estimates, with the estimates of the complex model at the leaves. In the case of shrinkage for HMM, subsets of states having similar word emission distributions are connected to a common parent. Internal nodes in turn can share a common parent, thus encoding weaker similarities between the corresponding groups of states. Word emission probabilities associated with states high in the hierarchy become simpler than those for states below, with the top of each hierarchy corresponding to the uniform distribution. The "shrinkage-based" parameter estimate is defined as a linear interpolation of the estimates in all distributions from the leaf to the root. The corresponding mixture weights are optimized by running EM on a held out dataset.

HMMs and Structure Learning. The two recently discussed HMM-based systems start with a predefined model structure, and learning is used only in estimating the model parameters. For tasks in which the entities to be extracted are densely represented inside a document, as is the case with headers and research paper references, a single HMM containing states for all entity types may be more appropriate. Variability in the relative ordering of the fields can be captured in the model by allowing the same field to be represented by more than one state. Learning the structure of such a model is the focus of the approach described in (Seymore, McCallum, & Rosenfeld, 1999).

2.2.2 Discriminative Models

We have already distinguished between IE approaches based on token classification and approaches based on *phrase classification*. Another useful dichotomy, orthogonal to the previous one, is that of *generative* vs. discriminative models. All HMM models reviewed here are generative in the sense that they try to model both the observation and hidden state sequences. However, in most application of HMMs, the observations are given, the task being that of "decoding" the hidden state sequence. Therefore, a major drawback of generative models is that modeling effort is spent on observations, instead of being focused entirely on describing the state sequence. The attempt to model the observations while keeping the inference tractable has led to the *output independence assumption*, which stipulates that the current observation, given the current state, is independent of previous observations. Usually, in text applications, observations correspond to words, and consequently the output independence assumption is not fair enough. The mismatch between model assumptions and data becomes even more pronounced if overlapping features, such as word capitalization and part-of-speech, are added as observations. Another inadequacy (McCallum, Freitag, & Pereira, 2000) is due to the way parameters are estimated. In an HMM, parameters are set to maximize the likelihood of the observation sequence, while the task is that of predicting the state sequence given the observations. All these mismatches and limitations are eliminated in discriminative approaches, in which the conditional probability of state sequences given the observations lies at the core at the model.

2.2.3 Maximum Entropy Models

The Maximum Entropy (MaxEnt) (Berger, Della Pietra, & Della Pietra, 1996) principle has been widely used to create discriminative probabilistic models for natural language tasks. The classification problem is viewed in terms of a random process that produces an output value y from a finite set Y, based on the contextual information x, a member of a finite set X. In a token classification scenario, this means associating a tag y to each text token, whereas the context x is derived from the text centered at the current token position. In maximum entropy modeling we are looking for a probability distribution p(y|x) that satisfies a set of constraints $C_i \in C$ derived from a collection of user specified features $f_i(x, y) \in F$. Each feature is expressed as a binary function based on the context x at the current token position and its proposed classification y. For example, a useful feature in tagging for named entity recognition is the capitalization of the token to be classified, and it can be expressed as follows:

$$f_i(x, y) = \begin{cases} 1 & \text{if current token is capitalized & } y = Inside, \\ 0 & \text{otherwise.} \end{cases}$$

The constraint C_i associated with a feature function f_i is expressed simply by imposing that the expected value of f_i under the target distribution p(y|x) be the same as the expected value of f_i under the empirical distribution $\tilde{p}(x, y)$ (derived from the training data):

$$C_i
ightarrow \sum_{x,y} ilde{p}(x,y) f_i(x,y) = \sum_{x,y} ilde{p}(x) p(y|x) f_i(x,y)$$

Out of a potentially infinite number of probability distributions p(y|x) satisfying a particular set of constraints, the maximum entropy principle dictates that we select the most "uniform" distribution, where

a formal measure for the "uniformity" of a distribution is given by the information theoretic notion of conditional entropy (Cover & Thomas, 1991):

$$H(Y|X) = -\sum_{x,y} ilde{p}(x) p(y|x) \log p(y|x)$$

Based on the concept of duality from constrained optimization, it can be shown that the distribution p(y|x) satisfying the constraints C_i , and which also minimizes the conditional entropy H(Y|X), is a member of the exponential family:

$$p(y|x) = \frac{1}{Z(x)} exp\left(\sum_{i} \lambda_i f_i(x, y)\right)$$

where $Z(x) = \sum_{y} exp(\sum_{i} \lambda_{i} f_{i}(x, y))$ is a normalizing constant. An additional compelling justification for the maximum entropy principle is that the resulting distribution is also the model which, among all log-linear models of the above form, maximizes the likelihood of the parameters given the training sample.

In (Ratnaparkhi, 1996), the authors describe a maximum entropy approach to part-of-speech tagging in which they introduce a feature template f(a, b) which relates the tags of two consecutive tokens:

$$f\langle a,b\rangle(x_t,y_t,y_{t-1}) = \begin{cases} 1 & \text{if } y_{t-1} = a \& y_t = b \\ 0 & \text{otherwise.} \end{cases}$$

They also define a similar feature template relating the tags of three consecutive tokens. Computing the highest probability label for each token, from left to right, does not necessarily lead to the most likely sequence of tags. To alleviate this, the authors use a beam search procedure, in which they consider tokens from left to right, keeping at each position the five sequences of tags concentrating the most probability mass. A more rigorous approach, which was later used in maximum entropy models for named entity recognition (Chieu & Ng, 2003), is to use a Viterbi-like algorithm for decoding, which guarantees finding the most likely labeling of the entire sequence of words.

Maximum Entropy Markov Models. This new type of features, relating tags in consecutive positions, suggests a class of maximum entropy models in which binary features may include a test on the class of the previous token, besides conditioning on the observed input context and the mandatory test on the class of the current token. Each such feature is uniquely identifiable by a condition g on the observed input x_t and the possible instantiations a and b for the current and previous tags, y_t and y_{t-1} , as follows:

$$f\langle g, a, b \rangle(x_t, y_t, y_{t-1}) = \begin{cases} 1 & \text{if } g(x_t) = 1 \& y_{t-1} = a \& y_t = b, \\ 0 & \text{otherwise.} \end{cases}$$

One "extreme" case is that when for any given input feature g, for each valid combinations of tags $\langle a, b \rangle$, the above defined compound feature $f\langle g, a, b \rangle$ is included in the model. This is a maximum entropy model in which the same set of input features g is associated with transitions between any two hidden states a and b. It can be shown that this type of model is in fact equivalent with a Maximum Entropy Markov Model (MEMM) (McCallum et al., 2000), which means that the same generic system that is currently used for learning a MaxEnt model, can also be used for learning an MEMM model by simply providing it with the appropriate set of features.

An MEMM (McCallum et al., 2000) creates an maximum entropy model for each state in the model. Thus, for a given state s', the framework learns an exponential model corresponding to the probability of transitioning to another state s from s', given the observation sequence o, i.e. p(s|s', o). Consequently, if



Figure 5: Unrolling an MEMM as a directed graphical model

the set of states is S, the MEMM will contain |S| exponential models. Finding the most likely sequence of states in this context can be done efficiently using a Viterbi-like algorithm. The procedure for learning the parameters is the same as in the MaxEnt case i.e. using Improved Iterative Scaling (Della Pietra, Della Pietra, & Lafferty, 1997) or a gradient based method (the likelihood function is concave, and the gradient is simply the difference between observed and expected feature counts).

2.2.4 Conditional Random Fields

A fundamental problem with MEMMs and other discriminative Markov models based on directed graphical models is that they are biased toward states with few successor states. This is the "label bias problem" (Lafferty et al., 2001), which in a more general form stipulates that states with low entropy next-state distributions will take little notice of observations. The maximum entropy model from (Ratnaparkhi, 1996) is subject to this problem too, as some of the features it uses are indirectly associated with transitions (they contain conditions on labels of consecutive tokens). The reason for this behavior stems from the fact that the same probability mass is allocated for modeling the labeling decision at each position in the sequence. A principled solution to this problem is that of Conditional Random Fields (Lafferty et al., 2001), where a single probability distribution is learned, one that models the joint probability of a label sequence given a sequence of observations. Informally, this can be viewed as a finite state model with unnormalized transition probabilities. Therefore, some transitions may contribute more than others to the overall score, depending on the corresponding observations.



Figure 6: Unrolling a CRF as an undirected graphical model

Inference in CRFs can be done efficiently by accommodating the corresponding forward-backward or Viterbi algorithms used for HMMs (Rabiner, 1989). Learning the CRFs parameters can be cast as an optimization problem – the likelihood function is concave, thus a global maximum can be found efficiently using standard procedures, such as Improved Iterative Scaling (Della Pietra et al., 1997), or gradient based methods. We have started the list of token classification approaches with HMM models, which are generative and can be represented as directed graphical models. We have argued that conditional models are more appropriate for the tagging task, one of their benefits being that they allow for arbitrary, potentially overlapping features over the observation sequence. Consequently, we have described Maximum Entropy models, a class of conditional models which we have further shown that it subsumes Maximum Entropy Markov Models, a particular type of conditional Markov models. Although these conditional models offer increased representational power when compared with HMMs (their generative counterpart), they are all plagued by the "label bias problem". This is particularly troublesome, as the problem does not occur with HMMs. The solution came in the form of Conditional Random Fields, a type of undirected graphical models especially suited for labeling sequences, which overcomes the label bias problem by modeling the joint probability over the entire label sequence given the observation sequence. In the next sections we describe a generic type of undirected graphical models called Relational Markov Networks (RMNs) (Taskar et al., 2002) which can model more general types of label correlations, and are consequently a suitable framework for our initial approach to "collective information extraction".

2.3 Markov Random Fields

Graphical models offer an intuitive representation of conditional independence between domain variables. They come in two main flavors:

- **Directed Models** well suited to represent temporal and causal relationships (Bayesian Networks, Neural Networks, HMMs)
- Undirected Models appropriate for representing statistical correlations between variables (Markov Networks such as CRFs, RMNs, Boltzman Machines)

Markov Random Fields (Markov Networks) are a special class of undirected graphical models. Below is their definition, based on the following notation:

- V = a set of vertices used to denote random variables
- G = (V, E) an undirected graph
- N(v) = the set of neighbors of vertex $v \in V$

Definition 1 V is said to be a Markov Random Field with respect to G if for any vertex, its value depends only on its neighbors i.e. $P(V_i|V - V_i) = P(V_i|N(V_i)), \forall V_i \in V$

For the discriminative version, assume X is the set of observed variables, and Y is the set of hidden variables, such that $V = X \cup Y$.

Definition 2 *V* is said to be a Conditional Markov Random Field with respect to G if $P(Y_i|X, Y - Y_i) = P(Y_i|X, N(Y_i)), \forall Y_i \in Y$

Markov Random Fields characterize the underlying undirected graphical model via a local property, namely the Markov assumption. On the other hand, Gibbs Random Fields, which are going to be defined next, use a global property to characterize the corresponding graphical model. The corresponding notation follows below:

- V = a set of vertices which stand for random variables
- G = (V, E) an undirected graph
- C(G) = the set of cliques in G
- V_c = the set of vertices in a clique $c \in C$
- $\phi = \{\phi_c : V_c \to R_+, c \in C(G)\}$ a set of *clique potentials*

Definition 3 V is said to be a Gibbs Random Field with respect to G if $P(V) = \frac{1}{Z} \sum_{c \in C(G)} \phi_c(V_c)$, where Z is a normalization constant.

Thus, a Gibbs Random Field is specified numerically by associating potentials with cliques in the graph. A clique potential is a function on the set of possible configurations of the clique, that associates a positive number with each configuration. The joint probability distribution over all vertices in the graph is obtained by taking a product over the clique potentials.

For the discriminative version, assume X is the set of observed variables, and Y is the set of hidden variables, such that $V = X \cup Y$, and similarly, for every clique $c \in C(G)$, let $V_c = X_c \cup Y_c$.

Definition 4 V is said to be a Conditional Gibbs Random Field with respect to G if $P(Y|X) = \frac{1}{Z(X)} \sum_{c \in C(G)} \phi_c(X_c, Y_c)$, where Z(X) is a normalization constant.

Therefore, whereas a Markov Random Field is an undirected graphical model characterized by a local property, a Gibbs Random Field is an undirected graphical model constrained by a global property e.g. the Gibbs distribution. The following theorem stipulates that the two types of graphical models are in fact equivalent.

Theorem 1 (Hammersley & Clifford, 1971) V is a (conditional) MRF with respect to G if and only if V is a (conditional) GRF with respect to G.

Consequently, one can create a Markov Random Field by specifying an underlying probability distribution that factorizes into potentials over all maximal cliques in the graph.

2.4 Relational Markov Networks

Relational Markov Networks (Taskar et al., 2002) are conditional Markov random fields augmented with a set of *clique templates*. A clique template specifies which vertices are to be connected in a clique, associating the same clique potential with all cliques that it creates in the graph. Thus, a clique template provides at the same time a procedure for creating edges in the graph, and a mechanism for tying parameters (clique potentials) in the model.

In (Taskar et al., 2002), the RMN framework was introduced in order to model correlations between the class labels of hyperlinked web pages – pages which are hyperlinked tend to have the same label. The clique template responsible for this type of correlations is detailed below:

- Clique Creation Add an edge (a 2-node clique) between the labels of any two hyperlinked web pages.
- Clique Potentials To all edges created by this template, associate the same potential function φ. If the number of possible class labels is N, then φ can be specified as an N × N table of positive real values i.e. φ : {1, 2, ..., N} × {1, 2, ..., N} → R₊.

Figure 7 shows a sample RMN, where the above clique template creates eight edges between the labels Y of hyperlinked web pages X. The same potential ϕ is associated with all these edges. Other clique templates are responsible for creating edges between each label Y_i and the corresponding local context features in X_i .



Figure 7: An RMN unrolled, with cliques between hyperlinked web pages.

The CRFs, as previously illustrated in Figure 6, are therefore a particular type of RMNs, in which clique templates create 3-node cliques between any two consecutive labels, Y_{t-1} and Y_t , and their corresponding contextual features X_t .

Given a set of potentials, doing inference with RMNs may refer to two things:

- 1. Computing the marginal probabilities for all hidden variables, or a proper subset of them.
- 2. Computing the most probable assignment of values to all hidden variables in the model.

For tree-structured models, the belief propagation algorithm (Pearl, 1988) computes the marginals over all hidden variables in time linear with the number of nodes and edges in the underlying graph. For graphs with cycles, however, exact inference algorithms, such as the join-tree algorithm, have a running time exponential in the size of the largest clique in the triangulated graph. An alternative to exact inference is to do approximate inference using loopy belief propagation, which has shown reasonable performance in many practical applications (Murphy et al., 1999).

Learning with RMNs means computing the clique potential for each potential template, given training data where both the content attributes and the labels are observed. One alternative is to use a gradient based method in a Maximum Likelihood (ML) or Maximum A Posteriori (MAP) setting. For the last type of estimation, a "shrinkage" prior over the parameters is used, typically a zero-mean Gaussian. Because, in both cases, the objective function is concave, the optimization procedure is guaranteed to find a global maximum. An alternative learning method is to use *stochastic gradient ascent* in the form of a Voted Perceptron (Collins, 2002). In this case, the objective function is contave on the Most Probable Explanation (MPE) labeling, instead of computing the full feature count expectation. Nevertheless, inference is needed in both learning scenarios, either for computing marginals over subsets of hidden variables, of for deriving the MPE labeling.

Viewed from the RMN perspective, CRFs are a special type of linear-chain undirected graphical models, and, as with any linear-chain or tree-structured graphical models, both exact inference and parameter estimation can be solved efficiently.

3 Completed Research

Of all IE systems mentioned in the previous section, that of (Seymore et al., 1999) is able to model influences between various types of entities based on the order in which they occur in the document – in headers of research papers, for example, the author's name usually comes after the title. This type of order-based correlations is captured by learning an HMM structure in which the same entity type may be associated with multiple states in the model, while the set of transitions reflects the order in which various entity types occur in the training data.

There have been some previous attempts to use global information from repetitions, acronyms, and abbreviations during extraction. In (Chieu & Ng, 2003), a set of global features are used to improve a Maximum-Entropy tagger; however, these features do not fully capture the mutual influence between the labels of acronyms and their long forms, or between entity repetitions. In particular, they only allow earlier extractions in a document to influence later ones and not vice-versa.

In this section we are going to introduce a collective approach to Information Extraction which will allow the incorporation of arbitrary correlations between the labels of potential extractions from the same document. For this, we shall use the RMN framework to do extraction by phrase classification.

3.1 Candidate Entities

Doing phrase classification requires a set of phrases to start with. Throughout this document, we will use the terms *candidate entities*, *candidate extractions*, or *candidate phrases* to refer to the set of phrases that are to be classified as being valid extractions or not. Considering as candidate entities all contiguous word sequences from a document would lead to a quadratic number of phrases, which would adversely affect the time complexity of the extraction program. Various heuristics exist however which can significantly reduce the size of the candidate set, and some of them are listed below:

- **H1:** In general, named entities have limited length. Therefore, one simple way of creating the set of candidate phrases is to compute the maximum length of all annotated entities in the training set, and then consider as candidates all word sequences whose length is up to this maximum length. This is also the approach followed in SRV (Freitag, 1998).
- H2: In the task of extracting protein names from Medline abstracts, we noticed that, like most entity names, almost all proteins in our data are base noun phrases or parts of them. Therefore, such substrings are used to determine candidate entities. To avoid missing options, we adopt a very broad definition of base noun phrase – a maximal contiguous sequence of tokens whose POS tags are from {"JJ", "VBN", "VBG", "POS", "NN", "NNS", "NNP", "NNPS", "CD", "-"}, and whose last word (the head) is tagged either as a noun, or a number. Candidate extractions then consist of base NPs, together with all their contiguous subsequences headed by a noun or number.
- H3: The CoNLL 2003 English corpus (Tjong Kim Sang & De Meulder, 2003) contains four types of named entities: persons (PER), locations (LOC), organizations (ORG), and other (MISC). A more appropriate heuristic in this case is to consider as candidates all sequences of proper names, potentially interspersed with prepositions, commas, conjunctions or definite articles.

Table 1 below shows the candidate entities generated by H1 and H2 on a fragment from a Medline abstract. Similarly, Table 2 shows candidate entities generated by H1 and H3 on a fragment from a CoNLL document. Both H2 and H3 are strong heuristics, in the sense that they drastically reduce the number of

candidate entities. In the next sections, we shall focus on the task of extracting protein names from Medline abstracts.

	"the control of human ribosomal protein L22 ($rpL22$) "						
	\circ the \circ the control \circ the control of \circ the control of human \circ the control of						
H1	human ribosomal o o ribosomal o ribosomal protein o ribosomal protein L22						
	\circ ribosomal protein L22 (\circ ribosomal protein L22 (rpL22 \circ \circ L22 \circ L22 (\circ						
◦ L22 (rpL22 ◦ L22 (rpL22) ◦ ◦ rpL22 ◦ rpL22) ◦) ◦							
H2 • control • human ribosomal protein • human ribosomal protein L2							
	o ribosomal protein o ribosomal protein L22 o protein L22 o L22 o rpL22 o						

Table 1: Candidate Extractions: Medline.

" I	"Israel gave Palestinian President Yasser Arafat permission on Thursday"						
	∘ Israel ∘ Israel gave ∘ Israel gave Palestinian ∘ Israel gave Palestinian						
H1	$President \circ \circ \textbf{Palestinian} \circ Palestinian President \circ Palestinian President Yasser$						
• Palestinian President Yasser Arafat • • Yasser • Yasser Arafat • Pres							
	Yasser Arafat permission $\circ \dots \circ$ on \circ on Thursday \circ Thursday \circ						
H3	∘ Israel ∘ Palestinian ∘ Palestinian President ∘ Palestinian President Yasser						
	 Palestinian President Yasser Arafat President President Yasser 						
	 President Yasser Arafat Yasser Yasser Arafat Arafat 						

Table 2: Candidate Extractions: CoNLL.

3.2 Entity Features

The set of features associated with each candidate is based on the feature templates introduced in (Collins, 2002), used there for training a ranking algorithm on the extractions returned by a maximum-entropy tagger. Many of these features use the concept of *word type*, which allows a different form of token generalization than POS tags. The *short type* of a word is created by replacing any maximal contiguous sequences of capital letters with 'A', of lower-case letters with 'a', and of digits with '0'. For example, the word *TGF-1* would be mapped to type *A-0*.

Consequently, each token position *i* in a candidate extraction provides three types of information: the word itself w_i , its POS tag t_i , and its short type s_i . The full set of features types is listed in Table 3, where we consider a generic candidate extraction as a sequence of n + 1 words $w_0 w_1 \dots w_n$.

Each feature template instantiates numerous features. For example, the candidate extraction 'HDAC1 enzyme' has the head word HD=enzyme, the short type $ST=A0_a$, the prefixes PF=A0 and $PF=A0_a$, and the suffixes SF=a and $SF=A0_a$. All other features depend on the left or right context of the entity. Feature values that occur less than three times in the training data are filtered out.

3.3 The RMN Framework for Entity Recognition

Given a collection of documents D, we associate with each document $d \in D$ a set of candidate entities d.E, in our case a restricted set of token sequences from the document (Section 3.1). Each entity $e \in d.E$ is

Description	Feature Template			
Head Word	$w_{(n)}$			
Text	$w_{(0)} - w_{(1)} - \dots - w_{(n)}$			
Short Type	$s_{(0)}$ $-s_{(1)}$ $-\cdots$ $-s_{(n)}$			
Bigram Left	$w_{(-1)} - w_{(0)} = w_{(-1)} - s_{(0)}$			
(4 bigrams)	$s_{(-1)} - w_{(0)} = s_{(-1)} - s_{(0)}$			
Bigram Right	$w_{(n)}-w_{(n+1)}$ $w_{(n)}-s_{(n+1)}$			
(4 bigrams)	$s_{(n)}-w_{(n+1)}$ $s_{(n)}-s_{(n+1)}$			
Trigram Left	$w_{(-2)} - w_{(-1)} - w_{(0)} \dots$			
(8 trigrams)	$s_{(-2)}$ - $s_{(-1)}$ - $s_{(0)}$			
Trigram Right	$w_{(n)}$ - $w_{(n+1)}$ - $w_{(n+2)}$			
(8 trigrams)	$s_{(n)}$ - $s_{(n+1)}$ - $s_{(n+2)}$			
POS Left	$t_{(-1)}$			
POS Right	$t_{(n+1)}$			
Prefix	$s_{(0)} s_{(0)} - s_{(1)} \dots$			
(n+1 prefixes)	$s_{(0)}$ - $s_{(1)}$ -···- $s_{(n+1)}$			
Suffix	$s_{(n)}$ $s_{(n-1)}$ $\overline{s_{(n)}}$			
(n+1 suffixes)	$s_{(0)} - s_{(1)} - \cdots - s_{(n+1)}$			

Table 3: Feature Templates.

characterized by a predefined set of boolean attributes e.F (Section 3.2), the same for all candidate entities. One particular attribute is e.label which is set to 1 if e is considered a valid extraction, and 0 otherwise. In this document model, labels are the only hidden variables, and the inference procedure will try to find a most probable assignment of values to labels, given the current model parameters.

Each document is associated with an undirected graphical model, with nodes corresponding directly to entity attributes, one node for each attribute of each candidate entity in the document. The set of edges is created by matching *clique templates* against the entire set of entities d.E. A clique template is a procedure that finds all subsets of entities satisfying a given constraint, after which, for each entity subset, it connects a selected set of attribute nodes so that they form a clique.

Formally, there is a set of clique templates C, with each template $c \in C$ specified by:

- 1. A matching operator M_c for selecting subsets of entities, $M_c(E) \subseteq 2^E$
- 2. A selected set of features $S_c = \langle X_c, Y_c \rangle$, the same for all subsets of entities returned by the matching operator. X_c denotes the observed features, while Y_c refers to the hidden labels.
- A clique potential φ_c which gives the compatibility of each possible configuration of values for the features in S_c, s.t. φ_c(s) ≥ 0, ∀s ∈ S_c.

Given a set E of nodes, $M_c(E)$ consists of subsets of entities whose attribute nodes S_c are to be connected in a clique. In previous applications of RMNs, the selected subsets of entities for a given template have the same size; however, some of our clique templates may match a variable number of entities. The set S_c may contain the same attribute from different entities. Usually, for each entity in a matching set, its label is included in S_c . All these will be illustrated with examples in Sections 3.4 and 3.5 where the clique templates used in our model are described in detail.

Depending on the number of hidden labels in Y_c , we define two categories of clique templates:

- Local Templates are all templates $c \in C$ for which $|Y_c| = 1$. They model the correlations between an entity's observed features and its label.
- Global Templates are all templates $c \in C$ for which $|Y_c| > 1$. They capture influences between multiple entities from the same document.

After the graph model for a document d has been completed with cliques from all templates, the probability distribution over the random field of hidden entity labels d.Y given the observed features d.X is given by the Gibbs distribution:

$$P(d.Y|d.X) = \frac{1}{Z(d.X)} \prod_{c \in C} \prod_{G \in M_c(d.E)} \phi_C(G.X_c, G.Y_c)$$

$$\tag{1}$$

where Z(d.X) is the normalizing partition function:

$$Z(d.X) = \sum_{Y} \prod_{c \in C} \prod_{G \in M_c(d.E)} \phi_C(G.X_c, G.Y_c)$$
⁽²⁾

3.4 Local Clique Templates

As described in the previous section, the role of local clique templates is to model correlations between an entity's observed features (see Table 3) and its label. If, after filtering, we are left with h distinct boolean features f_i , one way to model these correlations is to introduce h local (clique) templates $LT_1, LT_2, ..., LT_h$. A template LT_i would then be defined as follows:

- 1. The matching operator M_i is set to match any single-entity set $\{e\}$.
- 2. The collection of attributes S_i corresponding to a singleton entity set $\{e\}$ is defined to be $S_i = \langle X_i, Y_i \rangle = \langle \{e.f_i\}, \{e.label\} \rangle$. This amounts to introducing in the RMN graph *h* attribute nodes for each candidate entity, which are to be connected by the *h* local templates to the corresponding entity label node. The 2-node cliques created by all *h* templates around one entity are illustrated in Figure 8.
- 3. The potential ϕ_i associated with all 2-node cliques created by template LT_i would consist in a 2 × 2 table (as both *e*. f_i and *e*.*label* have cardinality 2 assuming only one entity type is to be extracted, we need only two values for the label attribute).



Figure 8: RMN generated by local templates.

Each entity has the label node connected to its own set of h binary feature nodes. This leads to an excessive number of nodes in the model, most of which have value zero. The number of nodes can be reduced if, for each entity, we include in the graphical model only those nodes for which the corresponding feature variable has value 1. Consequently, the table associated with the local potential will be reduced from 4 to 2 values, specifying now the compatibility between that feature and the two possible values for the entity label.

Factor Graphs. An alternative, useful representation for Markov random fields is provided by factor graphs (Kschischang, Frey, & Loeliger, 2001). These are bipartite graphs which express how a global function of many variables (the probability P(d.Y|d.X) in Equation 1) factors into a product of local functions (the potentials $\phi_C(G.X_c, G.Y_c)$ in Equation 1). Factor graphs subsume many different types of graphical models, including Bayesian networks and Markov random fields. The sum/max-product algorithm used for inference in factor graphs generalizes a wide variety of algorithms including the forward/backward algorithm, the Viterbi algorithm, and Pearl's belief propagation algorithm (Pearl, 1988). To obtain the factor graph for a given Markov random field, we copy all original nodes from the MRF, referred henceforth as *variable nodes*, and create a *potential node* for each instantiated clique potential. Each potential node is then linked to all variable nodes from the associated clique.

In the case of local clique potentials, given that all feature nodes have value 1, we can eliminate them from the equivalent factor graph representation. What is left then is a variable node for the entity label, together with nodes for potential functions, one potential node for each entity feature whose value has been observed to be 1. As an example, Figure 9 shows that part of the factor graph which is generated around the entity label for 'HDAC1 enzyme' (with variable nodes figured as empty circles and potential nodes figured as black squares).



Figure 9: Factor Graph for local templates.

Note that the factor graph above has an equivalent RMN graph consisting of a one-node clique only, on which it is hard to visualize the various potentials involved. There are cases where different factor graphs may yield the same underlying RMN graph, which makes the factor graph representation preferable.

3.5 Global Clique Templates

Global clique templates enable us to model hypothesized influences between entities from the same document. They connect the label nodes of two or more entities, which, in the factor graph, translates into potential nodes connected to at least two label nodes. In our experiments we use three global templates:

Overlap Template (OT): No two entity names overlap in the text i.e if the span of one entity is $[s_1, e_1]$ and the span of another entity is $[s_2, e_2]$, and $s_1 \le s_2$, then $e_1 < s_2$.

Repeat Template (RT): If multiple entities in the same document are repetitions of the same name, their labels tend to have the same value (i.e. most of them are protein names, or most of them are not protein

names). Later we discuss situations in which repetitions of the same protein name are not tagged as proteins, and design an approach to handle this.

Acronym Template (AT): It is common convention that a protein is first introduced by its long name, immediately followed by its short-form (acronym) in parentheses.

3.5.1 The Overlap Template

The definition of a *candidate extraction* from Section 3.1 leads to many overlapping entities. For example, 'glutathione S - transferase' is a base NP, and it generates five candidate extractions: 'glutathione', 'glutathione S', 'glutathione S - transferase', 'S - transferase', and 'transferase'. If 'glutathione S - transferase' has label-value 1, the other four entities should all have label-value 0, because they overlap with it.

This type of constraint is enforced by the overlap template as follows:

- 1. The M_{OT} operator matches any two overlapping candidate entities $\{e_1, e_2\}$.
- 2. The set of attributes S_{OT} selected by this template for two overlapping entities $\{e_1, e_2\}$ is $S_{OT} = \langle X_{OT}, Y_{OT} \rangle = \langle \emptyset, \{e_1.label, e_2.label\} \rangle$. This translates in the factor graph into a potential node connected to the two selected label nodes.
- 3. The potential function ϕ_{OT} is set so that at most one of the overlapping entities can have label-value 1, as illustrated in Table 4.

ϕ_{OT}	$e_1.label = 0$	$e_1.label = 1$
$e_2.label = 0$	1	1
$e_2.label = 1$	1	0

Table 4: Overlap Potential.

Continuing with the previous example, because 'glutathione S' and 'S - transferase' are two overlapping entities, the factor graph model will contain an overlap potential node connected to the label nodes of these two entities.

An alternative solution for the overlap template is to create a potential node for each token position that is covered by at least two candidate entities in the document, and connect it to their label nodes. The difference in this case is that the potential node will be connected to a variable number of entity label nodes. However this second approach has the advantage of creating fewer potential nodes in the document factor graph, which results in faster inference.

3.5.2 The Repeat Template

We could specify the potential for the repeat template in a similar 2×2 table, this time leaving the table entries to be learned, given that assigning the same label to repetitions is not a hard constraint. However we can do better by noting that the vast majority of cases where a repeated protein name is not also tagged as a protein happens when it is part of a larger phrase that *is* tagged. For example, 'HDAC1 enzyme' is a protein name, therefore 'HDAC1' is not tagged in this phrase, even though it may have been tagged previously in the abstract where it was not followed by 'enzyme'. We need a potential that allows two entities with the same text to have different labels if the entity with label-value 0 is inside another entity with label-value 1. But a candidate entity may be inside more than one "including" entity, and the number of including entities may vary from one candidate extraction to another. Using the example from Section 3.5.1, the candidate entity 'glutathione' is included in two other entities: 'glutathione S' and 'glutathione S - transferase'.

In order to instantiate potentials over variable number of label nodes, we introduce a *logical OR clique* template that matches a variable number of entities. When this template matches a subset of entities $e_1, e_2, ..., e_n$, it will create an auxiliary OR entity e_{OR} , with a single attribute $e_{OR}.label$. The potential function ϕ_{OR} is set so that it assigns a non-zero potential only when $e_{OR}.label = e_1.label \lor e_2.label \lor ... \lor$ $e_n.label$. The cliques are only created as needed, e.g. when the auxiliary OR entity is required by repeat and acronym clique templates.

Figure 10 shows the factor graph for a sample instantiation of the repeat template using the OR template. Here, u and v represent two same-text entities, $u_1, u_2, ..., u_n$ are all entities that include u, and $v_1, v_2, ..., v_m$ are entities that include v. To avoid clutter, all entities in this and subsequent factor graphs stand for their corresponding label features. The potential function ϕ_{RT} can either be preset to prohibit unlikely label configurations, or it can be learned to represent an appropriate soft constraint. In our experiments, it was learned since this gave slightly better performance.



Figure 10: Repeat Factor Graph.

Following the previous example, suppose that the phrase 'glutathione' occurs inside two base NPs in the same document, 'glutathione S - transferase' and 'glutathione antioxidant system'. Then the first occurrence of 'glutathione' will be associated with the entity u, and correspondingly its including entities will be u_1 = 'glutathione S' and u_2 = 'glutathione S - transferase'. Similarly, the second occurrence of 'glutathione' will be associated with the including entities will be v_1 = 'glutathione antioxidant' and v_2 = 'glutathione antioxidant system'.

3.5.3 The Acronym Template

One approach to the acronym template would be to use an extant algorithm for identifying acronyms and their long forms in a document, and then define a potential function that would favor label configurations in which both the acronym and its definition have the same label. One such algorithm is described in (Schwartz & Hearst, 2003), achieving a precision of 96% at a recall rate of 82%. However, because this algorithm would miss a significant number of acronyms, we have decided to implement a softer version as follows: detect all situations in which a single word is enclosed between parentheses, such that the word length is at least 2 and it begins with a letter. Let v denote the corresponding entity. Let $u_1, u_2, ..., u_n$ be all entities that end exactly before the open parenthesis. If this is a situation in which v is an acronym, then one

of the entities u_i is its corresponding long form. Consequently, we use a logical OR template to introduce the auxiliary entity u_{OR} , and connect it to v's node label through an acronym potential ϕAT , as illustrated in Figure 11.



Figure 11: Acronym Factor Graph.

For example, consider the phrase 'the antioxidant superoxide dismutase - 1 (SOD1)', where both 'superoxide dismutase - 1' and 'SOD1' are tagged as proteins. 'SOD1' satisfies our criteria for acronyms, thus it will be associated with the entity v in Figure 11. The candidate long forms are u_1 = 'antioxidant superoxide dismutase - 1', u_2 = 'superoxide dismutase - 1', and u_3 = 'dismutase - 1'.

3.6 Inference in Factor Graphs

There are two problems that need to be addressed when working with RMNs:

- 1. Inference: Usually, two types of quantities are needed from an RMN model:
 - The marginal distribution for a hidden variable, or for a subset of hidden variables in the graphical model.
 - The most probable assignment of values to all hidden variables in the model.
- Learning: As the structure of the RMN model is already defined by its clique templates, learning refers to finding the clique potentials that maximize the likelihood over the training data. Inference is usually performed multiple times during the learning algorithm, which means that an accurate, fast inference procedure is doubly important.

In our setting, given the clique potentials, the inference step for the factor graph associated with a document involves computing the most probable assignment of values to the hidden labels of all candidate entities:

$$d.Y^* = \arg\max_{d.Y} P(d.Y|d.X)$$
(3)

where P(d.Y|d.X) is defined as in Equation 1. A brute-force approach is excluded, since the number of possible label configurations is exponential in the number of candidate entities. The sum-product algorithm (Kschischang et al., 2001) is a message-passing algorithm that can be used for computing the marginal distribution over the label variables in factor graphs without cycles, and with a minor change (replacing the sum operator used for marginalization with a max operator) it can also be used for deriving the most probable label assignment. In our case, in order to get an acyclic graph, we would have to use local templates only. However, it has been observed that the algorithm often converges in general factor graphs, and when it converges, it gives a good approximation to the correct marginals. The algorithm works by altering the

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belief at each label node by repeatedly passing messages between the node and all potential nodes connected to it (Kschischang et al., 2001).

The time complexity of computing messages from a potential node to a label node is exponential in the number of label nodes attached to the potential. Since this "fan-in" can be large for OR potential nodes (and also for the second solution to overlap potential nodes), this step required optimization. Fortunately, due to the special form of the OR and overlap potentials (high degree of sparsity), and the normalization before each message-passing step, these special cases can be computed in linear-time. For example, the formulae for computing the OR messages for the sum-product algorithm are shown in Figure 12 (to avoid clutter, *e* and ϕ stand for e_{OR} and ϕ_{OR} respectively).

$$\mu_{\phi \rightarrow e} \stackrel{e}{\underset{\mu_{e \rightarrow \phi}}{\longrightarrow}} \qquad \mu_{\phi \rightarrow e}(0) = \prod_{i=1}^{n} \mu_{e_i \rightarrow \phi}(0)$$

$$\mu_{\phi \rightarrow e}(1) = 1 - \mu_{\phi \rightarrow e}(0)$$

$$\mu_{\phi \rightarrow e_i}(0) = \mu_{\phi \rightarrow e}(1) + \frac{\mu_{\phi \rightarrow e}(0)}{\mu_{e_i \rightarrow \phi}(0)} (\mu_{\phi \rightarrow e}(0) - \mu_{\phi \rightarrow e}(1))$$

$$\mu_{\phi \rightarrow e_i}(1) = \mu_{\phi \rightarrow e}(1)$$

Figure 12: Messages in OR Factor Graph.

3.7 Learning Potentials in Factor Graphs

Following a maximum likelihood estimation, we shall use the log-linear representation of potentials:

$$\phi_C(G.X_c, G.Y_c) = exp\{\mathbf{w}_c \mathbf{f}_c(G.X_c, G.Y_c)\}$$
(4)

Let w be the concatenated vector of all potential parameters w_c . One approach to finding the maximumlikelihood solution for w is to use a gradient-based method, which requires computing the gradient of the log-likelihood with respect to potential parameters w_c . It can be shown that this gradient is equal with the difference between the empirical counts of f_c and their expectation under the current set of parameters w.

$$\nabla L(w,D) = \sum_{d \in D} f_c(d.X, d.Y) - \sum_{d \in D} \sum_{d.Y'} f_c(d.X, d.Y') P_w(d.Y'|d.X)$$
(5)

The expectation in the second term is expensive to compute, since it requires summing over all possible configurations of candidate entity labels from a given document. To circumvent this complexity, we used Collins' voted perceptron approach (Collins, 2002), which can be seen as approximating the full expectation of $\mathbf{f}_{\mathbf{c}}$ with the $\mathbf{f}_{\mathbf{c}}$ counts for the most likely labeling under the current parameters \mathbf{w} .

$$\nabla L(w,D) \approx \sum_{d \in D} f_c(d.X,d.Y) - \sum_{d \in D} f_c(d.X,d.Y_w)$$
(6)

The Voted Perceptron algorithm is detailed in Table 5. At each step i in the algorithm, inference is performed using the current parameters w_i , so that we get the most likely labeling $d.Y_i$. The parameters are then updated

based on the difference between the features counts computed on the ideal labeling d.Y and those computed on the current most likely labeling $d.Y_i$. The final set of parameters is the average taken over the parameters at all steps *i* in the algorithm.

Input: a set of documents <i>D</i> , number of epochs <i>T</i> ,
learning rate η .
set parameters $w_0 = 0$
set counter $i = 0$
for $t = 1T$
for every document $d \in D$
$d.Y_i = arg \max_{d.Y'} P_{w_i}(d.Y' d.X)$
$w_{i+1} = w_i + \eta * [f(d.X, d.Y) - f(d.X, d.Y_i)]$
i = i + 1
Output: $w = \frac{1}{T D } \sum_i w_i$

Table 5: The Voted Perceptron Algorithm.

In all our experiments, the perceptron was run for 50 epochs, with a learning rate set at 0.01.

3.8 Experimental Results

We have tested the RMN approach on two datasets that have been hand-tagged for human protein names. The first dataset is Yapex¹ which consists of 200 Medline abstracts. Of these, 147 have been randomly selected by posing a query containing the (Mesh) terms protein binding, interaction, and molecular to Medline, while the rest of 53 have been extracted randomly from the GENIA corpus (Collier, Park, Ogata, Tateisi, Nobata, T.Ohta, Sekimizu, Imai, Ibushi, & Tsujii, 1999). It contains a total of 3713 protein references. The second dataset is Aimed² which has been previously used for training the protein interaction extraction systems in (Bunescu, Ge, Kate, Marcotte, Mooney, Ramani, & Wong, 2004). It consists of 225 Medline abstracts, of which 200 are known to describe interactions between human proteins, while the other 25 do not refer to any interaction. There are 4084 protein references in this dataset. We compared the performance of three systems: LT-RMN is the RMN approach using local templates and the overlap template, GLT-RMN is the full RMN approach, using both local and global templates, and CRF, which uses a CRF for labeling token sequences. We used the CRF implementation from (McCallum, 2002) with the set of tags and features used by the Maximum-Entropy tagger described in (Bunescu et al., 2004). All Medline abstracts were tokenized and then POS tagged using Brill's tagger (Brill, 1995). Each extracted protein name in the test data was compared to the human-tagged data, with the positions taken into account. Two extractions are considered a match if they consist of the same character sequence in the same position in the text. Results are shown in Table 6 which give average precision (P), recall (R), and F-measure (F) using 10-fold cross validation.

$$P = \frac{\#\text{correct extractions}}{\#\text{extractions}}, \quad R = \frac{\#\text{correct extractions}}{\#\text{annotated extractions}}, \quad F = \frac{2P \times R}{P + R}.$$

These tables show that, in terms of F-measure, the use of global templates for modeling influences between possible entities from the same document significantly improves extraction performance over the

¹URL:www.sics.se/humle/projects/prothalt/

²URL: ftp.cs.utexas.edu/mooney/bio-data/

Yapex				Aimed			
Method	Precision	Recall	F-measure	Method	Precision	Recall	F-measure
LT-RMN	70.79	53.81	61.14	LT-RMN	81.33	72.79	76.82
GLT-RMN	69.71	65.76	67.68	GLT-RMN	82.79	80.04	81.39
CRF	72.45	58.64	64.81	CRF	85.37	75.90	80.36

Table 6: Extraction Performance on two protein datasets.

local approach (a one-tailed paired t-test for statistical significance results in a p value less than 0.01 on both datasets). There is also a small improvement over CRFs, with the results being statistically significant only for the Yapex dataset, corresponding to a p value of 0.02. We hypothesize that further improvements to the LT-RMN approach would push the GLT-RMN performance even higher. The tagging scheme used by CRFs, in which each token is assigned a tag, is essentially different from the RMN approach, where candidate extractions are either rejected or accepted. In the tagging approach used by CRFs, extracted entities are available only after tagging is complete, thereby making it difficult to account for influences between them during tagging.

Figures 13 and 14 shows the precision-recall curves for the two datasets. These were obtained by varying a threshold on the extraction confidence, which is the posterior probability that its label is 1 as computed by the sum-product algorithm.



Figure 13: Precision Recall Curves on Yapex.

We also explored using a global template that captured the tendency for candidate entities whose phrases are coordinated to have the same label. An example is shown in Figure 15 where the two entities 'eNOS' and 'NOSIP are coordinated through the conjunction 'and'. This template did not improve performance since detecting whether two NPs are coordinated is difficult, and the methods we tried introduced too many false coordinations.



Figure 14: Precision Recall Curves on Aimed.



Figure 15: Coordinated phrases eNOS and NOSIP have the same entity label.

In order to evaluate the applicability of our method to other types of narrative, we also tried it on the CoNLL 2003 English corpus (Tjong Kim Sang & De Meulder, 2003) which contains four types of named entities: persons (PER), locations (LOC), organizations (ORG), and other (MISC). Consequently the number of label values increased from two to five (with a label-value of 0 to indicate none of the four categories). For the global approach we used the same overlap template and a modified version of the repeat template in which the OR potential was replaced with a different type of potential (SEL) that allows at most one of the including entities to have a non-zero label-value. The SEL variable (replacing the OR variable) is forced to have label-value 0 if all including entities have label-value 0, otherwise it selects the one label-value that is not 0. The resulting repeat template, besides handling exact repetitions, is also able to capture correlations between entity types, when one entity repetition is included in another entity with a potentially different type. For example, it is common in this corpus to have country names repeated inside organization names in the same document, as is "Japan" in "Bank of Japan", or "Japan Aluminium Federation".

The overall results are shown in Table 7, with the global approach exhibiting improvement over the local approach, albeit less pronounced than in the biomedical domain. This results are still under some of the best published results on the same corpus - however no dictionaries were used in our experiments, and no custom feature selection was performed – the feature templates were the same as those used in the biomedical extraction.

Method	Precision	Recall	F-measure
LT-RMN	82.15	78.13	80.09
GLT-RMN	83.17	81.44	82.30
CRF	81.57	80.08	80.82

Table 7: Extraction Performance on CoNLL.

4 Current Research

The sum-product algorithm is guaranteed to do exact inference for factor graphs without cycles. However, it happens very often that the factor graphs generated by our approach contain cycles, even when only the overlap template is used (i.e. the LT-RMN model). For example, if $w_1w_2w_3$ is a sequence of three nouns, the entire sequence, together with all its subsequences, will become candidate entities. We shall use the letter *e* to denote these candidate entities, as follows:

- unigram entities: $e_1 \leftarrow w_1, e_2 \leftarrow w_2, e_3 \leftarrow w_3$
- bigram entities: $e_{12} \leftarrow w_1 w_2, e_{23} \leftarrow w_2 w_3$
- trigram entities: $e_{123} \leftarrow w_1 w_2 w_3$

The corresponding set of overlapping pairs is $\{(e_1, e_{12}), (e_1, e_{123}), (e_2, e_{12}), (e_2, e_{23}), (e_2, e_{123}), (e_3, e_{23}), (e_3, e_{123}), (e_1, e_{12}, e_{12}), (e_1, e_{123}), (e_{12}, e_{123}), (e_{23}, e_{123})\}$. The overlap template will create a two-node clique between the two nodes from each overlapping pair in the RMN factor graph, as illustrated in Figure 16. Of all cycles contained in this graph, we have emphasized using thick lines the cycle $e_1 - e_{12} - e_2 - e_{123} - e_1$, with the corresponding factor graph illustrated on the right.



Figure 16: RMN and factor graph with cycles due to overlap clique potentials

In the next section, we are going to show that exact inference can be done for the local model (LT-RMN) in linear time, based on the junction-tree algorithm (Cowell, Dawid, Lauritzen, & Spiegelhalter, 1999) and the sparsity of a different version of the overlap potential.

4.1 Local models

The original overlap template used in our approach creates an edge between any two overlapping entities. The constraint that entities should not overlap can also be enforced with a different type of overlap template, as follows. For a token position i in the document, let E_i be the set of candidate entities whose span includes that position. The overlap template is then defined so that, for each i, it connects the labels of all entities in E_i in a clique, with an associated potential that is non-zero only when at most one entity from E_i has label-value 1. Thus, if $|E_i| = n$, then the corresponding overlap potential can be specified as a table with 2^n entries, of which n + 1 are set on 1, the rest being set to 0. This makes the potential table very sparse (a linear number of non-zero entries), a fact that will be used later in the inference algorithm. Notice that this version of the overlap template can match a variable number of entities, depending on the number of overlap template can.

Continuing with the same noun phrase example $w_1w_2w_3$, the sets of overlapping entities corresponding to the three token positions are:

- 1. $E_1 = \{e_1, e_{12}, e_{123}\}$
- 2. $E_2 = \{e_2, e_{12}, e_{23}, e_{123}\}$

3. $E_3 = \{e_3, e_{23}, e_{123}\}$

The overlap template creates three cliques, corresponding to the three sets E_1 , E_2 , and E_3 . This results in the same graph as that from Figure 16, containing numerous cycles, which again means that the belief propagation algorithm (or the sum-product algorithm in the equivalent factor graph) is not guaranteed to result in exact inference.

4.1.1 Exact, linear time inference

The junction tree algorithm (Cowell et al., 1999) is a generalization of the sum-product algorithm that can be used for exact inference in general graphs. It is based on the junction tree representation, which is a singly connected graph whose nodes are clusters of nodes from the original graph. The usability of the junction tree algorithm is however limited by the fact that its time complexity is exponential in the size of the largest cluster, which can get very large, especially when the original graph has cycles.

Definition 5 (Cowell et al., 1999) H = (H.V, H.E) is a cluster graph for G = (G.V, G.E) if $H.V \subseteq 2^{G.V}$ e.g. any vertex in H is a cluster of vertices from G.

Definition 6 (*Cowell et al., 1999*) A cluster graph *H* is a **junction tree** for *G* if it has the following three properties:

- 1. singly connected: there is exactly one path between each pair of clusters.
- 2. covering: for each clique A of G there is some cluster C such that $A \subseteq C$.
- 3. **running intersection:** for each pair of clusters C and C' that contain a vertex $v \in G.V$, each cluster on the unique path between C and C' also contains v.

In order to create a cluster graph having the running intersection property, one needs to *triangulate* the original graph. *Triangulation* refers to adding sufficient additional edges such that the graph contains no *chordless* cycles i.e. cycles of four or more distinct vertices without a short-cut. The cluster nodes in the junction tree are simply maximal cliques in the triangulated graph. It is usually the process of triangulation which leads to arbitrarily large cliques in the triangulated graph, which translates into arbitrarily large cluster nodes in the junction tree. Fortunately, as the next theorem asserts, the graphs created by the overlap template are already triangulated (e.g. *chordal*):

Theorem 2 Let $w_1w_2...w_n$ be a word sequence, arbitrarily long (it may be the entire sequence of words from a document). Let E be an arbitrary set of candidate entities, and E_i the set of overlapping entities at position i, where $1 \le i \le n$. Let G be the graph created by the application of the overlap template e.g. the result of creating n cliques, one clique for each E_i , for all $1 \le i \le n$. Then the overlap graph G is a chordal graph.

For example, it can be verified easily that the overlap graph in Figure 16 is a chordal graph. We do not include the proof of this theorem here, as it will not be used directly in creating the junction tree associated with an overlap graph. Instead, we are going to create a particular cluster graph and show that it is a junction tree for the overlap graph by verifying directly the three properties from Definition 6, as in the following theorem:

Theorem 3 Keeping with the notation from Theorem 2, let H be a cluster graph for G, defined as follows:

- $H.V = \{E_i | 1 \le i \le n\}$ e.g. the sets of overlapping entities are vertices in the cluster graph.
- $H.E = \{(E_i, E_{i+1}) | 1 \le i \le n-1\}$ e.g. connect clusters corresponding to consecutive positions only (resulting in a list of clusters).

Then H *is a junction tree for* G.

The result of applying this procedure on the overlap graph in Figure 16 is illustrated in Figure 17. Ellipses denote *cluster nodes*, while rectangles (*separator nodes*) are used to show the intersection between adjacent cluster nodes. It can be easily verified that this is a junction tree.



Figure 17: Sample junction tree.

Proof of Theorem 3. The first two properties from Definition 6 are obviously verified. What is left for proof is the running intersection property. Let E_i and E_j be two cluster nodes. Assume without loss of generality that i < j. Let e be a vertex in the original graph G such that $e \in E_i$ and $e \in E_j$. Let e.l and e.r be the left and respectively right boundaries of entity e in the word sequence. Using the definition of the overlapping sets, $e \in E_i \Leftrightarrow e.l \leq i \leq e.r$, and $e \in E_j \Leftrightarrow e.l \leq j \leq e.r$. Let E_k be a cluster node on the path between E_i and E_j . Because of the way in which H was created, k should be a position between i and j i.e. i < k < j. At this point, we have these three inequalities:

- $e.l \leq i \leq e.r$
- $e.l \leq j \leq e.r$
- i < k < j

Based on these inequalities, we get that $e.l \le i < k < j \le e.r$, therefore e.l < k < e.r. This implies that $e \in E_k$. As E_k was an arbitrary cluster on the path between E_i and E_j , this means that H has the running intersection property. Therefore, H is a junction tree for G.

Both Theorem 2 and Theorem 3 are important because they show that the size of the largest cluster in the junction tree (i.e. its *width*) is actually the size of the largest overlapping clique in the original graph. Because the inference algorithm using junction trees is in general exponential in this size, and because the size of the largest overlapping clique can be linear in the number of candidate entities, this means that exact inference using the generic junction tree algorithm is still exponential in the number of candidate entities. However, as Theorem 3 shows, for any overlapping graph, there exists a junction tree whose clusters are exactly the overlapping cliques. Because of the special form of the overlap clique potential (a sparse table, with only n + 1 non-zero entries, where n is the size of the clique), the messages sent between two adjacent *cluster nodes* in the junction tree can be computed in time linear in the size of the cluster. We therefore have an exact inference algorithm based on message propagation, where:

- The computation of any message takes time linear in the size of the adjacent cluster nodes.
- Assuming a two-phase propagation schedule (Jensen, Lauritzen, & Olesen, 1990), the total number of messages is twice the number of cluster nodes.
- Assuming the length of any candidate entity is less than a maximum length (as is the norm in information extraction), the sum of all cluster sizes in the junction tree is linear in the total number of candidate entities.

Based on the three facts above, the overall time complexity of the message propagation algorithm in the junction tree structure from Theorem 3 is linear in the number of candidate entities.

4.1.2 Learning Algorithm

Because the overlap template potential is fixed, the only potential values that need to be learned are those used by local templates. Based on the same notation as in Section 3.7, we use the log-linear representation for a local template potential $\phi_c = exp(\mathbf{w_c f_c})$. Being an exponential model, the gradient of the log-likelihood objective function $L(\mathbf{w}, D)$ with respect to the weight vector $\mathbf{w_c}$ is the difference between the observed and expected counts of the feature vector $\mathbf{f_c}$:

$$\nabla_c L(w,D) = \frac{\partial L(w,D)}{\partial w_c} = \sum_{d \in D} f_c(d,X,d,Y) - \sum_{d \in D} \sum_{d,Y'} f_c(d,X,d,Y') P_w(d,Y'|d,X)$$
(7)

$$= \sum_{d \in D} f_c(d.X, d.Y) - \sum_{d \in D} E_w[f_c(d)]$$
(8)

While the first term in Equation 8 is easy to compute, the second term is usually expensive to compute in general graphical models. In the current setting however, we'll make use of the fact that all potentials involved are local potentials. Therefore, we can write:

$$f_c(d.X, d.Y) = \sum_{e \in d.E} f_c(e.X, e.Y)$$
(9)

Consequently, the expectation term in Equation 8 can be rewritten as follows:

$$E_w[f_c(d)] = \sum_{d,Y} \sum_{e \in d,E} f_c(e,X,e,Y) P_w(d,Y|d,X)$$
(10)

$$= \sum_{e \in d.E} \sum_{d.Y} P_w(d.Y|d.X) f_c(e.X, e.Y)$$
(11)

$$= \sum_{e \in d.E} \sum_{e.Y} \left(\sum_{d.Y \sim e.Y} P_w(d.Y|d.X) \right) f_c(e.X, e.Y)$$
(12)

$$= \sum_{e \in d.E} \sum_{e.Y} P_w(e.Y|d.X) f_c(e.X, e.Y)$$
(13)

The expression $d.Y \sim e.Y$ above refers to all labelings of d consistent with a particular entity label e.Y. The term $P_w(e.Y|d.X)$ in the last equation is the marginal distribution for an entity label e.Y, which can be easily computed after running belief propagation in the junction tree, by selecting a cluster node containing e and marginalizing the cluster distribution over all other entities from the same cluster. Because the junction tree algorithm computes all clusters' marginal distributions at once, this means that computing the expectation term $E_w[f_c(d)]$ takes time linear in the number of candidate entities d.E. Based on the last equation, the final formula for the gradient is:

$$\nabla_{c}L(w,D) = \sum_{d \in D} \sum_{e \in d.E} \left(f_{c}(e.X,e.Y) - \sum_{e.Y'} P_{w}(e.Y'|d.X) f_{c}(e.X,e.Y') \right)$$
(14)

with a total computation time linear in the number of candidate entities.

Based on this formulation, any gradient-based method can be used for maximizing the likelihood function. In our implementation we used L-BFGS, a limited-memory quasi-Newton method (Liu & Nocedal, 1989), which has shown very good performance elsewhere (Sha & Pereira, 2003).

In conclusion, we have introduced a discriminative model for information extraction based on phrase classification, in which exact inference is linear in the number of candidate phrases, and where both ML and MAP learning can be done efficiently. The overall approach is simple, and can be summarized as follows:

1. Candidate Entities: Based on the generic heuristic H1, or alternative domain-specific heuristics, create a set of candidate entities d.E, for each document in the corpus, $d \in D$.

- 2. Junction Tree: Assemble the set of candidate entities into cluster nodes E_i , one node for each token position *i* in the document. Each cluster E_i contains candidate entities that span over position *i*. Link cluster nodes corresponding to consecutive positions. The result is a list of cluster nodes, which by Theorem 3 is a junction tree for the original overlap graph. Depending on the set of candidate entities, some positions in the document may result in empty clusters, which split the junction tree into two or more smaller trees. This is also the case when the document is split in sentences first because no entity can belong to two different sentences, each sentence will have its own separate junction tree.
- 3. Cluster Potentials: Initialize each cluster potential with an overlap potential. Due to sparsity, if the cluster size is n, the cluster potential can be represented using only n + 1 numbers. Multiply all local template potentials for each entity into one and only one cluster potential. If there is more than one cluster containing the entity, choose one at random.
- 4. **Inference:** Run a message propagation algorithm on the resulting set of junction trees, using a twophase propagation schedule.
- 5. Learning: Use a gradient based method in a ML or MAP setting, based on the gradient formula in Equation 14.

Compared with CRFs, this has the additional benefit of allowing the incorporation of phrase-based features. In a recent work, (Cohen & Sarawagi, 2004) have introduced a conditional version for segmental semi-Markov models (Ge, 2002) to achieve a similar aim, showing that the phrase classification approach can lead to better performance vs. CRFs, especially when training data is small, because of a more natural use of phrase based features, such as similarities with existing dictionaries. Compared with their work, where sentences are modeled as Markov sequences of segments, our approach is more direct in modeling the extraction task as one of phrase classification. We explicitly model the entire set of candidate entities by including a node for each entity label in the graphical model. This has the advantage of allowing the incorporation of *global* correlations between labels of different entities from the same document, as will be detailed in Section 5.1.

4.1.3 Experimental Results

In Tables 8 and 9 we compare the performance of the (approximate) inference algorithm in factor graphs (FG) with that of the (exact) inference algorithm in junction trees (JT) on the two protein data sets. In both cases, exact inference leads to better results.

Method	Precision	Recall	F-measure
LT-RMN (FG)	70.79	53.81	61.14
LT-RMN (JT)	72.08	57.46	63.95

Table 8: Extraction Performance on Y

Method	Precision	Recall	F-measure
LT-RMN (FG)	81.33	72.79	76.82
LT-RMN (JT)	81.76	75.11	78.29

Table 9: Extraction Performance on Aimed.

5 Proposed Work

5.1 Restricted global models

The global phrase model is an extension of the local phrase model. While in the local model the overlap clique template is the only template introducing cliques between the labels of different entities, in the global model any type of global clique template can be used. Unfortunately, adding new cluster nodes in the junction tree representation so that it has the *covering* property with respect to the global phrase model may easily break the *singly connected* property. Sometimes adding just one cluster node results in a cycle, as illustrated in Figure 18. The new cluster node E may have been introduced, for instance, in order to model the repetition of two candidate entities $u \in E_i$ and $v \in E_j$.



Figure 18: Introducing a cycle in the junction tree.

However, if the set of candidate entities is the result of applying a strong heuristic, like H2 or H3 from Section 3.1, then some of the positions in the text will result in *empty clusters*, which have the benefit of breaking cycles, as illustrated in Figure 19. There, the cluster for position k is empty. As defined in Theorem 3, a cluster node E_k contains all candidate entities overlapping at position k in the text. Then E_k is an *empty cluster* if no candidate entities overlap at position k. If H2 is used, this may be because the word at position k was tagged as a preposition, verb, or other part of speech different from those used by H2.



Figure 19: Empty clusters break cycles.

This means that, depending on the heuristic used to create the set of candidate entities, at least a subset

of the global label correlations can be introduced in the model such that the inference remains linear in the number of entities. Both H2 and H3 give rise to many empty clusters. In general, both heuristics result in models composed of many small junction trees, which can therefore be connected into larger junction trees using global label correlations.

5.1.1 Exact, linear time inference

Even though they allow updating the model with global dependencies such that it remains tractable, heuristics H2 and H3 have the drawback that sometimes they may miss real entity names. In the case of H2 for example, a candidate entity cannot contain parentheses, however our corpus contains a few entity names like 'V (1a) receptor', or 'interleukin 10 (IL-10) receptor', which violate this assumption. Because the local phrase model may be able to learn patterns like "allow a close parenthesis in an entity name if it is followed by the word *receptor*", we propose to eliminate domain-specific heuristics like H2 and H3 and to assign their role to the local phrase model. The extraction process will then proceed in two steps:

- 1. Local Extraction. Using only the local phrase model and the generic heuristic H1, perform exact inference based on the junction-tree representation. At each position k in the text, the probability that the corresponding cluster is "empty" e.g. $P(e.label = 0, \forall e \in E_k)$ is readily available as one of the marginals computed during inference. Eliminate from the junction-tree all cluster nodes for which this probability is above a predefined threshold $\tau \in [0, 1]$. This will result in a forest of junction-trees that will be the input for the next step. From the remaining clusters remove all candidate entities that were contained in any of the eliminated clusters.
- Global Extraction. Connect separate junction-trees through cluster nodes corresponding to global correlations, as illustrated in Figure 19. Stop when no global correlation can be added without introducing a cycle in the model. Perform exact inference in the resulting junction-tree, this time in order to recover the most probable assignment of labels to candidate extractions, and output this as the final extraction.

Thus, instead of coming up with a domain-specific heuristic to restrict the set of candidate extractions, we can use the local phrase model to automatically learn the heuristic. The threshold τ could be set up based on development data. Lower values for τ will break the junction-tree into many smaller trees, which will allow many global correlations to be used. However, eliminating too many cluster nodes increases the risk of eliminating true entities. On the other hand, a value of τ too close to 1 will eliminate only a few clusters from the junction-tree, and correspondingly very few global correlations can be used. Another question is that of which subset of global correlations to be used in Step 2. For the repeat template, an obvious choice, better than randomly choosing global correlations, is to include in the selected pairs of repetitions as many "locally extracted" entities as possible, giving priority to repetitions for which the local phrase model assigns conflicting labels (by "locally extracted" entity we mean an entity that would be extracted by the local phrase model).

5.1.2 Learning Algorithm

Learning the parameters for the global phrase model mirrors the two steps used in inference:

1. Local Model. Learn first the parameters for the local model, exactly as described in Section 4.1.2.

2. Global Model. Keeping the local parameters fixed, learn the global parameters e.g. the potentials for the global templates. The training data is first made *consistent* with the set of candidate entities that resulted from the application of the local model with threshold τ . This means eliminating from training data all entities not contained in the set of candidate entities. This step will ensure that the likelihood of the global parameters with respect to the training data is not identically zero. Then global parameters are estimated so that they maximize the log-likelihood using a procedure similar to that used for the local parameters.

A definite short term goal is to implement the restricted global model, and to compare its performance with that of the factor graph implementation of the unrestricted global model from Section 3.5. We expect that the potential loss in accuracy caused by the model ignoring some of the global correlations will be compensated by the increase in accuracy due to exact inference. There are also numerous approximate inference methods that exploit the structure of the graphical model and the associated potentials in order to obtain results closer to the exact version, so that the inference step is still tractable. In (Yedidia et al., 2000), the authors introduce generalized versions of the belief propagation algorithm, where messages are passed between sets of nodes, at additional computational cost. Depending on the choice of the basic sets of nodes, the resulting inference algorithm is shown to lead to significantly better approximations versus the original belief propagation algorithm. We intend to experiment with this and other alternative approximate inference methods, by suitably adapting them to our approach to information extraction.

5.2 Collective classification for WSD and POS tagging

Information Extraction is only one of the natural language tasks that can benefit from document level correlations - two other obvious examples are word sense disambiguation (WSD) and part-of-speech (POS) tagging. In WSD, given a text document, the task is to annotate all content words from the document with the appropriate sense, selected from a sense inventory such as WordNet. POS tagging can be seen as a coarse version of WSD, where the sense inventory is limited to a set of syntactic categories like noun, adjective, verb, adverb, pronoun, preposition, conjunction, etc. (for an exhaustive list, see (Santorini, 1990)). The one-sense-per-discourse hypothesis (Yarowsky, 1995) specifies that multiple occurrences of the same word in one document tend to have the same sense. This can be easily modeled in an RMN model by connecting any pair of repeated words through an undirected edge. For the task of POS tagging, current state-of-the-art results are obtained with algorithms like CRFs (Lafferty et al., 2001) that assume a Markov dependency between consecutive tags. This means that by adding an edge between the tags of two repeated words, the resulting graphical model may contain cycles, which are hard to accommodate in the inference algorithm. Besides resorting to approximate inference, we can also use linear time exact inference in a restricted version of the global model, as described in Section 5.1. This would lead to the graphical model depicted in Figure 20, where black nodes correspond to high confidence tags, and arcs connect tag nodes for pairs of repeated words.



Figure 20: High confidence tags break cycles.

Given the already competitive performance of POS tagging algorithms, the advantage of using document

level correlations between tags in POS tagging should be most evident when the learned model is applied on documents whose narrative is different from that in the training corpora. One example would be training the POS tagger on the Penn Treebank Corpus (Marcus, Santorini, & Marcinkiewicz, 1993) and testing it on gold-standard POS tagged Medline abstracts from the GENIA corpus (Collier et al., 1999).

5.3 Using the web to improve information extraction

The web can provide additional evidence regarding the class of a candidate entity. If the task is that of recognizing protein names, and x is a candidate entity, then one may argue that the phrase pattern "x is a protein" is likely to have been used in a document on the web, if x is indeed a protein name. Consequently, a search engine should be able to return that document when searching for the phrase "x is a protein". Another reasonable hypothesis is that the more hits a search engine gets for this pattern, the more likely it is that "x" is a protein name.

There is however a problem with the pattern above – the predicate nominal "protein" is often too general, and consequently it is not used to introduce a protein instance so often as one would expect. Instead, people tend to introduce a protein name x using a predicate nominal describing a subclass or a family of proteins, as in "x is an enzyme", or "x is a eukaryotic transcription regulatory factor". This is in agreement with the well known principles of categorization introduced by Rosch in (Rosch, 1978), which postulate a basic-level of categories that provide maximum information with minimum cognitive effort. Moreover, even when the generic term "protein" is used, it is often preceded by a modifier, as in "x is a giant protein", or by a sequence of modifiers, as in "x is a general eukaryotic protein". To account for an arbitrary sequence of modifiers preceding the name of a protein family, one would need a search engine capable to answer queries containing wild cards, which is beyond current technology. Ideally, we would need a search engine able to answer queries of the form "x is a np $\langle y \rangle$ ", where $np\langle y \rangle$ denotes any noun phrase headed by y. Such tools have only begun to appear, and their coverage is still very limited. One notable example is the Linguist's Search Engine (Resnik & Elkiss, 2003), which currently indexes a corpus of three million sentences from the Internet Archive.

To accommodate the query types supported by current search engines, we shall limit ourselves to using phrase patterns of the type "x is a p", where p is a particular class of proteins, instantiated from an already available ontology of proteins, such as that from the Gene Ontology Database (Gene Ontology Consortium, 2000).

The number of hits returned by the search engine for a particular pattern can be used to compute various quantities such as the *pointwise mutual information (pmi)*, which has been previously used for computing word association norms (Church & Hanks, 1990). For the pattern "x is a y", this amounts to computing:

$$pmi(x, y) = \frac{Hits('x \text{ is } a y')}{Hits('x') * Hits('y')}$$

If x_1 and x_2 are two candidate entities, and y is the generic name "protein", then the fact that $pmi(x_1, y) > pmi(x_2, y)$ could be used to assert that x_1 is more likely to be a protein name than x_2 . The same pmi measure has been used in conjunction with web or large corpora searches for measuring the similarity of pairs of words (Turney, 2001), for computing the semantic orientation of reviews (Turney, 2002), or for solving associative anaphora (Bunescu, 2003).

This, or other similar measures, could be integrated in the IE system as additional features. However, given the big number of candidate entities considered by the IE system when doing extraction, it would be highly inefficient to perform a web search for each of them. This leads to a scenario in which, first, the

normal IE system is applied on a document, after which the confidence for each extraction is updated based on the statistics collected by the search engine on the corresponding discriminative patterns.

5.4 Flexible use of external dictionaries

Named entity recognition can also benefit from the use of external dictionaries. In our previous work (Bunescu et al., 2004), we used a dictionary of more than 70,000 protein names to improve the performance of a maximum entropy tagger. One way of using the dictionary in a token classification approach is to find all dictionary entries that occur in the document and pre-tag the tokens of those occurrences with a special tag. Features can then be defined so that they take into account this tag. The main drawback of this approach is that the document may contain phrases which do not exist in the dictionary, yet they are very similar with entries in the dictionary. We would like to use the intuition that phrases which are similar with dictionary entries are likely to be valid named entities. For example, assume that the task is to extract protein names, and the current document contains the phrase 'USF-2', which is missing from the dictionary. Instead, the dictionary contains the similar entry 'USF-1'. This indicates that, at least from an orthographic point of view, 'USF-2' could be a protein name. In (Bunescu et al., 2004) we captured this by generalizing the dictionary entries - numbers, single Roman letters, and Greek letters were replaced with a corresponding generic tag, that would match any number, single Roman letter or Greek letter respectively. Following the example above, the entry 'USF-1' would be generalized to 'USF- $\langle n \rangle$ ', where the tag $\langle n \rangle$ can match any number. As a result, this generalized entry would eventually match the phrase 'USF-2' in the document. While this approach worked well for protein tagging, it has two main drawbacks:

- Incompleteness: The document may contain phrases which are similar with dictionary entries, yet they are not matched by any of the generalized dictionary entries.
- Specificity: The generalizations used are very domain specific. They may not be relevant for other domains, or, even if they can be used in another domain, they may be just a subset of the relevant generalizations.

Ideally, we would like to use a measure of the similarity of each candidate entity with entities in the dictionary, a measure that would be learned for each domain. This similarity measure cannot be used in a token classification approach, which lacks the concept of candidate entities, nevertheless it is very straightforward to integrate it in a phrase classification approach. There, the similarity of a candidate entity with entries in an external dictionary would be just another local feature for that entity. A similar approach has been used in (Cohen & Sarawagi, 2004), where the authors defined the similarity between a text segment and a dictionary to be the maximum similarity between that segment and entries in the dictionary. As for the actual similarity metric to use in our model, we conducted a preliminary experiment in which we used a cosine similarity metric. This did not lead to any improvement in accuracy, mainly because common tokens which occur in protein names, such as numbers or Greek letters, were getting too high a similarity with the dictionary. Yet these tokens cannot be protein names by themselves. This suggests an approach where the similarity metric is learned, so that tokens like these get a low similarity with the dictionary. We contemplate using adaptive similarity measures, based on the ideas introduced in (Bilenko & Mooney, 2003). One issue that needs to be addressed is the lack of "well defined" training examples. In (Bilenko & Mooney, 2003), the authors show how similarity metrics can be learned, assuming a training set composed of pairs of strings which are known to be similar (positive examples) or dissimilar (negative examples). This kind of training examples are missing in our approach, where all we are given is a dictionary of entity names, together with a set of candidate entities partitioned in two subsets: true entities and spurious entities. Nevertheless, we can assemble a weakly labeled dataset as follows:

- Negatives: Consider as negative examples all pairs consisting of a spurious candidate entity and a dictionary entry.
- Positives: Consider as positive examples all pairs consisting of a true candidate entity and a dictionary entry, such that they share a minimum number of tokens. This list of positive examples may be later replaced by, or augmented with, manually labeled pairs of similar entities.

For the special case of recognizing protein names, we can also use the fact that the protein dictionary is organized in clusters of names, where all names from a cluster are synonyms corresponding to the same protein. Therefore, we may consider as positive examples all pairs of synonyms from the same cluster which share a minimum number of tokens.

5.5 Feature selection

The task of feature selection refers to finding a subset of features, out of a usually large collection of features, such that they capture the relevant properties of the data. In a supervised learning setting, this reduces to choosing that set of features which best model the labels in the training data. In this case, the final aim of feature selection is to construct classifiers which are both compact and accurate. In Maximum Entropy models, features selection has been commonly done using a simple frequency-based cut-off. This is also the method that we used in our initial RMN model for information extraction (Bunescu & Mooney, 2004). There, we have ignored all features which do not occur at least three times in the training data. Recently, we have run an experiment in which we used only a subset of the features proposed in (Bunescu & Mooney, 2004) (also repeated in Section 3.2 of this proposal). More exactly, we used the following feature templates: the text of the candidate entity, its short type, the word and the POS tag preceding/following the entity, the first and last words in the candidate entity, and any word occurring inside the entity. This time we considered all features generated by the above templates in the training data, irrespective of their actual frequency. The results on the Yapex dataset are summarized in Table 10.

Method	Precision	Recall	F-measure
LT-RMN	73.90	54.90	63.00
GLT-RMN	74.16	63.48	68.40
CRF	72.45	58.64	64.81

Table 10: Extraction Performance on Yapex.

The results for both the local and global models are slightly better than those presented in Table 6 for the same dataset. This suggests that, besides making the model more compact, an appropriate feature selection algorithm may also lead to a non-negligible increase in performance. Likelihood-based feature induction algorithms, as introduced in (Della Pietra et al., 1997) and further extended to conditional random fields in (McCallum, 2003) can be seen as a particular way of doing feature selection. We intend to use feature induction in our setting, which means that we need to start with a set of atomic, local features, such as the word preceding an entity, the entity head, and others. New features, such as bigrams or trigrams, can be created from conjunction of atomic features. As explained in (McCallum, 2003), the set of features is incrementally updated by iterating the following four steps (initially, the set of features is empty):

1. Consider a set of newly proposed features. This may contain both atomic features or conjunctions of features.

- 2. Select for inclusion only those features which bring the highest gain in the likelihood function.
- 3. Train the weights for all included features.
- 4. Iterate back to the first step, until a stopping criterion is met.

5.6 Relation extraction and syntactic information

Until now we have focused on the task of named entity recognition. A natural next step is that of identifying relations between extracted entities. The types of relations can be very diverse, ranging from personaffiliation and organization-location, occurring frequently in newspaper corpora, to protein-protein interactions or subcellular-localization, which are common in biomedical corpora. Current best results for relation extraction are obtained with methods that make use of syntactic information. In (Ray & Craven, 2001), the authors show how information from the shallow parses of sentences can be represented into the states of a Hidden Markov Model, leading to increased extraction performance for two binary relations in the biomedical domain. This is however a generative model, and consequently subsequent approaches tried to exploit the additional strengths of discriminative models. In (Zelenko, Aone, & Richardella, 2003), the authors define convolution kernels on shallow parse trees where nodes are augmented with relation roles. Deeper syntactic information is used in (Culotta & Sorensen, 2004), where relation roles are used to annotate nodes in a dependency tree representation of sentences, with experimental results showing significant error reduction over the corresponding bag-of-words approach. Based on these ideas, we intend to create a discriminative model for the task of extracting protein-protein interactions from Medline abstracts – the challenge here is in coming up with a model that is robust enough to parsing errors. These errors are more numerous in the case of biomedical corpora, given that most state-of-the-art parsers are trained on newspaper corpora.

The two discriminative relation extractors referenced above assume that the entities used for instantiating relation roles have already been tagged. However, the same syntactic information that is used for extracting relations can also help in entity recognition. In Table 11 we illustrate the 'accuse' frame as instantiated in a set of sentences from the CoNLL 2003 corpus.

- 1) (Syria) [accused] (Israel) on Wednesday of launching a hysterical campaign ...
- 2) (Sharif) [accused] (Bhutto) of corruption and nepotism ...
- 3) (Ernesto Samper) [accused] (the government) of indifference ...
- 4) (Iran) [has accused] (Iraq) of violating the ceasefire ...
- 5) (Jordan) [has accused] (Iraq) and a local pro-Baghdad party for the country's ...
- 6) (China) [on Thursday *accused*] (Taipei) of spoiling the atmosphere ...
- 7) (India) [has often *accused*] (Pakistan) of abetting militancy in the valley ...
- 8) (Loyola de Palacio) [had earlier *accused*] (Fischler) at an EU farm ministers' meeting ...
- 9) (Judges on the island) [had accused] (Paris) of taking a lax stance ...
- 10) (Bob Dole) [Wednesday accused] (the Clinton administration) of ignoring ...
- 11) (Benjamin Netanyahu) [has accused] (opposition leader Peres) ...
- 12) (An Iraqi Kurdish guerrilla group) [on Saturday accused] (Iraqi government forces) of ...
- 13) (The ruling Socialist Party) [last week accused] (Serbia's opposition) of ...
- 14) (Freddy Pinas, a Surinamese-born visitor from the Netherlands,) [accused] (Brunswijk) ...
- 15) (Hasina, speaking to a group of engineers in Dhaka on Monday,) [accused] (the BNP) of ...

Table 11: 'accuse' frame instantiations in CoNLL.

The head of the 'accuse' verb phrase is figured in italics, while the heads of the argument phrases are figured in bold. Notice that in the 'accuse' frame, both arguments are either persons, organizations, country names or capital cities. If a syntactic parse is available, this constraint can be easily modeled by a binary feature which is set to one if the candidate entity is the head of an argument of 'accuse'. The corresponding weight is then given a large value only when the candidate entity is labeled either as a person, organization, country name or capital city. Without using a parse, this constraint is enforced much more weakly, because of the presence of long range dependencies – examples more to the bottom of the table have more words occurring between the argument heads and the verb 'accuse'. A binary feature which is set to one if the candidate entity is followed by the verb 'accuse' would be triggered correctly only for the first three sentences. The same feature would be triggered also for spurious candidate entities such as 'Thursday', 'Wednesday', 'Saturday', 'week' (sentences 6, 10, 12, and 13 respectively). Introducing a feature for the second word following the candidate entity leads to the same problem.

Based on the observations above, we intend to create a robust discriminative model that would use information derived from syntactic parses for both entity and relation extraction.

6 Conclusion

We presented a phrase–based approach to collective information extraction in which correlations between the labels of candidate entities in a document were modeled by a relational Markov network. Preliminary experiments on extracting named entities from biomedical and newspaper corpora demonstrate the advantages of our approach. We then argued for an alternative representation of document–level correlations based on junction–trees that would allow for efficient, exact inference. This has the potential of further increasing the extraction accuracy. In future work we intend to use the same approach for other natural language tasks that may benefit from the same type of correlations, such as part-of-speech tagging and word sense disambiguation. We also contemplate integrating in our system features originating from additional knowledge sources, such as external dictionaries, web statistics on discriminative patterns, or syntactic parsing. The huge number of potential features motivates the use of a feature selection algorithm. A natural next step is to augment the current system such that it performs both entity and relation extraction.

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