Feature Subset Selection in Conditional Random Fields for Named Entity Recognition

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Abstract
In the application of Conditional Random Fields (CRF), a huge number of features is typically taken into account. These models can deal with inter-dependent and correlated data with an enormous complexity. The application of feature subset selection is important to improve performance, speed and explainability. We present and compare filtering methods using information gain or $\chi^2$ as well as an iterative approach for pruning features with low weights. The evaluation shows that with only 3% of the original number of features a 60% inference speed-up is possible. The $F_1$ measure decreases only slightly.

1 Introduction
Feature selection is well established for many machine learning methods, for instance for feed-forward neural networks [2] or decision trees [17]. The main advantages are an improvement of prediction performance, faster training and prediction as well as a better understanding of the models [4]. Methods can be distinguished between filters not using the learning algorithm and wrappers using the learning algorithm as a black box [8]. An overview of approaches for classification tasks is given by Liu and Motoda [11], more specifically for text classification by Yang and Pederson [25].

Such feature selection methods are not well established for Conditional Random Fields [9], a Maximum Entropy-based method [1] for structured data. We propose methods coping with the demanding task of handling sequential data represented by a huge number of features. Reported numbers are for instance 1,686,456 for a gene name tagger [5]. Due to this high complexity, training and inference times can explode. These high numbers of features are generated by automated methods, i.e., for every token in the training set, features are generated. Then, all other tokens are tested for these features. This method is typically applied to determine the identity of words as well as for prefixes or suffixes of different length or for learning schemata of regular expression-like patterns [20].

Only a few approaches dealing with feature handling for CRFs are published. The work by McCallum [12] demonstrates a method for iteratively constructing feature conjunctions that would increase conditional log-likelihood if added to the model. An analysis of different penalty terms for regularization is shown by Peng and McCallum [15]. Goodman [3] presented a related analysis of exponential priors for Maximum Entropy models. The very recent work of Vail et al. [23, 22] shows feature selection in Conditional Random Fields by $L_1$-norm regularization in the robotics domain, a work which is related to the selection in Maximum Entropy models proposed by Koh et al. [7].

These methods incorporate the training procedure in the selection process. In contrast, we present different filter methods for feature selection to limit the complexity before starting the training. It is demonstrated how the sequential structure of text can be respected by filtering approaches originally developed for classification problems (especially in Section 3.1.2 and 3.1.3). These filter methods are compared to an iterative approach.

The paper is organized as follows. A short description of Conditional Random Fields is given in Section 2. We introduce different approaches for feature selection in Section 3, namely the adaption of classification methods to filter features and an iterative approach to remove features with low weights. In Section 4, an evaluation of the feature selection methods is given. The results show a reduction of complexity leading to improved speed and better explainability.

2 Conditional Random Fields and Sequential Data
Conditional Random Fields [9, 13] are a family of probabilistic, undirected graphical models for computing the probability $P_S(y|x)$ of a possible label sequence $\tilde{y} = (y_0, \ldots, y_n)$ given the input sequence $\tilde{x} = (x_0, \ldots, x_n)$. In the context of Named Entity Recognition, this observation sequence $\tilde{x}$ corresponds to the tokenized text.

The label sequence is encoded in a label alphabet $L = \{I<-entity>, O, B<-entity>\}$ where $y_i = O$ means that $x_i$ is outside an entity, $y_i = B<-entity>$ means that $x_i$ is the beginning and $y_i = I<-entity>$ means that $x_i$ is inside an entity.

In general, a CRF is given by

$$P(\tilde{y}|\tilde{x}) = \frac{1}{Z(\tilde{x})} \prod_{j=1}^{n} \Psi_j(\tilde{x}, \tilde{y}) \quad (1)$$
with normalization $Z(\vec{x}) = \sum_{\vec{y} \in Y} \prod_{j=1}^n \Psi_j(\vec{x}, \vec{y})$, where $\Psi_j$ are the different factors and $Y$ is the set of all possible label sequences. These factor functions combine different features $f_i$ of the considered part of the text and label sequence and usually correspond to maximal cliques on the independence graph.

For simplicity, we focus on the linear-chain CRF as a special case of the general CRF. The factors are given in the form

$$\Psi_j(\vec{x}, \vec{y}) = \exp \left( \sum_{i=1}^m \lambda_i f_i \left( y_{j-1}, y_j, \vec{x}, \vec{y} \right) \right). \tag{2}$$

Each feature $f_i(\cdot)$ is weighted by $\lambda_i \in \mathbb{R}$. These weights are the parameters to be learned in the model and later used in the iterative approach for feature subset selection (in Section 3.2). An example formulation of features $f_i(\cdot)$ is

$$f_i \left( y_{j-1}, y_j, \vec{x}, \vec{y} \right) = \begin{cases} 1, & \text{if } y_{j-1} = s'_i \text{ and } y_j = s''_i \text{ and } \varphi_k(x_j) \text{ and } \varphi_k(x_{j+1}) \text{ otherwise,} \\ 0, & \text{otherwise}, \end{cases} \tag{3}$$

where $s'_i, s''_i \in \mathcal{L}$.

There are two kinds of features. First, the ones representing the token sequence as a feature vector sequence\(^1\) (these features are referred to as $\varphi_k \in \mathcal{F}$). An example is that $\varphi_k(x_j)$ holds if and only if $x_j$ is a capital word. Second, the feature functions representing $\varphi_k(x_j)$ with label transitions, to be referred to as $f_j \in \mathcal{F}$.

Optimization of the parameters $\lambda_i$ is often performed with the limited-memory Broyden-Fletcher-Goldfarb-Shanno algorithm (L-BFGS, Nocedal, [14]) on the convex function $L(T)$ with the training data $T$, including a penalty term:

$$L(T) = \log P_{\lambda}(\vec{y} | \vec{x}) - \sum_{i=1}^m \lambda_i^2 / 2 \sigma_i^2. \tag{4}$$

### 3 Feature Selection Methods for CRFs

An exhaustive search to find the optimal feature subset is not possible due to the large number of features and comparatively long training times. Hence, a wrapper approach considering the CRF as a black box is impractical. We compare different approaches for feature subset selection of sequential data in CRFs, i.e., filtering methods (in Section 3.1) and an iterative method (in Section 3.2).

#### 3.1 Filter

To apply filter methods for classification tasks, the sequence data have to be represented as classification instances. For a pair of sequences $(\vec{y}, \vec{x})$ this is done with respect to the incorporated factors in the CRF. For every factor $\Psi_j(\vec{y}, \vec{x})$, an instance is built. The labels are all dependencies on $\vec{y}$, the features have the values at the corresponding position for $\vec{x}$.

\(^1\)For simplicity, we only consider boolean features.

For the factors in a linear-chain CRF as shown in Equation 2, the instance $T_i = (\vec{y}_i, \vec{x}_i)$ at position $j (0 < j \leq n)$ has the label $\Sigma_j = (y_{j-1}, y_j)$ from a set of all possible transitions $\Sigma_j \in \mathcal{L}^2$. The feature values are $\varphi_k(x_j)$. Instances are built for all positions in all training examples from $T$.

The features are ranked by measures presented below. The best $p_{\text{frac}}$ features (where $p_{\text{frac}}$ is a parameter specifying the percentage of kept features) are selected to represent the text data.

In the following, the number of generated instances is denoted with $h$, the number of instances with feature $\varphi(x_j)$ with value 1 with $h_1$ and with value 0 with $h_0$. The number of instances with label $\Sigma_j$ is $h(\ell)$, with feature values 1 or 0 of those with $h_j^1(\ell)$ and $h_j^0(\ell)$ respectively.

### 3.1.1 Simple Information Gain

Our first approach for measuring the quality of a feature is the use of information gain of a feature $I_G(\varphi(x_j))$ to differentiate between all possible labels $\Sigma_j$. It is defined as

$$I_G(\varphi(x_j)) = I \left( \frac{h_1}{h} \right) - R(\varphi(x_j)) \tag{5}$$

where $I(\cdot)$ is the information content

$$I(p_1, p_2) = -p_1 \log_2 p_1 - p_2 \log_2 p_2 \tag{6}$$

with probabilities $p_1$ and $p_2$. $R(\varphi(x_j))$ is the remainder of bits of information after testing feature $\varphi(x_j)$:

$$R(\varphi(x_j)) = \sum_{\ell=1}^{\lvert \mathcal{L} \rvert} \left( \frac{h_\ell(\ell)}{h} \cdot \log_2 \left( \frac{h_j^1(\ell)}{h_\ell(\ell)} \cdot \frac{h_j^0(\ell)}{h_\ell(\ell)} \right) \right). \tag{7}$$

For comparing the features it is sufficient to compute $R(\cdot)$ because $I(\cdot)$ is constant for one feature [18]. Determining this measure for $m$ features with $\lvert \mathcal{L} \rvert$ different labels lasts $\mathcal{O}(m \lvert \mathcal{L} \rvert)$.

Ranking the features with this approach cannot lead to differences between transitions in the CRF. Therefore, it is referred to as Simple IG.

#### 3.1.2 Information Gain One-Against-All

The limitation of Simple IG is the disregard of differences between transitions in CRF. To cope with that, we assign a list of the best $p_{\text{frac}}$ features to every transition $\Sigma_j$. In general, every clique in the graph has its own evaluation of features $\varphi(\cdot)$.

The remainder, the measure for the quality of a feature in Equation 7, changes slightly to

$$R_{\text{OAA}}(\varphi(x_j), \Sigma_j) = \left( \frac{h_\ell(\ell)}{h} \cdot \log_2 \left( \frac{h_j^1(\ell)}{h_\ell(\ell)} \cdot \frac{h_j^0(\ell)}{h_\ell(\ell)} \right) \right) + \left( \frac{h_\ell(\ell)}{h} \cdot \log_2 \left( \frac{h_j^1(\ell)}{h_\ell(\ell)} \cdot \frac{h_j^0(\ell)}{h_\ell(\ell)} \right) \right) \tag{8}$$

\(^2\)For the linear-chain CRF of order 1. In general, $\Sigma_j \in \mathcal{L}^{c+1}$ with order $c$ holds.
Another well-known and often incorporated ranking method is IG, One-Against-All (IG-OAA) where

\[ \phi(x_j) = 1 \quad \text{or} \quad \phi(x_j) = 0 \]

\[ \sum_{\ell \in \{1, \ldots, |\mathcal{L}|\} \setminus \ell} \bar{h}_0(\ell) \bar{h}_1(\ell) \bar{h}(\ell) \]

and \( \bar{h}_0(l) \) and \( \bar{h}_1(l) \) analogous. This approach is applied to rank the features \( \phi(x_j) \) for every transition \( \mathcal{L}_\ell \) separately. It is referred to as Information Gain One-Against-All (IG-OAA). The runtime is \( O(m|\mathcal{L}|) \) and therefore less than Simple IG.

### 3.1.3 \( \chi^2 \)-Statistics

Another well-known and often incorporated ranking method are \( \chi^2 \)-statistics [16]. The \( 2 \times 2 \) contingency table is defined for each feature \( \phi(x_j) \) and each transition \( \mathcal{L}_\ell \) compared to all other transitions (cf. Table 1). The \( \chi^2 \)-statistic is then computed by

\[
\chi^2(\phi(x_j), \mathcal{L}_\ell) = \frac{(\bar{h}_1(\ell) \cdot \bar{h}_0(\ell) - \bar{h}_0(\ell) \cdot \bar{h}_1(\ell))^2 \cdot h}{\bar{h}(\ell) \cdot \bar{h}(\ell) \cdot \bar{h}_0(\ell) \cdot \bar{h}_1(\ell)} \tag{9}
\]

Similar to IG-OAA, this is performed to rank the features \( \phi(x_j) \) for every transition \( \mathcal{L}_\ell \) separately. Therefore, the runtime is also \( O(m|\mathcal{L}|) \). We refer to this method as \( \chi^2 \) OAA.

### 3.1.4 Random

The most simple method is a random ranking and selection of features. This is used as a baseline to evaluate the other measures presented in the previous sections.

### 3.2 Iterative Feature Pruning

Training a CRF is commonly performed by the iterative algorithm L-BFGS to assign weights \( \lambda \) to all feature functions \( f_j \in \mathcal{F} \) such that \( \mathcal{L}(T) \) is maximized (compare to Equation 4). Typically, many weights are close to 0. The idea of Iterative Feature Pruning (IFP) is that feature functions with low absolute weight value have a low impact on the output sequence.

Based on this assumption, the algorithm (see pseudo code in Figure 1) starts with a fully optimized CRF using all features representing the training data (Line 2). The next step is the removal of features with lowest absolute value (3). The parameter \( p = 1 - \frac{|S|}{|\mathcal{F}|} \) specifies the percentage of features to be removed in each iteration (13–15). \( S \) is the set of remaining features after one iteration of IFP.

In each step, a retraining with L-BFGS is performed to allow an adaption of the model by adjusting the weights for the remaining features (16). After that, the pruning is repeated (17) until no features are left (11).

During this process, at each iteration of pruning, the current performance of the model is evaluated and stored (8). This information can be used to select the final feature set (Line 4, an heuristic is shown in Section 4.3) and to train a full model with the identified feature subset (Line 5).

To illustrate the process of IFP, it is shown exemplarily by means of extreme examples for one data set

1. function IFP(crf,trainData, valData, p)
2. crf = BFGS(crf, trainData, valData)
3. log = PRUNING-STEP(crf, trainData, valData, p)
4. crf = SELECTFEATURESET(log)
5. crf = BFGS(crf, trainData+valData, null)
6. end function

7. function PRUNING-STEP(crf, trainData, valData, p)
8. log ← EVALUATE(trainData, valData, crf)
9. \( \mathcal{F} = GETFEATURESET(crf) \)
10. if \( \mathcal{F} = \emptyset \) then
11. return log
12. end if
13. S = features with lowest weights such that
14. \( p = 1 - \frac{|S|}{|\mathcal{F}|} \)
15. \( \mathcal{F} = \mathcal{F} \setminus S \)
16. crf = BFGS(crf, trainingData)
17. return PRUNING-STEP(crf, trainData, valData)
18. end function

Figure 1: Iterative Feature Pruning Algorithm (starting with method IFP(\( \cdot \) ) in Line 1)

### 4 Results

In this section, the methods proposed in Section 3 are evaluated on the data sets and configurations of the CRFs described in Section 4.1. The hypotheses to be analyzed are:

Selecting a reasonable subset of features:

A1 Improves explainability of the CRF model,
A2 Improves training time and tagging time which is beneficial for developing as well as applying the model,
A3 Improves performance in \( F_1 \) measure or does not decrease it dramatically.

Additionally, we assume that one method is superior to all others:

B One method outperforms the others.

1CoNLL data set introduced in Section 4.1

2Results not shown here due to page limitation.
The evaluations in the following form the basis for the discussion of these hypotheses in Section 4.4.

4.1 Data Sets used for Evaluation

The results and evaluations are shown on the basis of two data sets with slightly different configurations of the CRF. Quantities of entities are given in Table 2.

The BioCreative 2 Gene Mention Task data (BC2) contains entities of the class Gene/Protein with the specialty of acceptance of several boundaries for entities [24]. We incorporate the configuration of the CRF as described in a participating system using only the shortest possible annotation as exact true positive per entity [6, 21].

The CoNLL data [19] is an annotation of the Reuters corpus [10] containing the classes person, organization, locations and misc. We use an order-one CRF with offset conjunction combining features of one preceding and succeeding token for each position in the text sequence. The feature set is fairly standard with Word-As-Class, prefix and suffix generation of length two, three and four as well as regular expressions detecting capital letters, numbers, dashes and dots separately and as parts of tokens. The combination of the provided sets “train” and “testa” is used for training and “testb” for testing.

For evaluating the inference time on a larger set, a uniform sample from the Medline\(^5\) database of 10,000 entries is used additionally. Each one comprises titles, author names, and abstracts. The number of tokens is 958869 for BC2 and 960744 for CoNLL (additionally to BC2 tokenization, splitting on all dots is performed for CoNLL data).

4.2 Cross-Validation on the Training Sets

As a basis for parameter selection (presented in Section 4.3) and to evaluate the impact of feature selection, 10-fold cross- validation is used.

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### Table 2: Numbers of labels in data sets, the different entity classes are added in CoNLL.

| Name  | Training Set | B- | I- | O   |
|-------|--------------|----|----|-----|
| BC2   |              | 18165 | 15017 | 382983 |
| CoNLL |              | 29466 | 13180 | 213499 |

| Name  | Test Set     | B- | I- | O   |
|-------|--------------|----|----|-----|
| BC2   |              | 6290 | 4801 | 128915 |
| CoNLL |              | 5654 | 2458 | 38554 |

\(^5\)http://www.nlm.nih.gov/databases/databases_medline.html
validation is performed on the training sets (Section 4.1). The results are shown in Figure 3. The curves depict the average $F_1$ measure\(^6\) of the 10 partitions. The different numbers of features are detected with different parameters specified for the respective selection method. The transparent band around the line depicts the standard deviation for the according number of features.\(^7\) The significance of the difference of the methods is tested regarding the area under the curves in Figure 3 via Welch’s t-test with a significance level of $\alpha = 0.05$.

Comparing the results on the two data sets, the methods lead to similar results whereas the differences are clearer on CoNLL data. All approaches outperform the random selection significantly. The approach of $\chi^2$ OAA is worse than the conceptionally similar IG OAA and the more naive Simple IG on BC2 data.

IG OAA outperforms all other filtering approaches. Assuming the goal to reach the highest possible $F_1$ measure, IFP leads to better results than IG OAA on both data sets. Only if an extremely small number of features remains (fewer than about 50), IG OAA leads to better results. The superiority of IFP to $\chi^2$ OAA is significant ($p = 0.02$) on the CoNLL data set.

### 4.3 Results on independent test sets

We need to find the parameter assignment to determine the feature subset in the final model. For the filter approaches, the parameter is $p_{\text{thresh}}$. For IFP, the meaningful number of features at which the pruning has stopped is to be detected. Based on the smoothed\(^6\) values of $F_1$ measure in 10-fold cross-validation (see Figure 3), we define two measures to automatically detect these parameters: The maximally accepted loss in $F_1$ measure is denoted with $\Delta$, the threshold for the gradient is $g$. The detection of the feature subset is performed via backward selection starting with high numbers of features. The first position on the curve for which the gradient is smaller than $g$ or the $F_1$ measure is smaller than $\Delta$ is selected. The values $g = 2 \cdot 10^{-10}$ and $\Delta = 0.02$ lead to the positions denoted by arrows in Figure 3. The results for these values are evaluated in the following. The advantage of backward selection to forward selection is that it may capture interacting features more easily [8].

A model is built on the full training set applying IFP or filtering with the detected parameters. In Figure 4 the results on independent test sets mentioned in Section 4.1 are depicted. The smallest numbers of features are achieved by IFP followed by IG OAA.

The $F_1$ measures decrease up to the accepted $\Delta = 0.02$ on BC2 data and to a lower amount for the CoNLL data. The best trade-off between $F_1$ measure and the number of features (depicted in third bar chart) is always achieved by IFP followed by IG OAA.

A smaller number of features should induce a faster model in training and inference. The time of evaluating $P_X(g\|z)$ (compare to Equation 1) on all training examples is shown in the fourth bar charts. This computation is crucial for training durations as it has to be performed many times. These numbers correspond roughly to the number of features, the durations are smaller than for the原始 model. The ratio is not linear due to the necessary forward-backward algorithm calls (whose runtime is quadratic in the number of possible labels) and dynamic memory allocation times.

These numbers naturally lead to dramatically reduced training times. Training the full CoNLL model lasts 5788 seconds, 2397 s for BC2 respectively. With the feature set detected by IFP, these numbers reduce to 2156 s and 670 s. This improvement is not helpful in practice, as the IFP procedure incorporates training the model. However, filtering via IG OAA improves overall training time as it is computationally inexpensive. It leads to 5230 s and 1002 s for training a model.\(^9\)

Reducing the number of features also leads to a faster inference\(^10\). The fifth bar charts show durations for tagging 10000 sampled abstracts from Medline. Best results are achieved by IFP (CoNLL: 10.52 s instead of 17.56 s, BC2: 2.42 s instead of 4.56 s), followed by the filtering methods which do not differ remarkably (IG OAA: CoNLL: 14 s, BC2: 2.65 s).

Summarizing, a reduced training iteration time of 76% of original time evaluating $P_X(g\|z)$ with a loss of only 1.7% $F_1$ (absolute value) on the BC2 data is possible with IFP. The tagging time is reduced to 53%. On the CoNLL data, a loss of only 0.57% in $F_1$ occurs with savings of even 54% of original computing time. Tagging time is reduced to 60%.

### 4.4 Discussion

Comparing the methods, the results are similar for the data sets: In 10-fold cross-validation, the random method is dominated by all other methods. Simple IG or $\chi^2$ OAA are second worst, depending on the data set. IFP is the best method, closely followed by IG OAA.

The Simple IG lacks the representation of different transitions in the features which is especially important for the CoNLL data with 4 entity classes of interest. No dictionaries with members of these classes have been used in the presented setting, so all classes are memorized with automatically generated features, hence, a large number of different features is needed for the different transitions in the CRF.

The method $\chi^2$ OAA always leads to worse results compared to IG OAA on the 10-fold cross-validation although it is a systematically similar approach. The reason is presumably the unbalancedness of the labels in the generated classification instances\(^11\) which is taken into account in the

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\(^6\)The relation to the numbers of features and the iteration durations is not linear as the needed numbers of training iterations differs.

\(^7\)Measuring only the computation, not the time to read the data from hard disk and to extract the features.

\(^8\)Transitions of intermediate terms (like $(O,O)$) are for instance much more frequent than those of beginnings of entities (like $(B,B)$).
information remainder (Equation 7 and 8) but not in $\chi^2$ (Equation 9).

IFP leads to better results than IG OAA for high $F_1$ measures. The reason is the limitation of IG OAA to use the same number of features (but not the same set) for each transition (specified by $P_{rune}$). This does not hold for IFP as it only relies on the model structure itself. The drawback is the higher computational cost due to the incorporated L-BFGS optimization.

The random method does not lead to good results, but it should be noted, that even this approach can remove 30\%–40\% without a dramatic decrease in $F_1$ measure. The reason are unessential redundancies in the full feature set.

Mainly all these results are reflected on the independent test sets. IFP or IG OAA has the best trade-off between $F_1$ measure and the number of features. Good inference speed-ups can be achieved with both methods, corresponding to the low numbers of features. However, IFP cannot be used to speed up training as it incorporates the training procedure itself. Hence, IG OAA is proposed for this instance. For reducing tagging time as well as for understanding the model, IFP should be used as it leads to the smallest numbers of features.

In Table 3, the percentages and numbers of remaining features accepting maximally 0.01 loss in $F_1$ measure are depicted. Much more features can be ignored in the BC2 than in the CoNLL setting. This can be an indicator for a possible improvement by implementation of features with better generalization properties. Detected relevant features representing words or important pre- and suffixes help understand the different entity classes of interest. The fact that noisy features are removed allows for an investigation of the remaining features which can be assumed as meaningful.

We conclude the results with an investigation of the hypotheses:

A1 The explainability of models applying feature selection is improved by the lower complexity. The need for many automatically generated features can be an indicator for a possible improvement by implementation of features with better generalization properties.

A2 Training and tagging time are decreased by the lower complexity of the CRF. To improve training time, the use of a filtering approach like IG OAA is proposed as these methods are computationally inexpensive. To improve tagging time, IFP should be used as it leads to the lowest numbers of features.

A3 A small decrease in $F_1$ measure has to be accepted for the benefit of a model with a considerable lower number of features.

B The recommendation for a method depends on the application: For improving training time, a filtering method should be used, preferably IG OAA as it shows best results. For improving tagging time, the computationally more expensive IFP can be applied.

12 Lists of all features of the models are available online to demonstrate the explainability comprehensively. http://www.scai.fraunhofer.de/ranlp-crf-fs.html
5 Conclusion and Future Work

A huge number of features is typically used to represent input text in CRFs. We presented different approaches for feature subset selection, novel adaptations of filtering to the sequential structure of text as well as an iterative method. The methods have been evaluated on two domains, showing a decrease of computing time and complexity of the model. The $F_1$ measure varies slightly.

Summarizing, IG OAA is the best filter approach, a lower number of features can only be achieved with Iterative Feature Pruning (IFP) with a similar $F_1$ measure. IFP relies only on the CRF structure itself, so it is able to deal with different numbers of features per transition in contrast to the One-Against-All methods. Its main disadvantage is its higher computing cost due to the incorporated training process. It is notable that IFP and IG OAA are methods taking the sequential structure of the text into account, IFP via using the model itself, IG OAA via different feature sets for different transitions. The method Simple IG, which does not select features with respect to transitions, leads to worse results.

Building a new Named Entity Recognizer often includes annotation of a corpus. It has to be investigated, how the need for features changes during the process of enriching the training set with examples (e.g. via active learning).

Another point is that the proposed methods allow for more complex feature generations (e.g. with more context information). It has to be studied if new features, which could not be implemented before due to an exhaustive memory consumption or run-time demand, could improve the state-of-the-art results.

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