Notes on Grobid

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1 Introduction

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1.1 Terminology

We briefly present important terms found in this report. Words contained in the same entry are used interchangeably.

- Grobid the Java-based tool
- Wapiti the principal CRF engine used by Grobid
- CRF++ a secondary CRF engine available to Grobid
- Token a character string or "word", delimited from other tokens by white space—the atomic unit of data.
- Observation a sequence of tokens subjecting to a sequence of classifications
- Class(affection)/label/annotation used interchangeably to refer to the prediction of a single token.
- Document/article -
- Corpus/training set the body of annotated documents used for creating a model
- Evaluation/test set the body of annotated documents used for evaluating a model
- Header/front matter the parts of the document containing identifying information, typically located at the very top of an article, prior to the introduction. Header matter is however often discontinuous, found after the start of the document body, in the first footnote or even at the very end of the article. Not to be confused with a headnote (the opposite of a footnote).

- 2 The Problem Automatic Metadata Extraction
- 3 Conditional Random Fields and Probabilistic Approaches
- 4 Results
- 5 Appendix

6 Notes - Work In Progress

Grobid (GeneRatiOn of BIbliographic Data) is an open source (Apache license) Java tool for applying conditional random fields (CRF) to the problem of metadata extraction of scientific papers. The tool is a wrapper that sits atop a CRF engine that performs the abstract CRF computations. Wrapper and engine are combined through the use of the Java Native Interface (JNI). Grobid is divided into four packages: grobid-home, grobid-core, grobid-trainer, and grobid-server. Grobid produces a set of models, each addressing a different part of the metadata extraction, and each trained separately on its own spectrum of features. In the case of citation processing, the models are applied according to a hierarchy, starting with the segmentation of the reference list, followed by the tagging of each part of each reference, and finally a set of micro-models to further classify these reference parts.

6.1 Engines

Grobid allows for a choice between two alternative linear-chain CRF engines (both written in C++): CRF++ (developed by Taku Kudo) and Wapiti (developed by Thomas Lavergne). The default option is Wapiti, a more recent engine that allegedly performs better. Both engines follow similar input and output formats, allowing an easy abstraction within Grobid over the choice of engine. Though we have access to the source code, the CRF engine will be largely black-boxed in this project, and we will instead concentrate on tuning and configuring Grobid.

6.2 grobid-home

The grobid-home package houses the models, configuration files and additional software components such as pdftoxml, which is used to extract both plaintext and OCR data from the PDF inputs. This package does not contain any source code.

6.3 grobid-core

The grobid-core package contains the majority of the source code, including that for modelling documents and interacting with the CRF engine. grobid-core also supplies the main entry points for using Grobid as a command line tool. Through this interface, we may use Grobid to batch process collections of PDF files.

6.4 grobid-trainer

grobid-trainer contains the source code for coordinating the training of the models, including preparing the training data from the TEI inputs. grobid-trainer provides the main entry point for training models and can also be used for training models through the command line.

6.5 grobid-server

grobid-server onsists of a web interface for Grobid, running on Jetty (Java) HTTP server. It allows a user to upload a PDF through a web form for processing. This feature is unlikely to be of any interest to this project.

6.6 Models

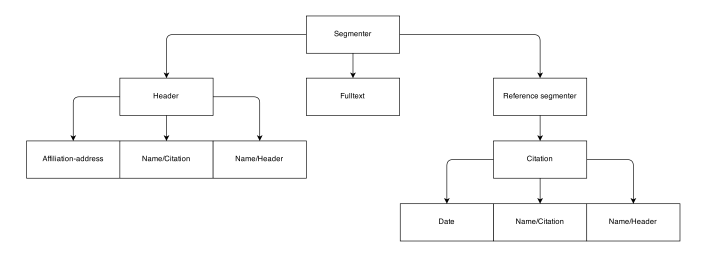


Figure 1: Cascade of models used by Grobid

Finding the header is typically easy, as we can assume it always to constitute the first page or so of the document. For this reason, when we call processHeader on its own, the Segmentation model is not called. On the other hand, though references are almost invariably at the end of an article, it is difficult to guess where it begins, and the reference model would suffer more from garbage than the header model. For this reason, the Segmentation model is called when we call processReferences, before calling the Reference-segmenter model to slice the reference list (identified by Segmentation) into individual citations.

Grobid manages the training, evaluation and usage of a set of models, each addressing a part of the information extraction. Higher level models such as the Header, Segmentation, and Reference Segmentation models may applied directly to PDFs, while the other, more specific models, such as the Date model, operate only on plaintext inputs. Though they have much in common, the models vary in the labels they assign, the features they exploit, and, due to the varying size of the vocabulary (compare say, the number of possible month names to the number of possible author names), the size (dimensionality) of the models. Calling processFullText runs all available models on a batch of PDF documents. The output of training is a model, which takes takes the form of a text or binary file, depending on the engine. These models are then "loaded" at prediction time for the labelling of new documents. Some models are in practice not used independently, rather, they form part of a "cascade" of models that progressively address finer subcomponents of the classification problem. As shown in Figure ??, reference extraction begins with

[...]

```
January january J Ja Jan Janu y ry ary uary LINESTART INITCAP NODIGIT 0 0 1 NOPUNCT I-<month> 1994 1994 1 19 199 1994 4 94 994 1994 LINEEND NOCAPS ALLDIGIT 0 1 0 NOPUNCT I-<year>
July july J Ju Jul July y ly uly July LINESTART INITCAP NODIGIT 0 0 1 NOPUNCT I-<month> 1996 1996 1 19 199 1996 6 96 996 1996 LINEEND NOCAPS ALLDIGIT 0 1 0 NOPUNCT I-<year>
[...]
```

Figure 2: Training input for Wapiti

```
# Capitalization
U50:%x[0,11]
U51:%x[1,11]
U52:%x[-1,11]
U53:%x[0,11]/%x[1,11]
U54:%x[-1,11]/%x[0,11]
```

Figure 3: Feature templates for Wapiti

segmentation models, which classify each line of a document, resulting in homogenous blocks of lines e.g. header, paragraph, figure, references. This information is then distributed to the other models, for example the Reference Segmentation model, which further breaks down the reference list into individual references. The Citation model then classifies the parts of each reference into classes, for example, date, affiliation, and author. Finally, the atomic subcomponents of these are classified by their respective models. Note that the citation branch of the hierarchy has the option of further cross-checking extracted references with the third-party CrossRef web service. Thus, the overall accuracy of the system is dependent on the combined accuracy of models.

6.7 Training

Both training and evaluation are performed on sets of XML documents following the Text Encoding Initiative (TEI) standard for representing electronic texts (http://www.tei-c.org/index.xml). This is also the output format for prediction, so there is consistency between input and output formats. It may appear paradoxical to *evaluate* on well-structured data, when the tool is ultimately intended to operate on unstructured PDF documents, that is, at prediction time. However, with closer inspection of the source code, an equivalence can be seen between:

- 1. applying an OCR tool (pdftoxml), tokenising the output and transforming to CRF input data
- 2. extracting tokens from TEI documents, and transforming to CRF input data

Both approaches yield the same input data for the CRF engine, and so evaluation is in fact equivalent to prediction, despite the initial difference in input formats. Figure ?? shows an excerpt from an input file to the CRF engine for training. These features are for inputs "January 1994" and "July 1996", for training the Date model. The features range from token identity, to a variety of prefixes and punctuation features. It should be noted that OCR information is only used in higher level models, that is, the Header and Segmentation models. The input for lower-level models such as Date is plaintext, and so features

are typically simple, but dictionary-based features, where information about a token is referenced in a dictionary resource within Grobid, are also used. Note the features shown are only those pertaining to the token itself. The full range of features (including those involving concatenations of the token's neighbours etc.) are defined by a set of feature templates. The feature templates for each model are contained in a separate file. An excerpt of this is shown in Figure??. These are given as a separate input to the CRF engine, and it is with these that the engine constructs all feature functions for the model. It is therefore vital that the feature extraction, which is generated by Grobid, is aligned with the template file, which is manually configured by the developer. As depicted in Figure ??, there is a strong coupling between these two parts of Grobid. The excerpt shown is from the Wapiti model, but the notation is the same for CRF++, which first standardised the syntax. This subset of five feature templates capture information about the capitalisation of a token and its neighbours. The notation has the structure, [identifier]: [%x][row, col], where row is the offset from the current token, and col indicates the feature index. Thus, "U50:%[0,11]", denotes that the feature template identified as "U50" takes the 11th feature for the current token (0 offset). This feature will be equal to 1 if a token is capitalised, and 0 otherwise. "U52:%[-1,11]" indicates the same thing, but based on the capitalisation of the previous token. "U54:%x[-1,11]/%x[0,11]" is a binary function for detecting the capitalisation of the current and the following token.

Now we may see an alignment with the mathematical model. Recall a linear chain CRF is expressed in the simplest case as,

$$p(\mathbf{y}|\mathbf{x}) = \frac{p(\mathbf{x}, \mathbf{y})}{\sum_{y'} p(\mathbf{x}, \mathbf{y}')},\tag{1}$$

where,

$$p(\mathbf{x}, \mathbf{y}) = \exp \left\{ \sum_{t} \sum_{i,j \in S} \lambda_{ij} \mathbb{1}_{\{y_t = i\}} \mathbb{1}_{\{y_{t-1} = j\}} + \sum_{t} \sum_{i \in S} \sum_{o \in O} \mu_{io} \mathbb{1}_{\{y_t = i\}} \mathbb{1}_{\{x_t = o\}} \right\},$$
(2)

Here **x** is a sequence of observations and **y** is a sequence of labels. S is the set of all labels, O is set of observations (the vocabulary of the tokens to be labelled). When the coefficients $\lambda_{ij} = \log p(y_t = i, y_{t-1} = j)$ and $\mu_{ij} = \log p(y_t = i, x_t = o)$, this joint distribution is equivalent to a Hidden Markov Model (HMM), with coefficients, λ_{ij} as transition probabilities and μ_{ij} emission probabilities. In this simple case, features are based solely on the token's identity, i.e. feature functions are an indicator function. For clarity, we may write,

$$p(\mathbf{x}, \mathbf{y}) = \exp \left\{ \sum_{i \in S} \sum_{j \in S} \lambda_{ij} F_{ij}(\mathbf{y}) + \sum_{i \in S} \sum_{o \in O} \mu_{io} F_{io}(\mathbf{x}, \mathbf{y}) \right\},$$
(3)

where $F_{ij} = \sum_t \mathbb{1}_{\{y_t=i\}} \mathbb{1}_{\{y_t=i\}} \mathbb{1}_{\{y_t=i\}} \mathbb{1}_{\{x_t=o\}}$. In a CRF, however, we may replace the indicator function for observations with any sort of function, typically binary, extracting rich features from a token. Thus, $F_{io} = \sum_t \mathbb{1}_{\{y_t=i\}} f_{io}(\mathbf{x})$. The set of functions, $\{f_{io}\}$, are the functions that we define in the feature template files. Note that, unlike an HMM, the vocabulary is not pre-defined, it is "discovered" through training on samples. Therefore, the number of actual features depends on the training set itself, whereas the feature template is fixed. Since we use indicator functions, which produce a feature for every observation, we may end up with an enormous number of features. Take the Date model for example: 5815 features are produced for a single block (not counting the one representing the label), and there are seven labels. As per our formulation in (??) we therefore have 7 * 7 "transition" features and 5815 * 7 "emission" features, totalling 40754 features. This is corroborated by the model output in Figure

```
* Initialize the model
* Summary
   nb train:
                 493
   nb labels:
                 7
   nb blocks:
                 5816
   nb features: 40754
* Train the model with l-bfgs
                        act=16482
  1] obj=1688,58
                                      err=25,80%/50,91% time=0,08s/0,08s
  2] obj=1221,30
                                      err=19,11%/35,50% time=0,05s/0,12s
                         act=15580
                                      err=17,20\%/33,67\% time=0,04s/0,17s
  3] obj=922,15
                         act=13869
  4] obj=638,04
                         act=10845
                                      err= 6,53\%/15,21\% time=0,04s/0,20s
  5] obj=478,72
                                      err= 5,68\%/13,59\% time=0,04s/0,24s
                        act=10582
  Γ
      6] obj=416,15
                         act=9926
                                      err= 3,77\%/ 9,53\% time=0,04s/0,28s
```

Figure 4: Output from training date model

```
8:u00:_x-3,

8:u01:_x-2,

8:u02:_x-1,

12:u03:November,

6:u04:19,

5:u05:,,

8:u06:2001,

17:u07:_x-1/November,

15:u08:November/19,

8:u09:19/,,

13:u0A:_x-2/_x-1,

8:u10:_x-2,

8:u11:_x-1,
```

Figure 5: Excerpt of model contents—the realisation of feature functions

??. Wapiti automatically constructs this vast feature space from the inputs we provide. In the Date model, the labels are I-<day>, I-<month>, I-<year>, I-<other>, <day>, <month>, and <other>. The I (probably) stands for "initial", as in training these are assigned to the first tokens of this class found in the string.

A model is typically a large file (as much as 100Mb). At the top of the file, the feature templates are declared, just as they are in the input. Because of this, that file is not required at prediction time. Following this the labels are declared. Then come two longer sections: first, the feature functions themselves as defined. Figure ?? shows the first 12 features produced from the first token in the first sample in the training set—"November". Because this is the first token in the string, we see the first three feature macros, which relate to the identity of the token's predecessors, remain unresolved. The fourth, however, shows the indicator for the token. This function will be true if a token is equal to "November". The fifth function is an indicator for if the token's successor is equal to "19", and so on. The final (and usually largest) section of the model file defines the non-zero weights for the feature functions. The weights are represented in scientific notation and in hexadecimal representation, presumably to avoid arithmetic underflow (a common problem when dealing with with the computation of HMMs and related models).

6.8 Evaluation

Training may be done with a split defined by the developer, which Grobid will use to set aside a proportion of the training data for evaluation. The evaluation of a model produced by training follows identical procedures, preparing the same input data. The output, however, is not a model but a the input data with labels. Grobid compares its predictions with the ground truth and outputs precision, accuracy and F1 scores as performance indicators, at the token, field, and instance levels. A token refers to a single contiguous string of characters (without spaces), a field is a block of contiguous tokens, and an instance is an entire document. Accuracy of an instance is therefore judged by the correctness of all tagging for the whole document, a difficult thing to achieve without any mistakes.

6.9 Prediction

Figure ?? shows the flow of information from input to output, as well as the relationship between training and prediction. When it comes to labelling (prediction), the starting point is a PDF document. With a third-party tool, pdftoxml, this is transformed into an XML file containing OCR information (font, style, orientation) for every token in the document. This information is stored in LayoutToken objects within Grobid. These tokens are arranged into blocks and features are extracted as they were for training and evaluation. Excepting the absence of tokens, the input is the same. The model created in the training phase is first loaded, and then the EngineTagger calls the CRF engine to label the inputs. Unlike for training, the feature template file is not required, as these have already been absorbed into the model file. After processing, Wapiti returns the same file with tags inserted. Grobid then further processes this information to transform it into the final TEI format.

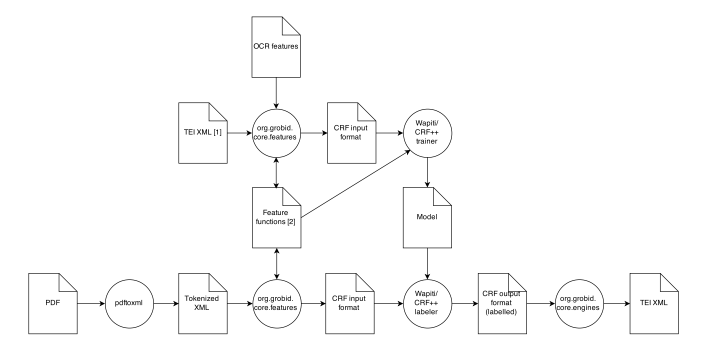


Figure 6: The flow of information and the interaction of training and prediction. [1] Training inputs follow and TEI standard. [2] There is a strong coupling between the features module and the feature functions file.

6.10 Other Functionality

In addition to the above, Grobid provides a means of producing training sets semi-automatically. This consists of applying the existing models on the training set to produce the XML inputs. Of course, each field must be checked against a ground truth and errors corrected before it is used as a training set. We intend to use this functionality to generate training data for benchmarking Grobid on HEP papers.

These methods are,

- 1. createTrainingHeader
- 2. createTrainingFulltext
- 3. createTrainingSegmentation
- 4. createTrainingReferenceSegmentation
- 5. createTrainingCitationPatent

Now, for example, when we run **createTrainingHeader** to create a header training set, grobid creates four files per pdf:

- 1. PDF-NAME.affiliation.tei.xml
- 2. PDF-NAME.authors.tei.xml
- 3. PDF-NAME.header.tei.xml
- 4. PDF-NAME.header

Files (1) and (2), are the training data for the affiliation-address and author models respectively. Files (3) and (4) are jointly required for training the Header model: file (3) simply contains the tagged data, but file (4) contains the feature (untagged) for each token, including OCR information. We must additionally store (4) as the OCR information cannot be derived from the plaintext TEI file. That the processing does not tag the blocks allows the programmer to intervens and correct and discrepancies with the ground truth. Thus the header files need only be corrected then moved to a separate folder for training. Notice training data for the segmentation models may be harder to generate automatically.

Notice that there is no method for creating citation training data, \mathcal{D}_{tr} , directly. This data is created as part of createTrainingFulltext. This creates four files also:

- 1. PDF-NAME.training.citations.authors.tei.xml
- 2. PDF-NAME.training.references.tei.xml
- 3. PDF-NAME.training.fulltext.tei.xml
- 4. PDF-NAME.training.fulltext

File (1) contains a refinement of the author names found in file (2), thus can be used as training data for the Name-citation model. File (2) contains the training data for the Citation model, in the format apparent in the training corpus. As should be expected, the name and date fields for this data are not resolved into their subcomponents, this task being left to the smaller models dedicated to these tasks. Files (3) and (4) are the files required for training the citation model.

Model	Labels
Header	<title>, <author>, <affiliation>, <reference>, <submission>, <abstract>, <ad-</td></tr><tr><td></td><td>dress>, <keyword>, <degree>, <pubnum>, <email>, <date>, <copyright>, <in-</td></tr><tr><td></td><td>tro>, <web>, <note>, <phone>, <dedication>, <entitle>, <grant>, <date-</td></tr><tr><td></td><td>submission></td></tr><tr><td>Affiliation address</td><td><pre><institution>, <other>, <settlement>, <department>, <postcode>, <contry>,</pre></td></tr><tr><td></td><td><pre><marker>, <region>, <addrLine>, <laboratory>, <postbox>, <other>, <null></pre></td></tr><tr><td>Name/Header</td><td><pre><forname>, <surname>, <marker>, <middlename>, <other>, <suffix>, <title></pre></td></tr><tr><td>Name/Citation</td><td><pre><surname>, <forname>, <other>, <middlename></pre></td></tr><tr><td>Citation</td><td><pre><journal>, <volume>, <other>, <issue>, <pages>, <date>, <author>, <title>,</pre></td></tr><tr><td></td><td> <booktitle>, <location>, <pubnum>, <note>, <publisher>, <editor>, <institu-</td></tr><tr><td></td><td>tion>, < tech>, < web>, < issue></td></tr><tr><td>Date</td><td><ohresidasis <ohresidasis <ohresidasis <ohresidasis <ohresidas <oh</td></tr><tr><td>Segmentation</td><td><pre><headnote>, <header>, <body>, <page>, <references>, <footnote>, <cover>,</pre></td></tr><tr><td></td><td><acknowledgement>, <annex></td></tr><tr><td>Reference-Segmenter</td><td><label>, <reference>, <other></td></tr><tr><td>Fulltext</td><td><pre><section>, <paragraph>, <citation_marker>, <other>, <table_marker>, <fig-</pre></td></tr><tr><td></td><td>ure_marker>, <figure_head>, <trash>, <figDesc>, <equation>, <item></td></tr></tbody></table></title>

Table 1: We have here excluded the Patent, Entities, and E-book models as these are experimental models not currently used by Grobid. Of the models listed, we will be focusing on Header, probably Segmentation, and perhaps Reference-Segmenter and Citation also. This is because these are likely to require special training and configuration for HEP papers. Others such as Name and Date are unlikely to improve with training on HEP papers.

```
<?xml version="1.0" encoding="UTF-8"?>
<tei xmlns="http://www.tei-c.org/ns/1.0" xmlns:xlink="http://www.w3.org/1999/xlink"</pre>
xmlns:mml="http://www.w3.org/1998/Math/MathML">
<teiHeader><fileDesc><sourceDesc><biblStruct><analytic>
<author>
<persName>
<forename>B</forename>.
<surname>GARRIGUES</surname>
</persName>
ΕT
<persName>
<forename>A</forename>.
<surname>MUNOZ</surname>
</persName>
</author>
</analytic></biblStruct></sourceDesc></fileDesc></teiHeader>
</tei>
```

Figure 7: Sample TEI (input and output) format for the Name-header model.

```
<bibl>
<label>[20]</label>
D. A. Hill, "Fields of horizontal currents located above the earth, " IEEE Transactions on Geoscience and Remote Sensing, vol. 26, no. 6, pp. 726-732, 1988. [21]
A. M. Badescu, V. Savu, and O. Fratu, "Preliminary tests in Unirea salt mine (Slanic Prahova, Romania), "Report 4, University Politehnica of Bucharest, Bucharest, Romania, 2011.
</bibl>
<br/>
M. Stefanescu, O. Dicea, and G. Tari, "Influence of extension and compression on</br>
```

Figure 8: Sample error made by the (cora-trained) reference-segmenter model: the model fails to spot the citation label for reference [21], and the references are consequently merged. The citation model will then fail to match the ground truth.

6.11 Pipeline

6.12 Training - References

To prepare Grobid for practical reference extraction, our task is twofold, involving the training of both the reference-segmenter and citation models. The overall success of the citation model in tagging an entire reference list depends on the reliability of the citation blocks fed to it by its parent model. To illustrate, even a citation model that is perfect in its own right will not help very much in the reference extraction if the reference-segmenter above it is incapable of finding the references. Note that, to be precise, there is a further dependency on the accuracy of the segmenter model, the most high-level model in Grobid's arsenal, which is responsible for identifying the reference section for the before it is segmented, however we will assume temporarily that this is accurate enough in practice. Similarly, the date and name models are assumed to be sufficiently accurate, insofar as they are supplied with the correct text by the citation model, and so improvement of these will not be a focus of the project. Suffice to say, these models are unlikely to require special training on HEP papers. *Instance-level* accuracy for the citation model, measured over a whole document, will be the number of correctly tagged references as a fraction of the references correctly found by the reference-segmenter.

By contrast, the training of these two models may be performed independently. It does not matter to the citation model if the references in its training set are not contiguous members of the same reference list; the citation model operates at the reference string level. In the same way, it does not matter to the date or name models whether the dates or names they are training on or tagging belong to the same document. For this reason, creating our training set (as per the pipeline) for the citation model is simplified: if, when preparing the set, we notice a ground truth reference is missing from the output, we can just ignore this and continue. On the other hand, it matters very much that the reference-segmenter identifies the correct number of references, as this is the goal of the model, and it has a knock-on effect for the citation model below it. See for example Figure ??, showing an error made by the pre-trained (Cora) reference-segmenter model shipped with Grobid. It is rather surprising that such obviously strong indicators as the square brackets in the citation label, "[21]", are insufficient to classify it as such, especially when this error seems to happen rather often. This may be the result of poor feature engineering, a limited test set, or a mixture of the two. To see how citation model would tag the correctly segmented reference "[20]", see Figure ??.

If we are to correct the segmentation done when creating the reference segmenter training set, it is useful

<bibl><author>D. A. Hill</author>, " <title level="a">Fields of horizontal
currents located above the earth</title>, " <title level="j">IEEE Transactions
on Geoscience and Remote Sensing</title>, vol. <biblScope type="vol">26
</biblScope>, no. <biblScope type="issue">6</biblScope>, pp. <biblScope type="pp">
726âĂŞ732</biblScope>, <date>1988</date>. [<editor>21] A. M. Badescu, V. Savu, and
O. Fratu</editor>, " <title level="a">Preliminary tests in Unirea salt mine</title>
(<pubPlace>Slanic Prahova, Romania</pubPlace>), " <note type="report">
Report 4</note>, <orgName>University Politehnica of Bucharest</orgName>,
<pubPlace>Bucharest, Romania</pubPlace>, <date>2011</date>.</bibl>

Figure 9: Citation tagging of the correctly identified reference number 20 as shown in Figure ??.

to note segmentation is done simply by placing the delimeters, i.e. the labels. If the delimeter is correctly placed, the entire reference is correctly segmented. We may therefore proceed by checking the delimeters. If one of them is missing, it indicates a problem such as shown in Figure ??, and further processing is required. Again, we are assuming the reliability of the OCR tool and the initial segmentation model to provide the reference-segmenter with the correct block of text as reference list.

6.13 Training - Segmentation

The segmentation model models the following

8:<header>, ÃċÅŞâĂIJ
6:<body>, ÃċÅŞâĂIJ
12:<references>, ÃċÅŞâĂIJ
10:<footnote>, ÃċÅŞâĂIJ
17:<acknowledgement>, ÃċÅŞâĂIJ
7:<annex>, ÃċÅŞâĂIJ
10:<headnote>, ÃċÅŞâĂIJ
8:I-<page>, ÃċÅŞâĂIJ

A publisher's cover page is put in the tags </titlePage>. This probably corresponds with </cover>
In an abuse of notation, training data leaves the </teiHeader> tags essentially empty and in fact per

</body> tags encapsulate blocks of text belonging to the body, but these are interrupted by </page>

<note place="footnote">is for footnotes, (ultimately transformed into <footnote>). It seems to get

<div type="acknowledgement"></div> identifies an acknowledgement section within the document.

</listBibl> indicates the reference list as a section.

</annex> is apparently for information outside the normal flow of the document. These might be appeared

Remember: tokens are lines in the segmentation model (as they are in fulltext)

one right, the tags in a window ranging from three left to three right, and the IOB suggestions for the previous two tokens. 2

3) In the reverse WPDV model, the direction of chunking is reversed, i.e. it chunks from the end of each utterance towards the beginning.

4) In the R&M WPDV model, Ramshaw and Marcus's type of IOB-tags are used, i.e. starts of chunks are tagged with a B-tag only if the

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nequence phrase type, the 141, and especially on one weak point: determination of the start position of NPs. I use separate WPDV models for each of the following cases:

1) Should a token now marked I-NP start a

³Cf. van Halteren (2000a). Also, the difference between training and running (correct IOB-tags vs model suggestions) leads to a low expected generalization quality of hill-climbing. I therefore stop climbing after a single effective step, but using an alternative climbing procedure, in which not only the single best multiplications/division is applied per step, but which during every step applies all multiplications/divisions that yielded improvements while the opposite operation did not.

Structural studies of genomes and proteomes are essentially limited to a small number of dimensions. The primary structure of a polynucleotide or a polypeptide is a one-dimensional entity, whereas folded structures can be traced in three-dimensions, and their temporal changes can be described by adding time as a fourth dimension. In contrast, functional properties can be expressed in numerous dimensions defined by the parameters investigated.

In addition to the eventual goal of evolving enzymes with

In addition to the eventual goal of evolving enzymes with tailor-made activities, we wanted to explore the distribution of members of a mutant library in functional substrate-activity space. This information could provide a basis for choosing the optimal variants from a suitable 'quasi-species' of a mutant library (Eigen et al., 1988) to be parents for a subsequent generation of mutants. The optimal parentage for a new generation is not limited to the mutants that have evolved the furthest in a desired direction, since their genetic background may be too narrow (Ness et al., 1999). Therefore, the identification of an entire group of near-optimal mutants is an important task.

near-optimal mutants is an important task.

The gene superfamily of glutathione transferases (GSTs) illustrates how similar structures can display widely different properties. GSTs were originally discovered as a group of detoxication enzymes catalyzing the conjugation of a wide variety of electrophilic substances with the sulfhydryl group of the tripeptide glutathione (GSH) (Josephy and Mannervik, 2006). The three-dimensional structure is highly conserved

Things become contentious when head- and footnotes appear in positions. It's important to recognise that Grobid is relying on the OCR tool for the correct sequencing of these elements as they appear on the page. This implies that if the OCR tool dependency were to change, we should revise the training set. It is possible, on the other hand, that the OCR tool provides a standard representation, and any alternative would provide Grobid with the same information. In any case, blocks of text e.g. footnotes etc. should not be moved around in the correction process, even if their places appears illogical. Only the tags themselves should be moved where required. This is best illustrated in the case of an article whose body is arranged into two columns of text. Where there are footnotes, these are considered to be part of the column, and so, though it may appear more logical to declare all the footnotes are at the base of the page and should all come after the body sections as a group, the tool will order them thus:

6.14 Evaluation

6.14.1 Why we use the F1 Statistic

Accuracy is defined to be, $\frac{TP+TN}{TP+FN+FP+TN}$, that is, the proportion of correct classifications to total classifications, where TP is the number $true\ positives$, the correctly predicted positive classes, where positives indicates any given class; TN is the number of $true\ negatives$, the correctly predicted $negative\ classes$; FN is the number of $false\ positives$, the incorrectly predicted positive classes; and TN is the number of $true\ negatives$. Accuracy can be a misleading statistic when we have uneven representations of classes in the dataset. In the event we have high bias on the number of negatives, we can achieve excellent accuracy simply by always predicting the negative class. For this reason, we consider other statistics too. $Precision\ is$ the proportion of positives correctly claimed, i.e., $\frac{TP}{TP+FP}$. $Precall\ is$ the proportion of positives correctly predicted to the total number of positive occurrences (equivalently, the accuracy over the positive class), i.e., $\frac{TP}{TP+FN}$. Notice that in the case described above, both precision and recall would be 0.

The F_1 statistic is a common measure used to assess binary classifiers, and is defined as,

$$F_1 = \frac{2 \times \text{precision} \times \text{recall}}{\text{precision} + \text{recall}},$$

that is, the harmonic mean of precision and recall (the "1" in F_1 indicates the two are evenly weighted).

¹Cf. http://ilk.kub.nl/.

²For unseen data, i.e. while being applied, the IOB suggestions used are of course those suggested by the model itself, not the true ones.

The F_1 statistic is a nice way of summarising both at once as it is simply the harmonic mean of the two. Furthermore, because of this, a large imbalance in recall and precision resuts in a lower F_1 score. It is necessary to be good in both precision and recall to have a good F_1 score; the harmonic mean of any data is always upper-bounded by its arithmetic mean

6.14.2 Evaluation in Grobid

In Grobid, evaluation is done at the token, field, and instance levels, that is, Grobid calculates the aforementioned statistics for individual token or words, and then for the classes themselves. Finally, it calculates the number of correct instances, that is, entire samples with no classification errors.

We supplement the Grobid evaluation output with a confusion matrix, and example of which is shown in Figure XX. Whereas the statistics allow us to compare one model to another, a confusion matrix can be used to see exactly which misclassifications are being made, which can in turn inform our feature engineering.

6.14.3 Modelling Collaborations

joseph@joseph-HP-Compaq-8200-Elite-CMT-PC:~/Desktop/grobid/grobid-trainer/resources/dataset/header../55004899.header.tei.xml:17: <note type="other">This is a reprint of a Cochrane review, prepared a ./55004899.header.tei.xml:23: <note type="copyright">Copyright Âl' 2009 The Cochrane Collaboration.

- ./header994.tei:11: <note type="other">for the STAR Collaboration<lb/></note>
- ./55000939.header.tei.xml:10: <titlePart>CoDoc: Multi-mode Collaboration<lb/> over Documents<lb/>
- ./header1002.tei:11: <note type="other">(for<lb/> the ALICE Collaboration)<lb/></note>
- ./header1015.tei:13: <note type="other"> for the ALICE Collaboration<lb/></note>
- ./55001560.header.tei.xml:33: <keyword>Keywords: Computer-Supported Cooperative Work, Collaboration
- ./header1020.tei:11: <note type="other">(for the ALICE Collaboration)<lb/></note>
- ./header355.tei:11: <titlePart type="main">Search for Chargino and Neutralino <1b/>Production Using
- ./header1006.tei:14: <note type="other">for the ALICE Collaboration<lb/></note>

These should be

This is from a search of 2749 CORA articles... from a set of only 36 HEP conference papers, we get at least as many again.

6.14.4 Training Data

Obtaining workable training data is a notoriously time-consuming task, given that a single data sample consists of a partially of fully annotated document. This challenge is often referred to with dismay in the literature [CITATION HERE]. Evidently, constructing a training set is an inherently manual task, involving the diligence and discretion of a human labourer to. Attempts to automate this task were found to be futile. This of course stands to reason, given that automating training set creation is a problem precisely equivalent to that we wish to solve, namely, automatic metadata extraction. Though there are some publishers that provide complete annotations of their publications, they typically impose a rigid format on the aricles. Building a training from highly select documents in such a way can only result in a homogeneous training set that is not representative of a real world corpus and can only underperform at prediction time. Thus, the researcher is effectively resigned to spend endless hours on manual training data creation. With Grobid, however, there is a shortcut; Grobid provides utilities for "creating" training

data, that is to say, for running an existing model on new articles and producing a "first pass" attempt at extracting the ground truth metadata. It is then the job of the researcher to correct and validate this output to ensure the gold standard. Thus, Grobid does most of the hard work in writing to an XML in the TEI format. This notwithstanding, the validation process takes an enormous amount of time.

As one would expect, the model that is most worth improving–segmentation–is the least effective at assisting in training data creation, and its mistakes are plentiful. Nevertheless, Grobid provides us with a good starting point by producing "predictions" for our sampled HEP papers. Though the shipped model is produced from as few as 120 articles, it is quite accurate at identifying the front matter (header) prior to the start of the body, and the reference section(s). The body matter is usually correctly classified (giving a high true positive (TP) rate), since with a limited training set, predicting the most frequent class can still give a relatively high likelihood. On the other hand, the number of false positives (FP) is significant, with footnotes, headnotes and page numbers regularly overlooked and absorbed into the body. Figure ?? shows this effect. Though in this project we are not interested in classifying page numbers etc., we are mindful of the potential use of our data to future research with Grobid, and we dutifully correct all mistakes we encounter.

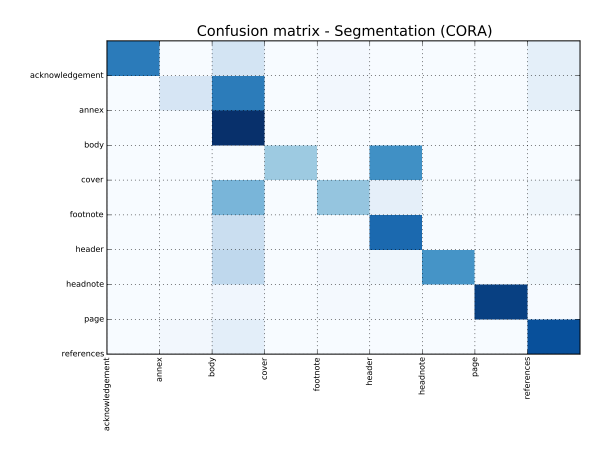


Figure 10: Heat map visualisation of the confusion matrix for the shipped segmentation model. The main diagonal shows the majority correctness of the model, Notably, classification for <body> gives a high true positive rate (TP) (observe the contrast in the <body> row), but a high number of false positives (observe the confusion in the <body> column), i.e. the false negatives of other classes.

The header model has its own problems. Despite being much more accurate (the developers have evidently spent far more time on this model—the shipped training set numbers over 2700 annotated articles, 20 times

as many as that of segmentation) [THIS COULD BE A FOOTNOTE].

7 Conclusions

Failing to admit the futility of attempting to automate training data creation, as well as the subsequent effort required to do it manually, was an enormous hindrance to the *déroulement* of the project.

[REFER TO GOALS HERE]

8 References

9 Appendices

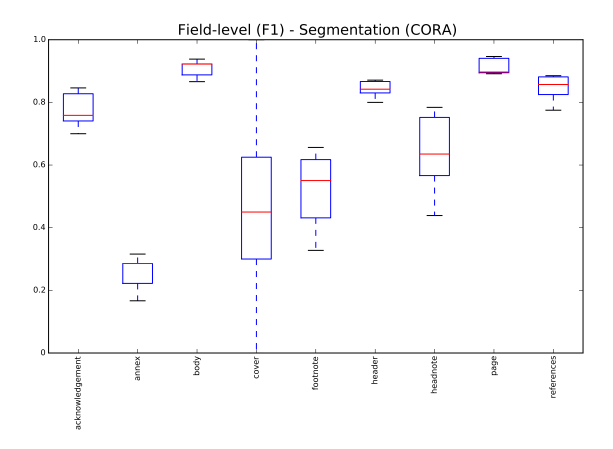


Figure 11: Boxplots of field level F1 scores obtained from 5-fold cross validation of segmentation model on CORA papers.

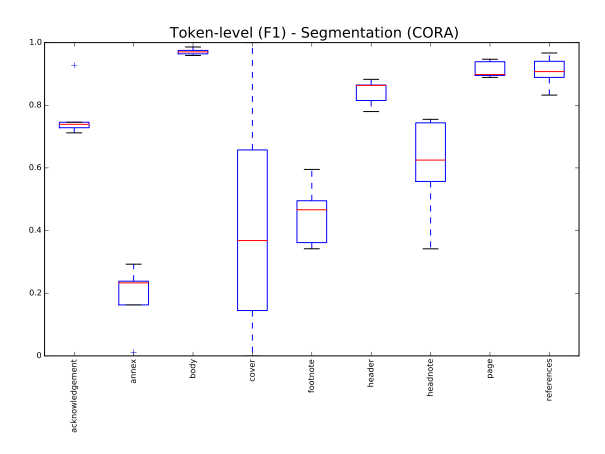


Figure 12: Boxplots of token level F1 scores obtained from 5-fold cross validation of segmentation model on CORA papers.