MSC ARTIFICIAL INTELLIGENCE MASTER THESIS

Enriching Textual Data with Document Structure For Text Classification

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July 12, 2018

42 EC 01 July 2017, 31 December 2017

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Abstract

Proceedings of parliamentary debates are often published as unstructured PDF files, making them unsuitable for indexing into a database or querying for specific information. Digitizing them into a structured format is challenging; doing so manually is labor-intensive, doing so through a rule-based system is error-prone. Manually annotating a small number of documents can provide a training set that allows a convolutional neural network to accurately classify the elements that denote the structure of the document (e.g. the start of a new speech), using purely the textual content of the document. Adding a preprocessing step that clusters pieces of text based on the physical layout of the document improves the classification performance in a minor, but statistically significant way.

1 Introduction

A healthy democracy relies on a transparent political process that is open to the common populace

hier valt vast iets leuks te quoten van een politiek filosoof

. A common step towards achieving this is by publishing the proceedings of the parliament's debates, such as the Bundestag in $Germany^1$ or the $Tweede\ Kamer$ in the Netherlands². These proceedings can be useful in various ways, for example:

- Double-checking whether a certain politician's actions in the parliament are consistent with their public stance.
- Using them as a source of data for text analysis, such as classifying political ideology[1].
- Tracking the change in ideological leaning of a politician or party over time[2].

In each of these cases it would be hugely beneficial if the data was properly indexed. If you want to know John Doe's stance on immigration, you would ideally simply query a database for speeches by John Doe regarding the topic of immigration without having to manually skim over hundreds of documents to find the relevant speeches. It is unfortunate then that many of these debates are published solely in unstructured formats, such as PDF or plain text. Projects such as Political Mashup³ handle this by writing systems to parse and then index these documents. The semantic information required for indexing is currently recovered using rule-based methods. In

¹https://www.bundestag.de/protokolle

²https://www.tweedekamer.nl/kamerstukken

³http://search.politicalmashup.nl/about.html



Figure 1 – The structure of a parliamentary debate is that of a shallow tree.

the case of a PDF document, this is fairly challenging. The data often gets transcribed by a human typist, compiled to a PDF, and then goes back into a PDF decompiler for easier processing; this adds a lot of places where minor variations can occur in the output even though the document itself uses a consistent layout. Dealing with this in a rule-based system entails using either highly general rules that lead to a large probability of false positives, or a large amount of highly specific rules which can quickly lead to a spaghetti-like mess of special cases and is very fragile to unseen cases.

I propose that by using a small number of manually annotated documents as a dataset, a machine learning algorithm can learn to classify fragments of text in a way that allows it to segment a document into its constituent parts, while being more robust to noise than its rule-based counterpart. While the structure of a parliamentary debate is tree-shaped (a general example in shown in fig. 1), this tree is generally both shallow and very rigidly structured. As a result, simply classifying the positions in the text where a new structural element begins is enough to reconstruct the full tree given the domain knowledge about this particular set of documents; some variations can occur depending on the particular conventions used by the creator of the documents, for example in how speech interruptions by noisy colleagues are handled. After the document's layout has been extracted, it is a matter of obtaining each element's metadata (such as the speaker and their political affiliation for each speech element). This step is outside of the scope of this thesis.

The common ways to do sentence classification (e.g. convolutional neural networks [3], recurrent neural networks or the simpler bag-of-words models) operate on sentences in a vacuum, considering only their linguistic contents

and ignoring any contextual information that might be contained in the layout in which the text might have been embedded. This is to be expected considering that most of the common datasets in this area really *are* just small bits of text in a vacuum; often-used datasets involve Twitter messages or short product reviews. In this case however, the sentences come from a document with a rich structure providing a lot of context. Anecdotally, as a human it is trivial to discern section headers in a document even when the document is in a foreign language; simply the fact that the section header might be printed in bold and centered rather than left-aligned gives it away. Incorporating this structural data into the learning process, using a clustering approach inspired by Klampfl *et al.* [4], will hopefully increase the performance of the system, either by simply scoring better on the used metrics, or perhaps more indirectly by requiring less data or training time to achieve the same score.

2 Problem Statement

The German parliament, called the Bundestag, publishes the proceedings of their meetings, as an effort to open up the political process to the common people. These proceedings have been continuously published since the inception of the German Empire in 1871, when the parliament was called the Reichstag, up to 1942; publishing resumed after the conclusion of World War 2 with the inception of the Bundestag in 1949. Of course, the further back in the time you go, the more difficult the documents become to use. While the more recent documents are fully digital, documents prior to 1998 are scans of physical documents and require optical character recognition, which becomes even more problematic in the documents from the 1800s where a thick Gothic font is used. Figure 2 shows a sample page from one of these proceedings; the left column contains a continuation of a speech from the previous page as well as two moderately sized speeches, while the right column contains a large number of very short speeches. Figure 3 shows the same page seen in fig. 2, but with a number of regions of interest (manually) colored in. Unfortunately this data is only available as PDF files, which in terms of internal representation are entirely unstructured. That means that none of the rich structure highlighted in fig. 3 is actually present in a computer-usable way.

The central problem in this thesis is the extraction of speeches from the documents, transforming each document into a structured series of speeches that can be serve as a useful entry point into further research. As a first step, the structural layout (as in fig. 3) will be extracted using unsupervised clustering algorithms. The textual contents of the document are then fed into supervised text classifier, where each piece of text is augmented with the corresponding layout information previously obtained. More details on

Präsident Dr. Norbert Lammert

(A) Ich darf bereits jetzt darauf aufmerksam machen, dass ich nach Schließen des Wahlgangs die Sitzung für die Auszählung der Stimmen unterbrechen werde. Stellen Sie sich bitte darauf ein, dass das etwa eine Stunde dauern kann, weil ja ein doch relativ komplexer Wahlgang ausgezählt werden muss.

Ich eröffne die Wahl

Liebe Kolleginnen und Kollegen, darf ich fragen, ob jemand im Saal ist, der seine Stimme noch nicht abgegeben hat? Oder hat jemand einen gesehen, den er dann nicht mehr gesehen hat und der seine Stimme noch abgeben könnte? – Dann schließe ich diesen Wahlgang und unterbreche die Sitzung bis zur Bekanntgabe des Ergebnisses der Wahl. Wir werden den Wiederbeginn der Sitzung rechtzeitig durch entsprechende akustische und optische Signale in den Immobilien des Bundestages ankündigen. Stellen Sie sich bitte darauf ein, dass es etwa eine Stunde dauern kann, bis wir diesen ja doch umfangreichen Wahlgang mit der gebotenen Sorgfalt ausgezählt haben.

Die Sitzung ist unterbrochen.

(Unterbrechung von 13.42 bis 14.52 Uhr)

Präsident Dr. Norbert Lammert:

Die unterbrochene Sitzung ist wieder eröffnet

(B) Liebe Kolleginnen und Kollegen, ich kann Ihnen das Ergebnis der Wahl der Stellvertreterinnen und Stellvertreter des Präsidenten bekannt geben: abgegebene Stimmkarten 626. Alle abgegebenen Stimmen waren gültig.

Von den abgegebenen Stimmen sind entfallen auf Peter Hintze 449 Jastimmen, 122 Neinstimmen und 51 Enthaltungen. In diesem Falle, was mich ein bisschen überrascht, waren 4 Stimmen ungültig. Das heißt, es gibt keine Stimmkarte, die insgesamt ungültig war, was ja doch auf eine gewisse Pfiffigkeit der neuen wie der alten Kollegen schließen lässt, aber bei einzelnen Wahlgängen ist das offenkundig anders. Noch einmal: 449 Jastimmen, 122 Neinstimmen, 51 Enthaltungen. Ich darf das mit Ihrem Einverständnis gleich mit der Frage an die jeweiligen Kolleginnen und Kollegen verbinden, ob sie die Wahl annehmen. Ich darf den Kollegen Hintze, der damit die notwendige Mehrheit erkennbar erreicht hat, fragen, ob er die Wahl annimmt.

Peter Hintze (CDU/CSU):

Ich bedanke mich. Ich nehme die Wahl an.

(Beifall bei der CDU/CSU sowie bei Abgeordneten der SPD und des BÜNDNISSES 90/DIE GRÜNEN)

Auf den Kollegen Johannes Singhammer sind bei 6 ungültigen Stimmen 442 Jastimmen, 115 Neinstimmen und 63 Enthaltungen entfallen. Auch er hat damit die notwendige Mehrheit eindeutig und klar erreicht. Ich darf ihn fragen, ob er die Wahl annimmt.

Johannes Singhammer (CDU/CSU):

Ich danke für den Vertrauensvorschuss und nehme die Wahl gerne an.

(Beifall im ganzen Hause)

Präsident Dr. Norbert Lammert:

Die Kollegin Edelgard Bulmahn hat bei wiederum 6 ungültigen Stimmen 534 Jastimmen erhalten.

(Beifall im ganzen Hause)

50 Kolleginnen und Kollegen haben mit Nein gestimmt, 36 haben sich der Stimme enthalten. Frau Bulmahn, ich darf auch Sie fragen, ob Sie die Wahl annehmen.

Edelgard Bulmahn (SPD):

Auch ich bedanke mich für das Vertrauen, und ich nehme die Wahl gerne an.

(Beifall im ganzen Hause)

Präsident Dr. Norbert Lammert:

Auf die vorgeschlagene Kandidatin Ulla Schmidt sind 520 Jastimmen entfallen.

(Beifall im ganzen Hause)

66 Kollegen oder Kolleginnen haben mit Nein gestimmt, 35 haben sich der Stimme enthalten. 5 Stimmen waren ungültig. Ich bin zuversichtlich, Frau Schmidt, dass Sie die Frage ähnlich beantworten wie die bisher angesprochenen Kolleginnen und Kollegen.

(D)

Ulla Schmidt (Aachen) (SPD):

Herr Präsident, Sie haben wie meistens recht. Ich nehme die Wahl an und bedanke mich für das große Vertrauen. Danke schön!

(Beifall im ganzen Hause)

Präsident Dr. Norbert Lammert:

Auf Petra Pau sind 451 Jastimmen entfallen,

(Beifall im ganzen Hause)

bei 113 Neinstimmen und 45 Enthaltungen. 17 Stimmen waren in diesem Wahlvorgang ungültig. Ich darf Frau Pau fragen, ob sie die Wahl annimmt.

Petra Pau (DIE LINKE):

Ja, Herr Präsident, ich nehme die Wahl gern an, und, liebe Kolleginnen und Kollegen, ich freue mich auf die weitere Zusammenarbeit.

(Beifall im ganzen Hause)

Präsident Dr. Norbert Lammert:

Schließlich darf ich noch das Wahlergebnis für Claudia Roth bekannt geben. Bei 14 ungültigen Stimmen hat sie 415 Jastimmen erhalten. Es gab 128 Neinstimmen und 69 Enthaltungen. Sie ist damit gewählt.

(Beifall im ganzen Hause – Claudia Roth [Augsburg] [BÜNDNIS 90/DIE GRÜNEN]:

Figure 2 – A sample page from one of the Bundestag proceedings.



Figure 3 – The same page as in Figure 2, hand-annotated with interesting regions that play a significant role in understanding the layout. The red regions are the headers the signify that someone is starting a speech, and contain information about who the speaker is. The blue regions are little interruptions (*Heiterkeits*), often signifying approval or displeasure (the regularly seen *Beifall* means applause). The green regions indicate plain blocks of text.

this process will be supplied in section 4.

As annotating data by hand is an expensive process, a big focus is on limiting the amount of required training data as much as possible; it would be preferable if the system was able to learn sufficiently from a handful (say, less than 5) of hand-annotated files.

2.1 Dataset

The dataset contains 11 hand-annotated documents, comprising 2052 positive samples and 76,944 negative samples. The average document contains about 200 positive samples. Seeing as the source documents are PDF files, some transformations are needed to extract text from them in a way that is suitable for use as a machine learning dataset. This is not as trivial as it might seem, given that PDF files only contain instructions for drawing certain characters at certain coordinates, with no internal concept of paragraphs, lines or even words.

For the supervised portion of the system, the PDF files are transformed using the pdftohtml utility from the Poppler PDF rendering library⁴. This utility takes a PDF file and uses a number of heuristics to output lines of text occurring inside the file. In this context, a line refers to any number of characters that occur on the same height on a page while preserving reading order (which is relevant when dealing with documents that have a two-column layout) and is explicitly not the same as a sentence. These lines are output in an XML format which includes metadata on the geometry of the line (position and size) and its font. An example of a portion of text from the dataset and the corresponding XML output from pdftohtml is shown in fig. 4. Since this process is based on heuristics, it can and does go wrong; there are instances of mistakes such as a single line being broken up into multiple XML nodes, or lines of two side-by-side columns being taken as a single XML node. This is not terribly common (the frequency of such errors is perhaps one per file on average), but it does make the dataset inherently noisy and is one of the issues that rule-based systems have trouble with.

For the unsupervised clustering, the PDF files are taken directly as input and clustering is done using individual characters as the basic unit. This is just as easy as clustering on the lines produced by pdftohtml, but has the benefit of not being dependent on the imperfect line-extraction heuristics.

⁴https://poppler.freedesktop.org/

Dr. Norbert Lammert (CDU/CSU):

Herr Alterspräsident, lieber Kollege Riesenhuber, ich nehme die Wahl geme an.

(Beifall im ganzen Hause – Abgeordnete aller Fraktionen gratulieren dem Präsidenten)

(a) A portion of the source PDF.

```
<text top="122" left="125" width="143" height="16" font="3">
    Dr. Norbert Lammert
</text>
<text top="122" left="269" width="83" height="17" font="4">
    (CDU/CSU):
</text>
<text top="142" left="125" width="328" height="17" font="4">
    Herr Alterspräsident, lieber Kollege Riesenhuber, ich
<text top="158" left="108" width="156" height="17" font="4">
    nehme die Wahl gerne an.
<text top="186" left="141" width="278" height="17" font="4">
    (Beifall im ganzen Hause Abgeordnete aller
</text>
<text top="203" left="158" width="242" height="17" font="4">
    Fraktionen gratulieren dem Präsidenten)
</\text{text}>
```

(b) XML created by running pdftohtml, corresponding to the PDF excerpt in Figure 4a. The contents of the text elements are used as inputs for the classification algorithm; the layout data contained in the properties is not used, as a separate software pipeline is used for the unsupervised clustering.

Figure 4 – A sample excerpt from a source PDF, along with its XML representation created by pdftohtml.

2.2 Research Question

A baseline model can be created by leaving out the clustering step, leaving us with two models:

- 1. A baseline model that classifies based on purely text
- 2. A model that classifies based on both text and layout information

There are two ways in which the second model can improve upon the baseline: either it performs better (using a metric such as the F1 score), or it requires less data to reach the same performance. This naturally leads to two research questions:

Research Question 1 Does augmenting a text classification system with layout information obtained by unsupervised clustering of the input data improve the F1 score of the classifier?

Research Question 2 Does augmenting a text classification system with layout information obtained by unsupervised clustering of the input data allow the classifier to reach its peak performance using less input data?

3 Related Work

The task handled in this thesis is in a way similar to that of wrapper induction, which is the process of inferring a wrapper (a program that extracts data into a usable form) from a web page. A fairly recent survey of the state of this field is done by Ferrara et al. [5], who note that a big problem is keeping up with the constantly changing nature of web pages. A novel approach to combat this is that of Gogar et al. [6]. They do wrapper induction by combining the textual content of a webpage with a screenshot of the rendered webpage in an effort to do wrapper induction on previously unseen web pages. The text is encoded in a way that maintains spatial information, a model they refer to as Text Maps or Spatial bag of words. The text maps and the screenshot are fed into separate convolutional networks, after which the output is combined for a final classification. In a test of extracting product names and prices from web pages, the system obtains very high score comparable to systems that do use site-specific initialization.

In terms of analyzing document structure, Klampfl et al. [4] introduce a method to analyze scientific articles, detecting blocks of text, labeling them (as e.g. section headers, tables or references) and determining the reading order — all in an unsupervised manner. The text block detection is done using a sequence of different clustering algorithms, while the labeling is done using a heuristic approach.

For text classification, various forms of convolutional neural networks are commonly used. The most basic architecture is described by Kim [3], where

the input is tokenized and the tokens are embedded into a higher-dimensional representation before passing them to the convolutional neural network. Using this same architecture, Zhang & Wallace [7] perform additional exploration of the parameters and their effects on various datasets. Comparable results are achieved by Zhang et al. [8] by operating on the character-level rather than the word-level, bypassing the overhead of using word embedding (either in extra training time or in finding suitable pretrained embeddings) as well as freeing the researcher from having another layer in their network to tune. All the previously mentioned architectures use a single convolutional layer; this is contrary to current trends in computer vision, where popular models such as ResNet[9] go as deep as 152 layers. This difference is explored by Conneau et al. [10], who take a character-level CNN and show that adding more layers improves performance, before leveling out at 29 layers. They hypothesize that the difference in effective depth between computer vision and language processing might be due to the difference in datasets. The common ImageNet dataset used in computer vision deals with 1000 classes; in contrast, sentiment analysis datasets vary between 2 and 25 classes. In addition, they note that the deeper networks do require a larger amount of data to train.

4 Methodology

As described in section 2, the input data comes in the form of a PDF file which can be converted into plaintext XML data containing lines of text along with each line's bounding box. There are two systems in play here; the main system is a convolutional neural network classifier that acts on the XML data. Further details on this are provided in section 4.2. The second system is the unsupervised preprocessor which acts on the PDF data and writes its output as an additional property into the XML data, which is elaborated upon in section 4.1. Figure 5 shows a high-level overview of the two systems and how they interact.

4.1 Unsupervised

The unsupervised algorithm is inteded to detect and classify blocks of text in the PDF file; Figure 6 shows an example. This approach is based on work by Klampfl et al. [4], and consists of two separate clustering steps. First, individual characters (the fundamental objects available in a PDF file) are clustered together into blocks of semantically relevant text. These could be, for example, paragraphs, section headers or page decoration. By using the bounding boxes of the blocks, they can be clustered based on their shape and some additional metadata (e.g. occurrence of font types and sizes). The rest of this section will go into details on the two clustering steps.

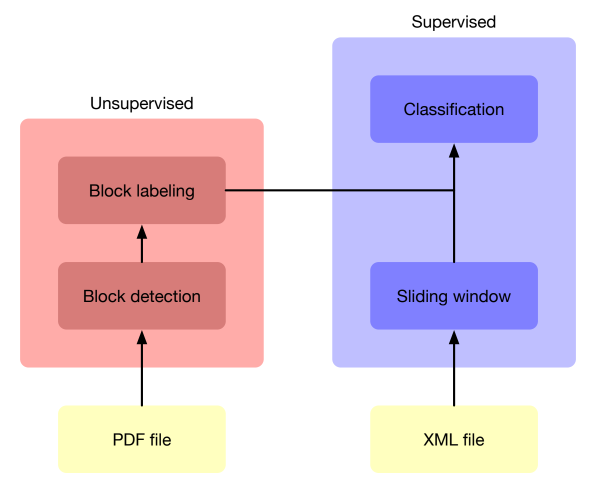


Figure 5 - A high-level overview of the system. The unsupervised block augments the input to the classifier.

4.1.1 Hierarchical Agglomerative Clustering

The first step is performed using hierarchical agglomerative clustering (HAC), an unsupervised bottom-up clustering algorithm that constructs a hierarchical tree of clusters (in this context referred to as a *dendrogram*). An example is shown in fig. 7. The algorithm gets fed the individual characters present in the PDF files, then iteratively groups the two closest clusters (the initial inputs being regarded as clusters of one element) together until only a single cluster remains. This process involves two parameters:

- 1. The distance function between two characters.
- 2. The distance function between two clusters of characters.

The first parameter is trivially chosen to be the Euclidian distance between the coordinates of the two characters. The second parameter is called the



Figure 6 – An example of clustered blocks of text. Blocks with the same outline color belong to the same cluster.

linkage and has several common options, the most basic of which are:

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• Single-linkage: The distance between clusters is based on the closest two elements:

$$d(A,B) = \min\{d(a,b) : a \in A, b \in B\}$$

• Maximum-linkage: The distance between clusters is based on the furthest two elements:

$$d(A, B) = \max\{d(a, b) : a \in A, b \in B\}$$

• Average-linkage: The distance between clusters is based on the average distance of its elements:

$$d(A,B) = \frac{1}{|A||B|} \sum_{a \in A} \sum_{b \in B} d(a,b)$$

Additionally there are more involved linkage criteria, such as the Ward method which minimizes the variance within clusters. Although these more

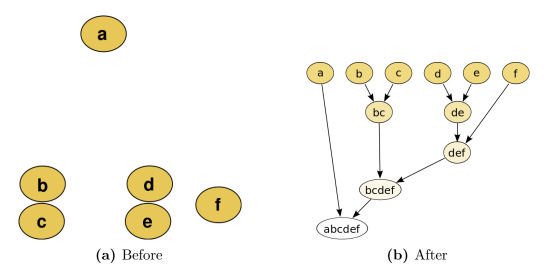


Figure 7 – An example of hierarcial agglomerative clustering, where the nodes are clustered by distance.

complex methods would generally be favored, in this specific context single-linkage clusters is actually the best option[4]. This is due to its tendency to form long, thin clusters; this mirrors the nature of text, in particular words and sentences (which are just long, thin strings of letters and words respectively). As an additional bonus, it is the most computationally efficient method. While the general time complexity for HAC is in $\mathcal{O}(n^3)$, clever algorithms exist for single-linkage clustering that fall in $\mathcal{O}(n^2)$ [11], making it far more usable on larger datasets.

After the dendrogram is constructed, it has to be cut at some level to obtain the desired blocks of text. Clustering can optionally be rerun using the newly found clusters as basic elements. This way, the document can incrementally be clustered from characters into words, words into lines, and finally lines into paragraphs. Both the level at which to cut the tree and the number of times to recluster are to be manually tuned based on the particular set of documents.

4.1.2 Clustering

The extracted blocks from the previous step are then clustered according to their similarity based on the following metrics:

- Width of the cluster
- Height of the cluster
- The ID of the most common font occurring in the cluster
- The size of the most common font occurring in the cluster

This is implemented two different ways, with their performance to be compared later on. The first method is a simple K-Means clustering, which works as follows:

- 1. Randomly initialize k cluster centroids.
- 2. Assign each observation to its nearest centroid.
- Recompute each centroid to be the mean of all of its assigned observations.
- 4. Repeat steps 2 and 3 until the assignments stop changing.

After the algorithm has converged, the clustering is defined by each observation's nearest cluster centroid; with k clusters, each observation is assigned a k-dimensional one-hot vector.

A more complex generalization of this is a mixture model. While K-Means has a hard boundary between clusters, in a mixture model each cluster is a probability distribution over the full space of observations. As a result, rather than ending up with a one-hot vector, with a mixture of k distributions each observation is assigned a k-dimensional vector whose ith element gives the probability of belonging to the ith cluster. Of course, this added information comes at the price of a far more involved algorithm required to compute the distributions.

4.2 Supervised

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After the data is augmented by the previously described clustering algorithms, it's fed into a convolutional neural network for classification. Since the documents have a dual column layout (meaning each line of text is pretty short) and classification is based on the lines from the document, a sliding window is used to supply more context. The text content of this window is then tokenized (such that words as well as each individual punctuation mark are a token) and fed into a standard convolutional neural network architecture [3], with the label (i.e. does or does not start a new speech) being provided by the center element of the window. First an embedding layer is used to learn a high-dimensional representation of the tokens (no pre-trained embeddings were used because of both the specialized political domain of the data as well as a lack of quality pre-trained German models), followed by a single convolutional layer, consisting of multiple filters each followed by 1-max pooling (a more in-depth explanation is provided in section 4.2.3). The output of the filters is then concatenated into a single feature vector, which is combined with the clustering data in one of two ways, which will both be evaluated:

1. The cluster types of each element in the window are added to the feature vector.

2. An additional smaller LSTM is applied to the cluster types of each element in the window, the output of which is concatenated with the feature vector.

The resulting feature vector is then fed into a fully-connected neural network. This network features one hidden layer with a ReLU activation ($\operatorname{relu}(x) = \max(0, x)$) and a single output node with a sigmoid activation. The layout of this system is detailed in fig. 8 along with its parameters and their baseline value.

The network is trained for 100 epochs, stopping early once no significant improvement has been made on a small validation set for 10 epochs in a row. The optimisation process is done using the Adadelta[12] algorithm⁵, with binary cross-entropy as the loss function and the training data delivered in batches. Regularisation is done through a combination of dropout[14] and an upper limit on the L2 norm of each weight vector[15]. Further explanations of the optimisation process and the regularisation methods are provided in section 4.2.1 and section 4.2.2 respectively.

4.2.1 Optimisation process

Optimisation is done using the Adadelta update rule[12]. This is easiest to explain by starting with the basic mini-batch stochastic gradient descent algorithm (SGD). At each iteration t, the network parameters θ are updated based on some calculation:

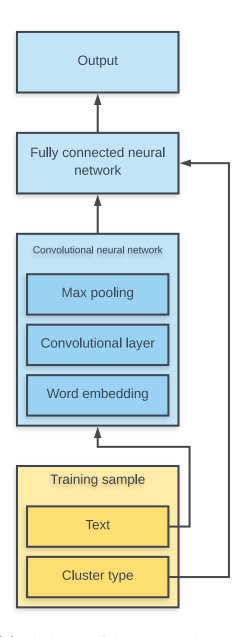
$$\theta_{t+1} = \theta_t - \Delta\theta_t \tag{1}$$

The difference between optimization algorithms is how $\Delta\theta$ is calculated. With SGD, it is simply

$$\Delta \theta_t = \mu \nabla_{\theta_t} \mathcal{L}(\theta_t) \tag{2}$$

where \mathcal{L} is the loss function (in this case, the binary cross entropy between the predicted labels and the true labels), and μ is an arbitrary learning rate between 0 and 1. This learning rate is a tricky parameter to set; too low and learning will take ages, too high and the network will fail to converge because the steps taken are too big. One way to improve this is by using the Adagrad[16] algorithm. While SGD uses a single learning rate for the entire parameter vector, Adagrad (which stands for "adaptive gradient") adapts the learning rate for each individual parameter; frequently updated parameters get a lower learning rate, while less frequently updated parameters

⁵Why not Adam[13]? The architecture[3] and survey paper[7] that this work is based on both date to roughly the year when the Adam paper was published, and thus likely predate the widespread acceptance of Adam as a de facto standard. Since a full analysis of different optimizers is outside the scope of this thesis and some quick informal tests showed no difference between using Adadelta or Adam, there is no reason to deviate from the survey.



Parameter	Default value
Window size	9
Word embedding size	300
Number of filters	100
Filter sizes	4, 5, 6
Activation	ReLU
Pooling type	1-max

(b) The default parameters used in the convolutional neural network.

Parameter	Default value		
Number of hidden layers	1		
Hidden layer size	50		
Hidden layer activation	ReLU		
Number of outputs	1		
Output activation	Sigmoid		

(c) The default parameters used in the fully-connected neural network.

Parameter	Default value
Batch size	50
Dropout rate	0.5
Learning rate	1.0
ρ	0.9
Maximum L2 norm	3

(d) The default parameters used for the optimisation process.

Figure 8 – The mod \emptyset and its parameters.

are updated with a higher learning rate. This is done by simply dividing the learning rate with the L2 norm of the sum of all previous gradients:

$$g_t = \nabla_{\theta_t} \mathcal{L}(\theta_t) \tag{3}$$

$$\Delta\theta_t = \frac{\mu}{\sqrt{\sum_{T=1}^t g_T^2}} g_t \tag{4}$$

This has been found to work very well, in particular in natural language processing and computer vision where features are often sparse, but it has two drawbacks:

- 1. A suitable value for the global learning rate μ has to be manually provided.
- 2. Since the learning rate is rescaled using a monotonically increasing sum of previous gradient magnitudes, the learning rate will converge to zero.

Adadelta nullifies these drawbacks by eliminating the global learning rate and restricting the accumulation of gradients to a window of recent updates. First of all, the sum over all previous gradients is replaced by an exponentially decaying average of the squared gradients, referred to as $E[g^2]$:

$$E[g^2]_t = \rho E[g^2]_{t-1} + (1 - \rho)g_t^2$$
(5)

Here ρ is a constant representing the rate of decay; this decaying sum serves as a more efficient approximation of an actual window of past gradients, which would require far more memory to store (considering both the huge number of parameters in a neural network and the memory constraints of running on a GPU). Substituting this into the Adagrad algorithm gives us:

$$\Delta\theta_t = \frac{\mu}{\sqrt{E[g^2]_t}} g_t \tag{6}$$

$$= \frac{\mu}{RMS[g]_t} g_t \tag{7}$$

The quantity $\sqrt{E[g^2]_t}$ is called the *root mean square* of g, which occurs often enough in optimisation algorithms that it is often abbreviated as RMS[g]. As a final step, the learning rate μ is eliminated by replacing it with a decaying average of the previous gradient updates, similar to eq. (5).

$$\Delta \theta_t = \frac{RMS[\theta]_{t-1}}{RMS[g]_t} g_t \tag{8}$$

4.2.2 Regularisation

Dropout[14] is a Regularisation method that is rather crude at first glance; on each batch update, each node in the neural network has a probability

p of outputting zero (i.e. the node being "disabled"). As a result, nodes cannot rely on the output of any other node being present, preventing coadaptation and as a result reducing the probability of overfitting on the training data. Dropout is only active during training; when running the network in evaluation mode, all nodes are active and the outputs are rescaled by a factor of 1-p to account for the now higher activation values. There is another way to view dropout. It is commonly known that neural network (or really any machine learning classifier) performance can be improved by training a large number of them and averaging their outputs. Since every combination of nodes disabled by dropout could be considered a unique neural network, dropout acts as computationally cheap approximation to averaging multiple networks.

In addition to dropout, a maximum L2 norm is imposed on each node's incoming weight vector. If the weight vector's L2 norm exceeds this limit, it is rescaled so that its new norm is equal to the maximum allowed norm; otherwise the weight vector is left alone:

$$\mathbf{w} = \begin{cases} \mathbf{w} & \text{if } ||\mathbf{w}||^2 \le n \\ n \frac{\mathbf{w}}{||\mathbf{w}||^2} & \text{otherwise} \end{cases}$$
 (9)

This differs from the more common L2-regularisation — where the combined L2 norm of all weight vectors is added to the loss function — in that the weights are not being continuously pushed towards zero, making it a milder form of regularisation that allows for a bit more complexity in the model.

4.2.3 Convolutional Neural Networks

In addition to explaining how a convolutional neural network works, this section will additionally go into why the choice was made to use a convolution neural network, rather than a recurrent neural network (which is more strongly associated with text processing) or any other machine learning algorithm. Starting with the basics, text is inherently troublesome because it produces 3-dimensional data: there is a feature vector for each word, and some (either variable or predetermined through padding) amount of words per text. This makes each training sample a 2-dimensional matrix, which then gets stacked in the depth dimension to produce 3-dimensional training data. This clashes with the tendency of usual machine learning algorithms to work on 2-dimensional data, assuming a feature vector for each sample rather than a matrix. There are three common methods to deal with this:

• Bag of words

- Convolutional neural networks
- Recurrent neural networks

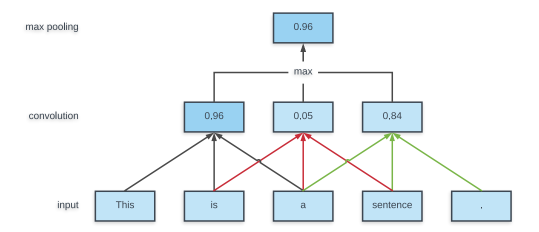


Figure 9 – A simplified convolutional neural network with one filter and max pooling.

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Aside from being completely different methods, they differ in a major way in how they handle the sequential nature of text (i.e. the words in a sentence have a meaningful order). The bag of words approach is the simplest in that it simply disregards this sequential nature entirely, instead creating what is essentially a histogram of word occurrences. This downsamples each sample from a feature matrix to a feature vector, allowing the use of conventional machine learning algorithms (most commonly support vector machines). While the sequential information can be kept to some degree by using histograms of n-grams rather than words (unigrams), this causes the size of the input data to scale exponentially with the value of n.

Convolutional neural networks (CNNs) work by taking a number of filters (sometimes called kernels or feature maps) of a specified size and convolving these over the input data. A simplified example using one filter is shown in fig. 9. In this example, the input text is convolved with a filter with a width of 3 and a stride of 1 — that is, each application of the filter considers three subsequent input elements, after which the window is shifted one space to the right. This filter is essentially a small neural network mapping three items to one output value, whose weights are reused for each application of the filter. Reusing the weights in this way (weight sharing) prevents the number of parameters in the network from spiraling out of control. [17] After the application of this convolution layer, the responses of the filter form a new sequence of roughly the same size as the input (minus a few due to missing the edges). The next step is to downsample this sequence by means of a max pooling layer, which maps all values within a window to the maximum value amongst those values. While conceptually similar to a convolution, this step generally does not involve overlap, instead either setting the stride to the same value as the window size (usually 2) or reducing the entire sequence to 1 value (1-max pooling). The reason for this is twofold:

- 1. It downsamples the data, reducing the amount of parameters required further on in the network.
- 2. It adds translation invariance to the feature detected by this filter. The example filter of fig. 9 reacts strongly to the first three words. Without the pooling layer, changing the location of these words in the input would similarly change the location of the high activation in the intermediate representation; this would be translation equivariance. The more aggressively the pooling is applied, the higher the degree of invariance (with full translational invariance being achieved with 1-max pooling).

This combination of convolution followed by pooling can be repeated multiple times as desired or until there is only a single value left as output from the filter. Finally, the outputs of all filters are concatenated and fed into a regular neural network.

While CNN architectures in computer vision are generally very deep, they tend to be very shallow in natural language processing; commonly just a single convolution followed by 1-max pooling [7].

4.2.4 Difference between convolutional and recurrent neural networks

Recurrent neural networks (in particular LSTMs or GRUs) are seemingly the most natural fit for language processing, since they process an entire sequence and are therefore fully conditioned on the word order (as opposed to the convolutional neural networks which tend to learn translation invariant ngram features). In spite of this, the decision to use convolutional neural networks is based on two factors:

- 1. In practise, the performance for classification tasks does not differ between the convolutional and recurrent neural networks.[18]
- 2. The computations in convolutional networks are highly independent of each other, allowing for great parallelization (in particular with regards to running on a GPU). In contrast, LSTMs are bottlenecked by the fact that each calculation is dependent on the previous calculations. As a result, CNNs achieve far higher training speeds.[19]

5 Experimental Setup

Experiments are focused on the difference in performance between the baseline CNN model without clustering information (referred to from here on as CNN) and the model augemented with clustering information (which

will be reffered to as CNN-cluster). Performance will be measured with regards to the following three metrics:

- 1. Number of training epochs until convergence
- 2. The F1 score or average precision metrics on a test set (see Section 5.2)
- 3. The number of training samples required to attain a specific F1/average precision score

In each case, the experiment will be repeated 10 times by means of 10-fold cross validation followed by a Student's T-test to gauge the probability of the following null hypothesis being true:

Null Hypothesis 1 Adding clustering information to the CNN model does not change the performance of the model.

5.1 Dataset

Referring back to Figure ??, the average document has between 100 and 300 positive samples. Since a secondary concern is to minimize the number of documents that would have to be annotated as training data, the tested dataset sizes will be very low, with the number of positive samples being one of 100, 200, 500 and 1000. Due to the relative abundance of negative samples and to prevent overfitting on the distribution of the labels, stratified sampling will be used to keep a 1:1 ratio of positive to negative samples. In addition to the size, the number of cluster types (the k in k-means) will be varied to examine its impact on the performance.

5.2 Testing performance

All models will be tested on a test set containing 1000 positive samples and 10000 negative samples, all of which are guaranteed not to be in the training set. Performance on this set is measured by constructing a precision-recall curve, and calculating two values:

- 1. The average precision (which is equivalent to the area under the curve)
- 2. The F1 score of the point on the curve maximizing the F1 score

5.3 CNN performance

Although less central to the thesis than the difference between the CNN and CNN-cluster models, some experimentation will be done with the parameters of the convolutional network in an attempt to optimize the performance. These parameters include the dimensionality of the word embeddings, the number of filters, the pooling strategy (1-max versus a smaller region) and the number of convolutional layers.

5.4 Generalisation

This particular dataset has the quirk that the performance of a rule-based system created based on recent documents decreases in performance when used on older documents, the older the document the worse it performs. This occurs despite the layout being visually the same all the way back to the 1950s. A number of files from old election periods has been labeled (and manually verified for correctness) in order to test

- 1. whether the CNN models handle this better than the rule-based system does.
- 2. Whether the clustering-augmented CNN model performs better on this task than the baseline CNN.

6 Evaluation

For evaluation, two models are considered:

- "Baseline", a basic convolutional neural network without using the clustering information.
- "K-Means", similar to the baseline model with the addition of clustering information into the final classification step.

These models are tested based on two scenarios that differ in the source of the test set. In the first scenario, training and testing data are both pulled from the same pool of data. In the second scenario, all the training data is pulled from the most recent 18th electoral period, while the testing data comes from a number of files from electoral periods 13, 14, 16 and 17. This makes the problem more difficult in two ways:

- 1. A change of electoral period means at least a partial change in parliamentary members, which reduces the ability of the model to essentially overfit on the names of members.
- 2. Although older documents use the same layout as newer ones, the PDF files were generated using different software (electoral period 13 started in 1994 and was the first period to use truly digital documents older documents are scanned versions of printed paper). This results in the flaky PDF decompiler making different mistakes than it does on the newer documents. This is a huge problem for rule-based systems; hopefully the probabilistic nature of machine learning makes it more robust to this issue.

In the first scenario, the experiments are run 10 times using a *shuffle* split algorithm; this is similar to k-fold cross validation, but without the restriction of having to use k evenly sized splits. Instead, for each iteration

n training samples are randomly sampled while the rest of the samples are used for testing. The second scenario uses the same method for generated a training set, but uses a separate, unchanging test set instead. Additionally, the sampling is stratified, meaning it maintains the ratio (1:1) of positive to negative samples in the dataset.

Before directly comparing the two models, a good baseline value has to be determined for the two parameters in the model that are the most difficult to determine:

- 1. the size of the sliding window applied to the input data
- 2. the number of clusters to detect in the blocks of text (i.e. the "k" in k-means)

Although there are more parameters, they are related to the convolutional neural network; this means they can be set to pretty good values using prior knowledge, as well as the fact that convolutional neural networks are not hugely sensitive to their parameters.

6.1 Sliding Window Size

The effect of the sliding window size is explored in ?????. Both cases show a clear downward trend as the window gets bigger, but peak at a different value. When testing on the full dataset a very small window size of 2 performs best, while testing on older electoral periods seems to require a bit more information and performs optimally at a size of 4. In addition, both testing scenarios show the same behaviour regarding the baseline and K-Means models: the K-Means model generally outperforms the baseline at suboptimal window sizes, but at the optimum their performance appears to be identical.

6.2 Number of Clusters

Results for the number of clusters are given in fig. 11. In this case, the window size was fixed to the previously determined optimum of 5. The Baseline model was not tested due to it not using the clustering data. Although the results seem all over the place with large confidence intervals, this is a bit of an illusion caused by the small range of F1 scores that they span. There is an odd spike at 7 clusters which is hard to explain, but might be caused by the probabilistic nature of the K-Means algorithm and the fact that the same clustering was used for each trial; the particular K-Means run when doing 7 clusters might have produced a below-average result.

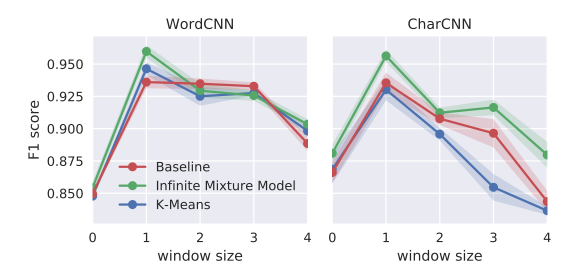


Figure 10 – These figures shows the F1 score with regards to the sliding window size for each model trained on 2000 training samples. The vertical axis shows the F1 score, averaged over 5 trials; the horizontal axis shows the size of the sliding window applied to the training data. The translucent bands around the lines indicates the confidence intervals, meaning that based on the observed F1 scores and assuming normality, the true mean is 95% likely to fall within that interval. The dots on the lines indicate measurements.

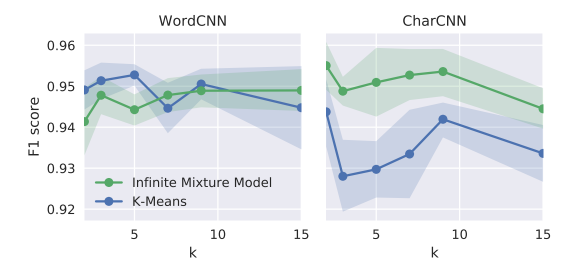


Figure 11 – This figure shows the performance with regards to the number of cluster types for each model trained on 1200 training samples with a window size of 5. The vertical axis shows the F1 score, averaged over 10 trials; the horizontal axis shows the number of cluster types considered in the final clustering step. The translucent bands around the lines indicates the confidence intervals, meaning that based on the observed F1 scores and assuming normality, the true mean is 95% likely to fall within that interval. The dots on the lines indicate measurements.

6.3 Training Set Size

The results of all three models tested with the basic parameters (fig. 8d) on various input sizes is shown in ??. The K-Means model outperforms the baseline at every input size, and as expected more data greatly improves the performance, rising sharply up to 800 samples and rising more gradually from then on. The precise values are given in table 1, along with p-values calculated through a two-tailed Student's T-test.

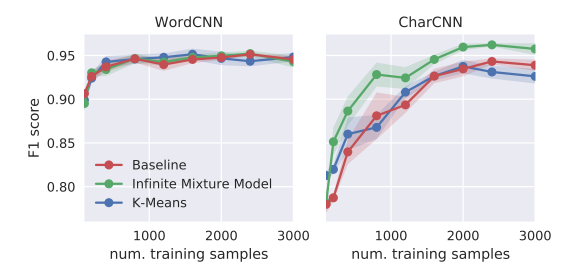


Figure 12 – This figure compares the performance of the baseline model, using only a CNN, to that of the model augmented with K-Means clustering. Tested on a test set of older electoral periods not present in the training set. The vertical axis shows the F1 score, averaged over 10 trials. The horizontal axis shows the number of samples used for training (at a 1:1 ratio of positive to negative samples). The vertical bars indicate confidence intervals, meaning that based on the observed F1 scores and assuming normality, the true mean is 95% likely to fall within the shown interval.

	Baseline		K-Means		
	mean	std	mean	std	
50	0.8263	0.0239	0.8347	0.0267	0.1981
100	0.8606	0.0222	0.8660	0.0235	0.3449
400	0.9268	0.0086	0.9363	0.0075	0.0048
800	0.9496	0.0032	0.9520	0.0042	0.0513
1200	0.9517	0.0038	0.9543	0.0035	0.0471
1600	0.9577	0.0046	0.9599	0.0052	0.0496

Table 1 – The F1 scores of both models compared at different sizes of the training set. The reported means and standard deviations are based on 10 repeated trials on randomized subsets of the data, with the higher mean between the two models bolded for emphasis. The last column indicates the probability that the scores for the K-Means model are drawn from the same distribution as those for the Baseline model (i.e. that their performance is the same and any observed difference is due to chance); probabilities below the usual 0.0.5 cutoff for significance are rendered italic.

7 Conclusion

Todo: conclusion

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