

Machine Learning Engineer Nanodegree

Capstone Project

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Definition

Project Overview

Of late, there has been a plethora of neural models for various NLP tasks. Recently, state-of-the-art text classification models such as Support Vector Machines have yielded to new neural models emerging almost every month. This project looks into just one task Text Classification.

Since the inception of word vectors, they are omnipresent in the NLP community. There has been a major shift from using n-grams (count) based models to word vectors based models. The latter models are able to capture semantic and syntactic information without much preprocessing, which played a part in obtaining competitive performance in the former models. Recurrent Neural Networks and Convolutional Neural Networks are responsible for major breakthroughs in computer vision, since then they have been adopted in NLP tasks and have proven effective in many tasks such as Part of Speech Tagger, Named Entity Recognition, and more. This project aims at providing a survey on most of the models considered for text classification in chronological order. This cannot be considered a comprehensive study on all the neural methods available, but can be used as a reference.

Deep learning in medicine has been applied in a variety of applications such as image-based assessments of traumatic brain injuries, identifying diseases from ordinary radiology image data, visualizing and quantifying heart flow in the body using any MRI machine, as well as analyzing medical images to identify tumors, nearly invisible fractures, and other medical conditions. A lot has been said during the past several years about how precision medicine and, more concretely, genetic testing is going to disrupt the way diseases like cancer are treated. However, this is only partially happening due to the large amount of manual work still required.

Project Statement

As a part of this project, the data used for evaluating all the models are taken from an active competition Personalized Medicine: Redefining Cancer Treatment [1] launched by Kaggle along with Memorial Sloan Kettering Cancer Center (MSKCC). This has been accepted by the NIPS 2017 Competition Track, because they need data scientists to help take personalized medicine to its full potential. Once sequenced, a cancer tumor can have thousands of genetic mutations. But the challenge is distinguishing the mutations that contribute to tumor growth (drivers) from the neutral mutations (passengers). Currently, this interpretation of genetic mutations is being done manually. This is a very time-consuming task whereby a clinical pathologist has to manually review and classify every single genetic mutation based on evidence from text-based clinical literature. For this competition, MSKCC is making available an expert-annotated knowledge base where world-class researchers and oncologists have manually annotated thousands of mutations. One needs to

develop Machine Learning algorithms that, using this knowledge base as a baseline, automatically classifies genetic variations.

Evaluation Metrics

Submissions are evaluated on Multi Class Log Loss between the predicted probability and the observed target. Multi Class Log Loss is the multi-class version of the Logarithmic Loss metric. Each observation is in one class and for each observation, a predicted probability for each class will be submitted. The metric is negative the log likelihood of the model that says each test observation is chosen independently from a distribution that places the submitted probability mass on the corresponding class for each observation.

$$\text{logloss} = -\frac{1}{N} \sum_i^N \sum_j^M y_{i,j} \log(p_{i,j})$$

where N is the number of observations, M is the number of class labels, \log is the natural logarithm, $y_{i,j}$ is 1 if observation i is in class j and 0 otherwise, and $p_{i,j}$ is the predicted probability that observation i is in class j .

Both the solution file and the submission file are CSV's where each row corresponds to one observation and each column corresponds to a class. The solution has 1's and 0's (exactly one "1" in each row), while the submission consists of predicted probabilities.

The submitted probabilities need not sum to 1, because they will be rescaled (each is divided by the sum) so that they do before evaluation. This metric is used to compare kaggle competition participants. But we are going to be using categorical accuracy which can serve as a good measure for this project to compare and contrast all the models used.

Analysis

Data Exploration

Models must be developed to classify genetic mutations based on clinical evidence (text). There are nine different classes on which to classify a genetic mutation. This is not a simple task since interpreting clinical evidence is challenging even for specialists. Therefore, modeling the clinical evidence (text) will be critical to success. Both training and test data sets are provided via two different files. One (training/test_variants) provides the information about the genetic mutations, whereas the other (training/test_text) provides the clinical evidence (text) that our human experts used to classify the genetic mutations. Both are linked via the ID field. Therefore, the genetic mutation (row) with ID=15 in the file training_variants was classified using the clinical evidence (text) from the row with ID=15 in the file training_text. Some of the test data is machine-generated. All the results of the classification algorithm must be submitted, and the machine-generated samples will be ignored. The following provides the file descriptions.

1. training_variants - a comma separated file containing the description of the genetic mutations used for training. Fields are ID (the id of the row used to link the mutation to the clinical evidence), Gene (the gene where this genetic mutation is located), Variation (the amino acid change for this mutations), Class (1-9 the class this genetic mutation has been classified on)

2. training_text - a double pipe (||) delimited file that contains the clinical evidence (text) used to classify genetic mutations. Fields are ID (the id of the row used to link the clinical evidence to the genetic mutation), Text (the clinical evidence used to classify the genetic mutation)
3. test_variants - a comma separated file containing the description of the genetic mutations used for training. Fields are ID (the id of the row used to link the mutation to the clinical evidence), Gene (the gene where this genetic mutation is located), Variation (the amino acid change for this mutations)
4. test_text - a double pipe (||) delimited file that contains the clinical evidence (text) used to classify genetic mutations. Fields are ID (the id of the row used to link the clinical evidence to the genetic mutation), Text (the clinical evidence used to classify the genetic mutation)
5. submissionSample - a sample submission file in the correct format

Data Snippets: Train Data Frame

	ID	Gene	Variation	Text
0	0	ACSL4	R570S	2. This mutation resulted in a myeloproliferat...
1	1	NAGLU	P521L	Abstract The Large Tumor Suppressor 1 (LATS1)...
2	2	PAH	L333F	Vascular endothelial growth factor receptor (V...
3	3	ING1	A148D	Inflammatory myofibroblastic tumor (IMT) is a ...
4	4	TMEM216	G77A	Abstract Retinoblastoma is a pediatric retina...

Data Snippets: Test Data Frame

	Gene	Variation	Sentences
0	[chek2]	[h371y]	[[the, incidence, of, breast, cancer, is, incr...
1	[axin2]	[truncating, mutations]	[[an, unselected, series, of, 310, colorectal,...
2	[wnt4]	[e216g]	[[mycosis, fungoides, and, szary, syndrome, ar...
3	[suc1a2]	[g118r]	[[regulated, progression, through, the, cell, ...
4	[braf]	[t599instt]	[[pilocytic, astrocytoma, pa, is, emerging, as...

Text attribute is considered to be the main text component on which the inference is performed. These statistics will be used to asses what can be considered an average length when modelling with different forms of documents in terms of words. The dataset contains 8989 records for training set and 989 records for test set. Some statistics on text being words are explored.

Statistics on Words with Document as Words	Train Data Frame Text Attribute	Train Data Frame Text Attribute
Mean	1485.407709	1633.366090
Standard Dev	845.216131	484.593756
Min	1.000000	1.000000

25th Percentile	919.000000	1337.000000
50th Percentile	1215.000000	1642.000000
75th Percentile	1887.000000	1920.250000
Max	7199.000000	6633.000000

Exploratory Visualization

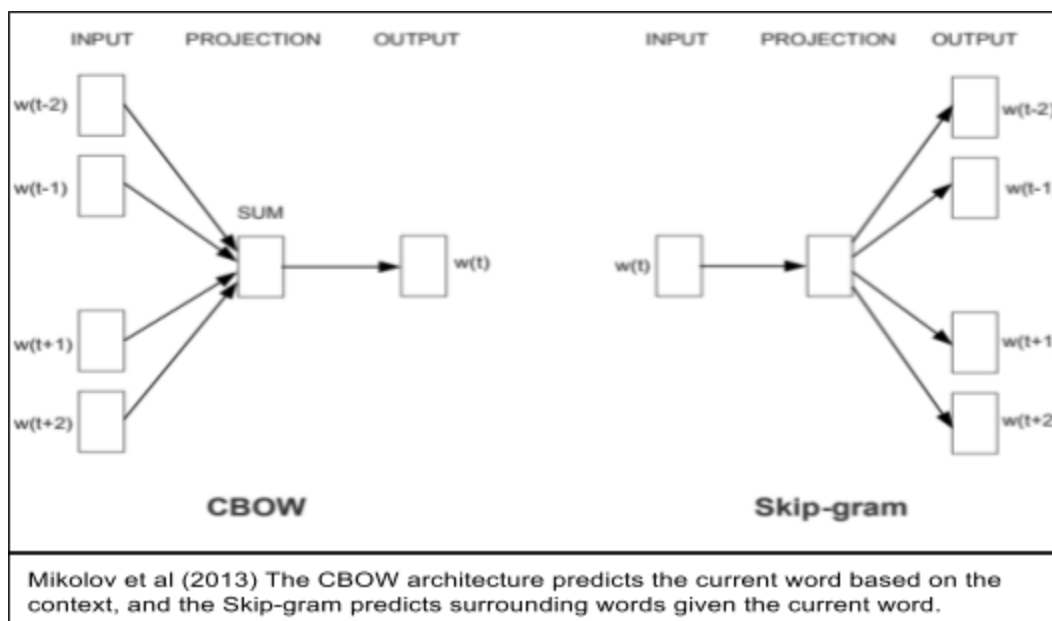
Models proposed for estimating continuous representation of words, such as Latent Semantic Analysis/Indexing and Latent Dirichlet Allocation, use documents for contextualizing information retrieval and are computationally expensive when used on large datasets. Recent models use words as contexts, which captures semantic similarity rather than semantic relatedness. Word embeddings created using unsupervised training on a corpus provide meaningful semantic similarity such as:

$$king - man + woman \approx queen$$

Word embeddings are low dimensional representations of words in vector space. They are also referred to as word vectors or distributed representation of words. Feed forward neural models such as the Continuous Skip-gram and the Continuous Bag of Words models introduced by Mikolov et al (2013) [4], provide better semantic and syntactic word similarities than count based models, with reduced computational cost.

The training methods and estimation of the loss function are beyond the scope of this project, but a quality reference can be found in Sebastian Ruder's blog post on word embeddings [5].

Joulin et al (2016) [6] [7] introduced the Fasttext [8] tool for word representation learning and text classification. This tool will be used to generate word vectors that form the basis for most of the deep models in which each the units considered are words. Multiple set of word vectors will be generated with varying dimensions (100,200,300), epochs (20, 50, 100) and models (skip gram and cbow). Sets of vectors generated from NLTK word2vec (trained by google), Glove [9] vectors from Stanford, and bionlp [10] word vectors based on medical text will also be used.



Lets visualize the word embeddings generated by training on the text collected from the dataset using skip gram model. These visualization are generated using tensorflow's tensorboard embeddings feature. The vocabulary is huge and the PCA accounting for just 8% of variations is not surprising since this is shows only 5000 words of 352220 words in the vocabulary.



t-SNE algorithm is run on the same set of words with perplexity 35 and learning rate of 0.1 (starts with 10 and manually reduced after viewing the changes in the dimensions of the word). The final visualization is presented below. The same word seems to be correlated differently based on their embeddings.



1. L2 norms, which help penalize larger model weights;

2. Batch Normalization, which helps with maintaining the internal covariance shift, and the ability to adapt to changing parameter distribution; and
3. Dropout layer, which helps reduce overfitting by randomly turning on/off the neurons and can be considered similar to training many models in reducing the variance overall.

When performing model training we mostly use ADAM optimizer for Stochastic Gradient Descent. Deeper discussion on regularization and training methods is beyond the scope of this project. Consider reading chapter 11 from book [13]

Deep model surveyed in this paper are listed below

1. CNN for Sentence Classification
2. DCNN for Modelling Sentences
3. VDDN for Text Classification
4. Multi Channel Variable size CNN
5. LSTM Deep Sentence Embedding
6. Multiplicative LSTM
7. Hierarchical Attention Networks
8. Recurrent Convolutional Network
9. C-LSTM Neural Networks
10. AC-BLSTM Networks

Benchmark

The best result from the count-based models described above will be considered as the benchmark model. The code categorical accuracy of 66% obtained from the sklearn's naive bayes model will be the benchmark result which will be used to evaluate the neural models. All the count based models will be displayed in the results section for comparison.

Methodology

Data Preprocessing

The foremost step in any text preprocessing is to decide on what character set the models will be working [2]. Since the data set is in English, character set "utf-8" is used to decode the text and special characters, not in selected unicode character set, are ignored. Also, the document contains only printable English characters; all others are removed. Looking at sample random documents, one can notice "figure", "table", "supplementary figure" and various iterations of these similar texts present throughout the data. All the texts are made lowercase for disambiguation.

Since in this project we will be generating/training word vectors based on skip gram and cbow models (will be discussed later) on the corpus, these unwanted texts contributing to the context words degrades the semantic relations. Hence, these texts will be removed using custom regex parsers. The hyperlink texts are not removed intentionally, which may provide some relation to the document.

After the above steps, the vocabulary is decided upon by dividing the text into words using the custom regular expression “`RegexTokenizer(r'\w+[-]*\w*')`”, which keeps the hyphenated words together since they may carry more information in medical texts. The delimiters will be space characters (tab, single or more space, newline). This will be the vocabulary used for all the models trained. For tokenizing a document into sentences there are multiple options available; here NLTK’s were chosen [3] (leading platform for building Python programs to work with human language data and contains wrappers for industrial-strength NLP libraries) as recommended sentence tokenizers (currently PunktSentenceTokenizer for the specified language).

For count based models, pre-processed text will be used. However, for neural models we need the ability to pass in the text in different forms. A DocumentGenerator Class helps in creating datasets as necessary in the following formats:

1. Document as Words: all document text as list of words
2. Document as Characters: all document text list of characters
3. Document as Sentence as Words: all document text as list of sentences that contain list of words
4. Document as Sentences as Characters: all document text as list of sentences that contain characters
5. Document as Sentences as Words as Characters: all document text as list of sentences that contain words represented as list characters

There are two data structures for maintaining vocabulary. One contains the list of vocabulary words and the other is a dictionary that maps the words to their position in the former. A corpus text is also formed with cleaned data that will be used for training word vectors.

Implementation

Models Based on n-grams

There are multiple n-gram count based classification models. This project uses these models as baseline models to assess whether the deep models are performing better. Bag of words model represents the text data by assigning each word an id and these ids will be used as features for the documents. These features represent number of occurrences of the word in the document. This can lead to issue of having too many features which could be handled with smaller vocabulary. The words not in the vocabulary will be replaced with a common id that does not occur in the vocabulary. This will result in very high dimensional sparse dataset. To scale down the impact of words with high frequency and dealing with differences for smaller and longer documents differently, we will be use Term Frequencies times Inverse Document Frequencies(tf-idf) with smoothing. There are many weighting schemes for using tf-idf, the one used in this project is of the form shown below

$$tf-idf(d, t) = tf(t) * idf(d, t)$$
$$idf(d, t) = \log\left(\frac{n}{df(d, t)}\right) + 1$$

Where n is total number of words in vocabulary, d is the document and t is the word in the document. $tf(t)$ represents the term frequency in the whole corpus. $df(d, t)$ represents the term frequency in the specific document.

After converting each document to the required tf-idf matrix, we use sklearn package to exercise all the necessary processing and functions to run the following machine learning algorithms:

1. Multinomial Naive Bayes classifier
2. Support Vector Machines
3. Softmax (Multi Class Logistic) Regression
4. K Nearest Neighbors
5. Passive Aggressive Classifier
6. Decision Trees
7. Adaboost Classifier
8. Extreme Randomization Trees
9. Random Forest Classifier

Catboost classifier from Yandex Technologies is also run on the dataset. Some models like sklearn's Gaussian Mixture Models and Xgboost were not considered since either they took more than 64GB of RAM or ran for more than 6 hours. Grid/Random searches are also not used on the above models considering the time factor. For all the runs, the vocabulary is not reduced in size since unique medical terms like gene names may carry more information.

Naive Bayes

Naive Bayes uses bayesian conditional probability model. The can be stated as

$$posteriordistribution = \frac{prior \times likelihood}{evidence}$$

The prior distribution on the words are generated using just word counts that exist in all the text data available, both training and testing. Then the likelihood estimates are calculated by grouping the the text data from training based on genetic mutations over which classification is performed. Then we can perform maximum likelihood estimation on the bayes formula.

$$P(genetic_mutation|text) = \frac{P(text|genetic_mutation) \times P(genetic_mutation)}{p(text)}$$

Since this is a maximum likelihood estimation, say we have m mutations, we can represent the solution as

$$genetic_mutation|text = argmax_m P(text|genetic_mutation) \times P(genetic_mutation)$$

We have omitted the $P(text)$ since it does not contribute the argmax operation.

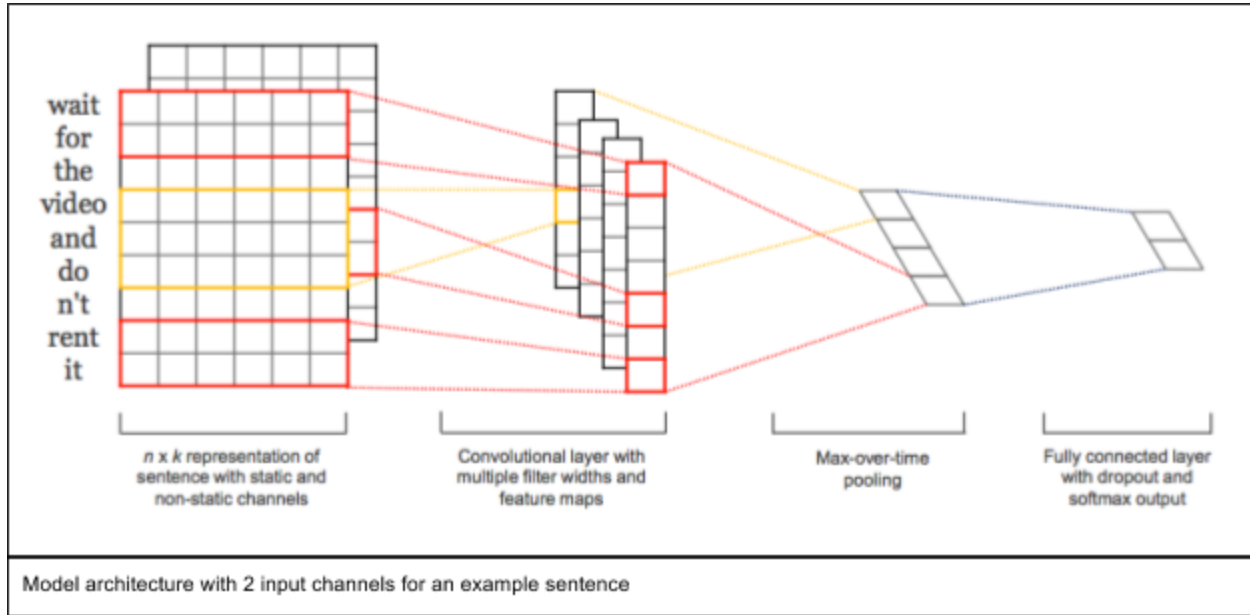
The solution to the naive bayes and other count based models can be seen [here](#)

Models Based on Convolutional Neural Networks

CNN for Sentence Classification

Yoon et al (2014) [14] proposed CNN models in addition to pre-trained word vectors, which achieved excellent results on multiple benchmarks. The model architecture as shown in the figure below maintains multiple channels of input such as different types of pre-trained vectors or vectors that are kept static during training. Then they are convolved with different kernels/filters to create sets of features that are then max

pooled. These features form a penultimate layer and are passed to a fully connected softmax layer whose output is the probability distribution over labels.



The paper presents several variants of the model:

1. CNN-rand (a baseline model with randomly initialized word vectors)
2. CNN-static (model with pre-trained word vectors)
3. CNN-non-static (same as above but pre-trained fine tuned)
4. CNN-multichannel (model with 2 sets of word vectors)

Keras model summary and code for CNN-non static can be found [here](#).

DCNN for Modelling Sentences

Kalchbrenner et al (2014) [15] presented Dynamic Convolutional Neural Network for semantic modelling of sentences. This model handles sentences of varying length and uses dynamic k-max pooling over linear sequences. This helps the model induce a feature graph that is capable of capturing short and long range relations. K-max pooling, different from local max pooling, outputs k-max values from the necessary dimension of the previous convolutional layer. For smooth extraction of higher order features, this paper introduces Dynamic k-max pooling where the k in the k-max pooling operation is a function of the length of the input sentences.

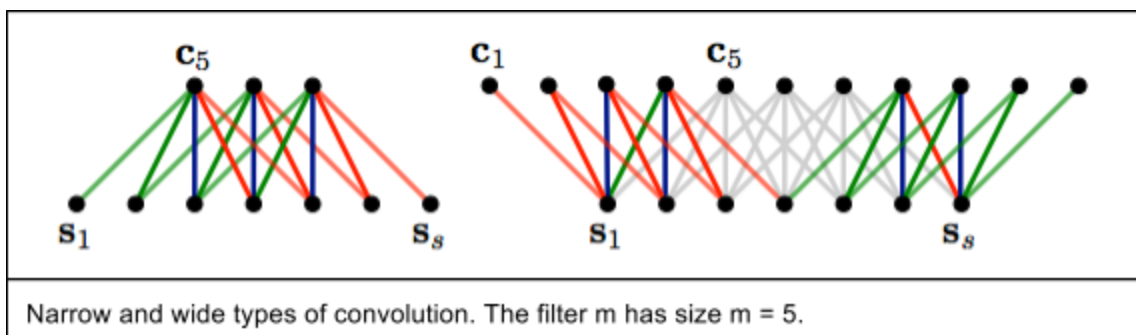
$$k = \max(k_{top}, \lfloor \frac{L-l}{L} \cdot s \rfloor)$$

where l is the number of convolutional layer to which pooling is applied, L is total number of convolutional layers, k_{top} is the fixed pooling parameter of the top most convolutional layer. The model also has a folding layer which sums over every two rows in the feature-map component wise. This folding operation is valid since feature detectors in different rows are independent before the fully connected layers.

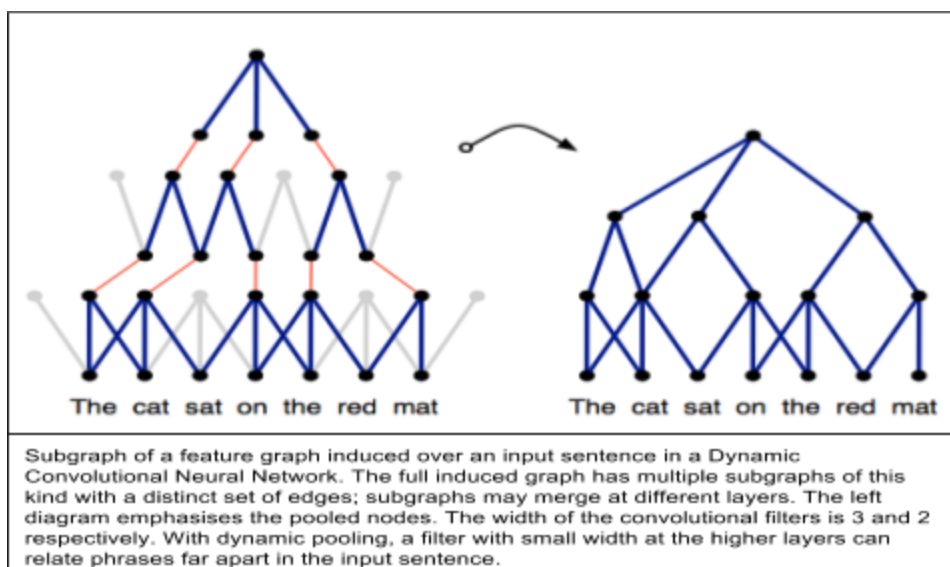
Wide convolutions are preferred for the model instead of narrow convolutions. This is achieved in the code using appropriate zero padding.

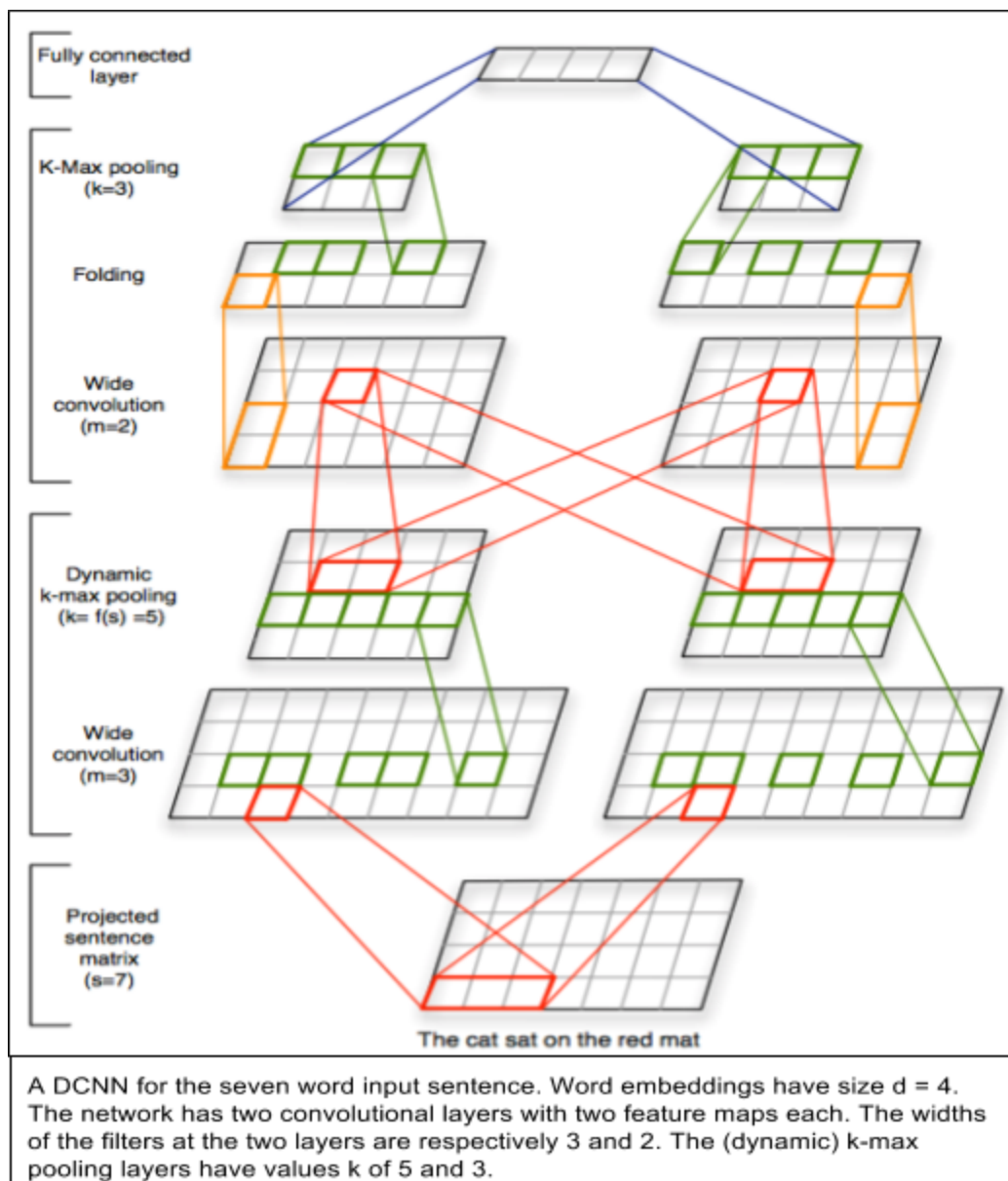
This network model's performance is related to its ability to capture the word and n-gram order in the sentences and to tell the relative position of the most relevant n-grams. The model also has the advantage of inducing a connected, directed acyclic graph with weighted edges and a root node as shown below.

Folding and K-max pooling layers are not readily available and have to be created using keras functional apis.



Keras model summary and code for DCNN can be found [here](#).





VDNN for Text Classification

Conneau et al (2016) [17] presented Very Deep CNN, which operates directly at the character level. This model also shows that deeper models perform better and are able to learn hierarchical representations of whole sentences.

The overall architecture of the model contains multiple sequential convolutional blocks. The two design rules followed are:

1. For the same output temporal resolution, the layers have the same number of feature maps, and
2. When the temporal resolution is halved, the number of feature maps are doubled. This helps reduce memory footprint of the model.

The model contains three pooling operations that halve the resolution each time resulting in three levels of 128, 256, 512 feature maps. There is an optional shortcut connection between the convolutional blocks that can be used, but since the results show no improvement in evaluation, that component was dropped in this project.

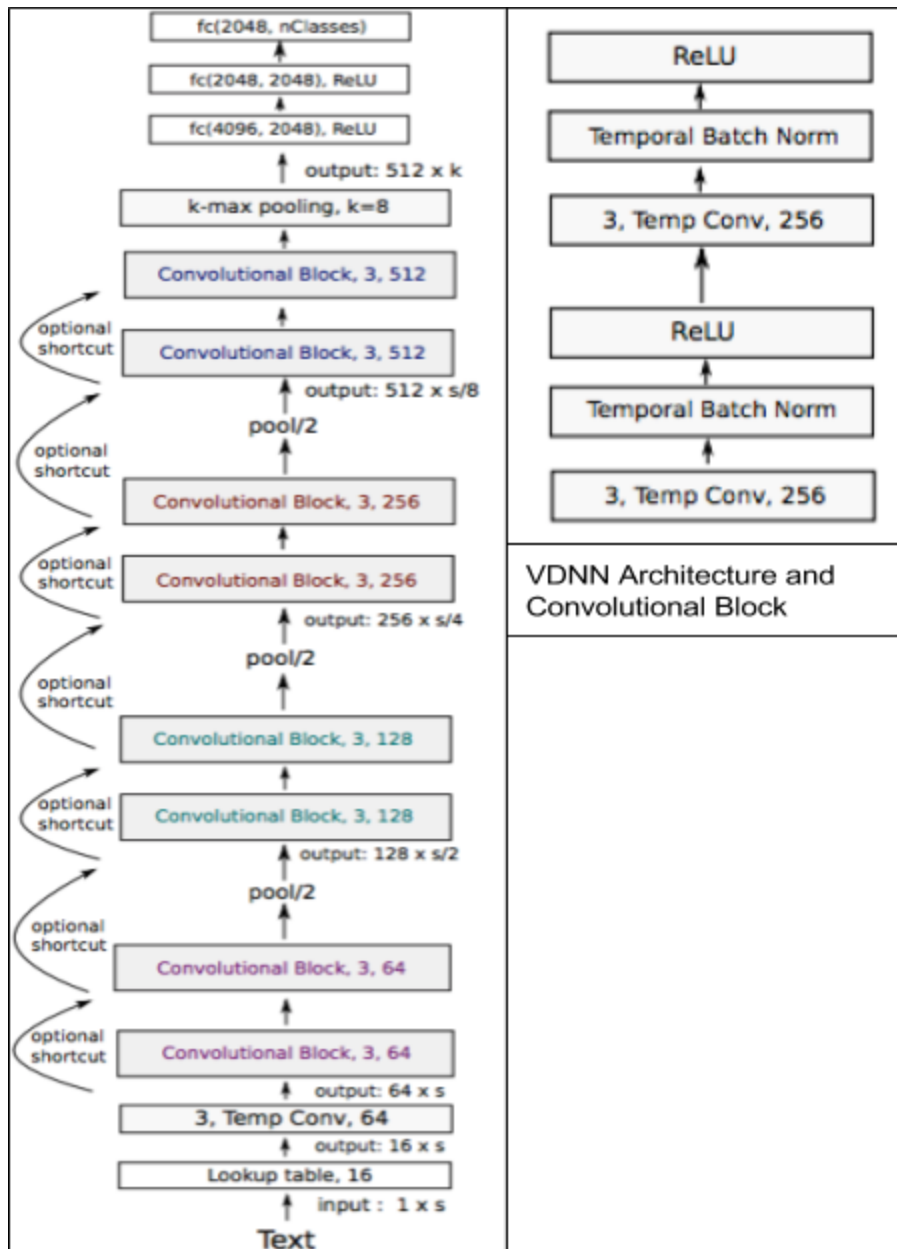
Each convolutional block is a sequence of convolutional layers, each followed by batch normalization layer and relu activation. The down sampling of the temporal resolution between the conv layers (K_i and K_{i+1}) in the block are subjected to multiple options:

1. The first conv layer K_{i+1} has stride 2
2. K_i is followed by k-max pooling layer where k is such that resolution is halved
3. K_i is followed by max pooling layer with kernel size 3 and stride 2

Characters used in character vectors for all the models in this paper are:

abcdefghijklmnopqrstuvwxyz0123456789-.,!?:'"/| # \$ % ^ & * ~ ' + = < > () [] { }

The VDNN architecture and each convolutional block are presented below. Keras model summary and code for VDNN can be found [here](#).



Multi Channel Variable size CNN

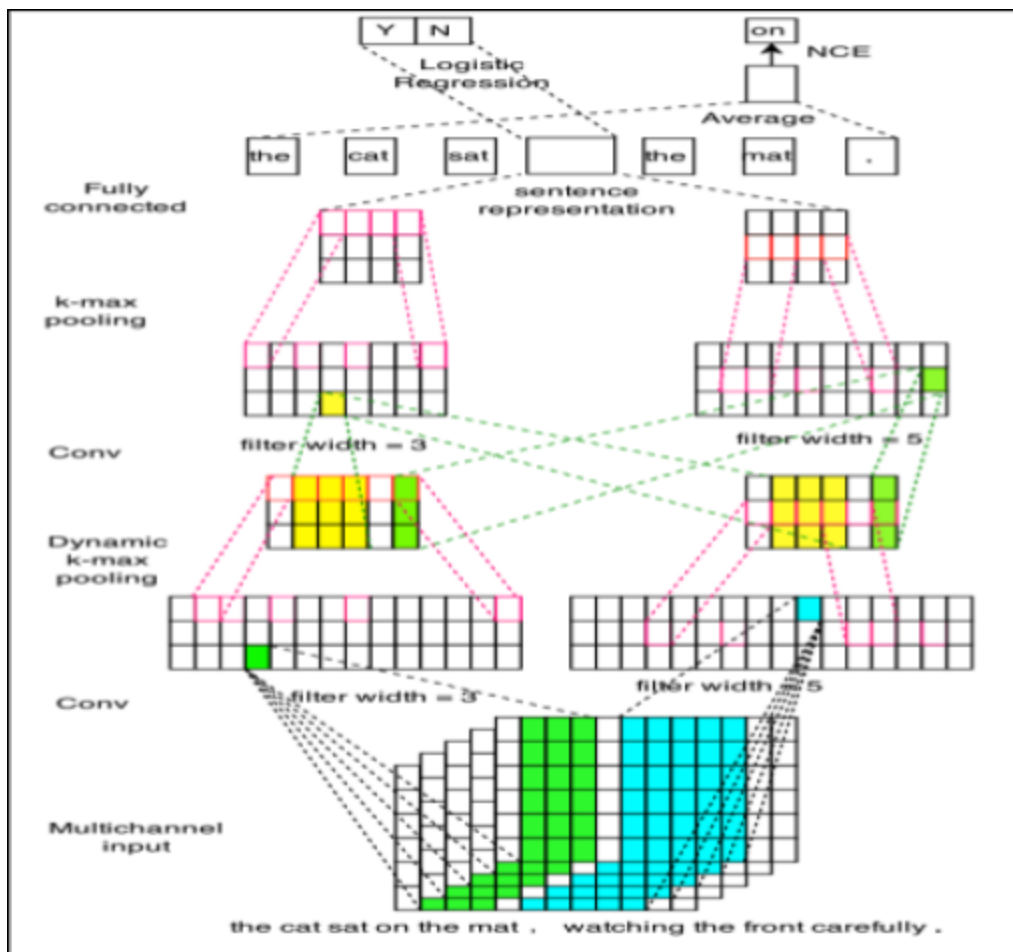
Yin et al (2016) [20] propose MV-CNN, which combines diverse versions of pre-trained word embeddings and extract features from multi-granular phrases with variable-size convolutions. Using multiple embeddings of the same dimension from different sets of word vectors should contain more information that can be leveraged during training.

The paper describes maintaining three dimensional embedding matrices with each channel representing a different set of text embeddings. This multi-channel initialization might help unknown words across different embeddings. Frequent words can have multiple representations and rare words (partially known words) can be made up with other words. The model then has two sets of convolution layers and dynamic k-max pooling layers followed by a fully connected layer with softmax (or logistic) as the last layer.

The paper describes two tricks for model enhancement. One is called mutual learning, which is implemented in this project. The same vocabulary is maintained across the channels and they help each other tune parameters while training. The other trick is to enhance the embeddings with pretraining just like a skip-gram model or autoencoder using noise-contrastive estimation.

Different sets of embeddings used for this model comes from glove, word2vec, custom trained vectors on the train and test corpus using fast text tool as discussed in the section [Word Vectors](#).

Keras model summary and code for MVCNN can be found [here](#).

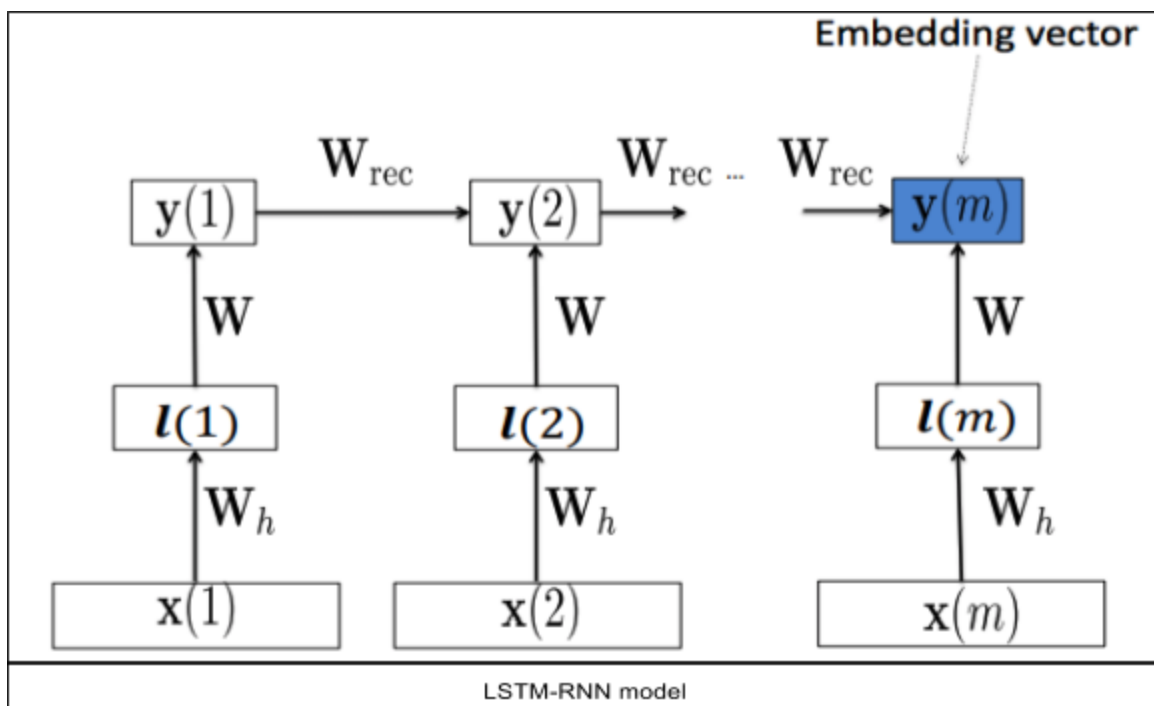


Models Based on Recurrent Neural Networks

LSTM Deep Sentence Embedding

Palangi et al (2015) [29] proposed LSTM-RNN for text classification. The model has shown to perform better than [Paragraph Vectors](#) for document/sentence embedding. As the model reads to the end of the sentence, the topic activation accumulates and the hidden state representation at the last word encodes the rich contextual information of the entire sentence. LSTM-RNN are effective against noise and can be robust in scenarios when every word in the document is not equally important and only salient words need to be remembered using limited memory.

This is a very simple model that takes all the words in the document sequentially and the final output gives the document embedding. The model does not use max pool layers to capture global contextual information.



Keras model summary and code for LSTM based embedding model can be found [here](#).

Multiplicative LSTM

Krause et al (2016) [30] proposed Multiplicative LSTM which is a hybrid RNN that gives the model more flexibility in choosing recurrent transitions for each possible input, which makes it more expressive in autoregressive density estimation. This model is proposed for sequential language modelling, but it was chosen for this project to see how it helps in text classification tasks. The authors argue that current RNNs have a hard time recovering from mistakes when predicting sequences. If RNN hidden state remembers erroneous information then it might take more time to recover and there will be a snowball effect. Some proposed solutions such as introducing latent variables and increasing memory will only result in complex intractable distribution over the hidden states and reinterpreting stored inputs. The multiplicative model provides more flexibility for transitions in these kind of situations.

Since this is a modification to existing RNN unit (LSTM or GRU), this has to be implemented as a keras Layer similar to keras LSTM layer. The equations for mLSTM are

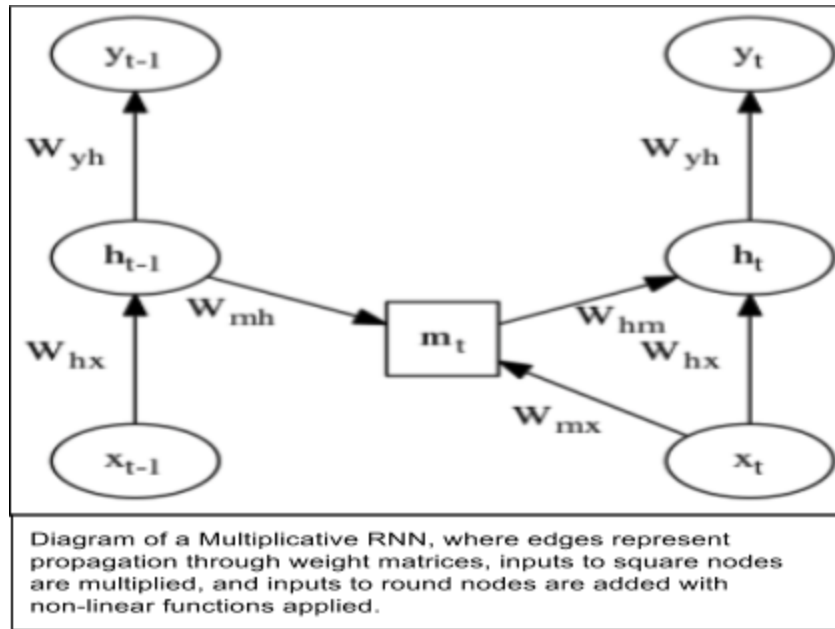
$$m_t = (W_{mx} \cdot x_t) \odot W_{mh} \cdot h_t$$

$$\hat{h}_t = (W_{hx} \cdot x_t) + W_{hm} \cdot m_t$$

$$i_t = (W_{ix} \cdot x_t) + W_{im} \cdot m_t$$

$$o_t = (W_{ox} \cdot x_t) + W_{om} \cdot m_t$$

$$f_t = (W_{fx} \cdot x_t) + W_{fm} \cdot m_t$$



Keras model summary and code can be found [here](#)

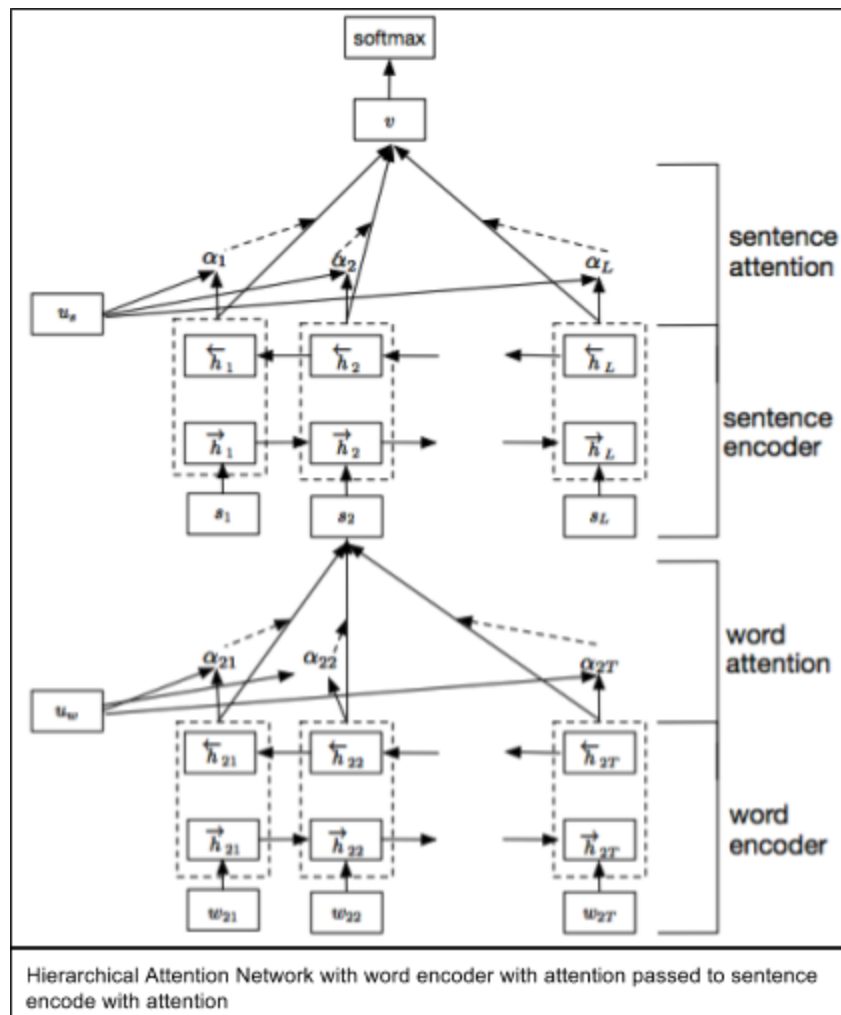
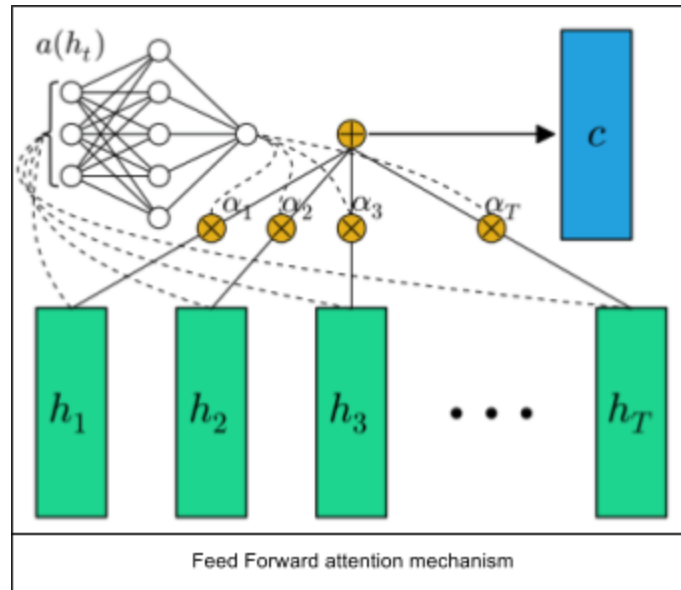
Hierarchical Attention Networks

Yang et al (2016) [31] proposed Hierarchical Attention Networks for document classification, which took advantage of a hierarchical structure that mirrors the hierarchical structure of the documents and two levels of attention mechanisms applied to differentially more and less important content when constructing document representation.

First let us look into what attention mechanism means. For this, let us look into the feedforward attention mechanism proposed by Raffel et al (2015) [32]. Attention mechanism allows more direct dependencies between the states of the model at different points in time. From the picture, the vectors seen in the hidden state sequence are fed into a learnable function that produces a probability vector, which in turn is used to find the weighted average of the output hidden states.

Similarly, Hierarchical Attention Networks proposes a two stage attention mechanism to encode documents. The intuition behind the model is that different words and sentences in a document are differentially informative. The document representations are constructed from each sentence and each sentence representation is constructed from words. Each sentence in a document can be considered as a word sequence that is fed into a generative LSTM layer with attention on top, and again the same procedure is followed for sentences with attention on top. Finally a softmax layer on top to classifying the document labels.

Keras does not provide apis for attention mechanism, hence a custom Attention layer is created to handle different levels of attentions.

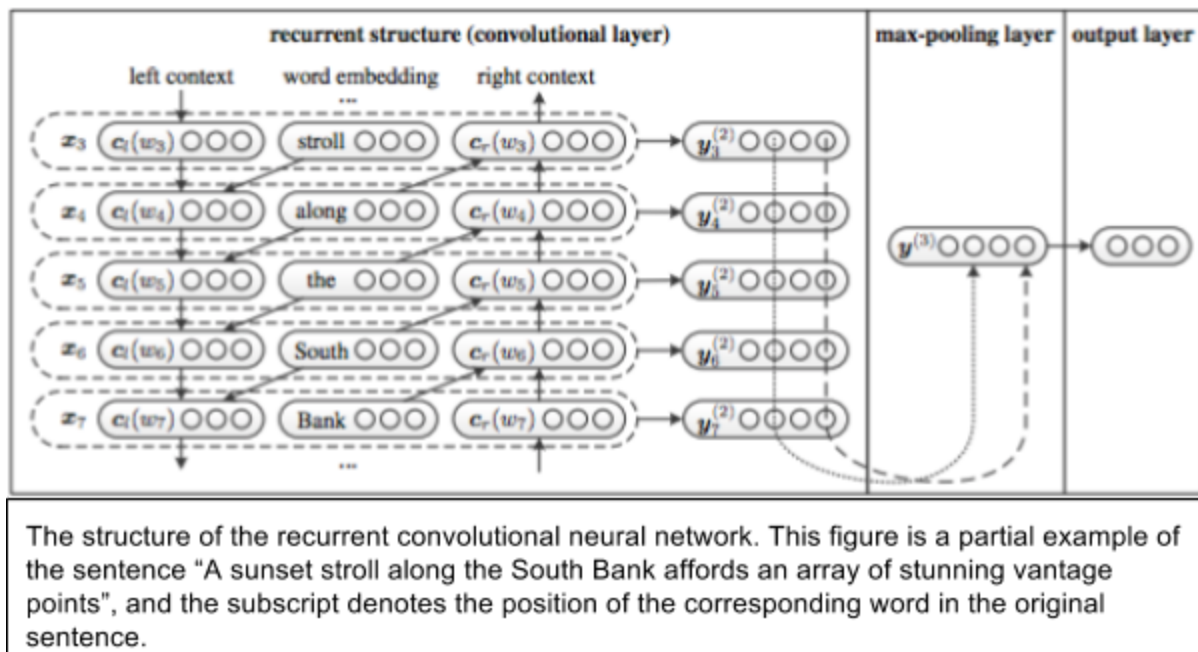


Keras model summary and code for HAN model can be found [here](#)

Models Based on CNN and RNN ensemble

Recurrent Convolutional Network

Lai et al (2015) [33] proposed RCNN model that addresses the limitation of unbiased CNN models with shorter window sizes and biased RNN models. The model has a bi-directional recurrent structure that reduces noise and captures semantic information to the greatest extent possible. Max pool layer on top of recurrent structure judges the features role in capturing key components necessary for classification.



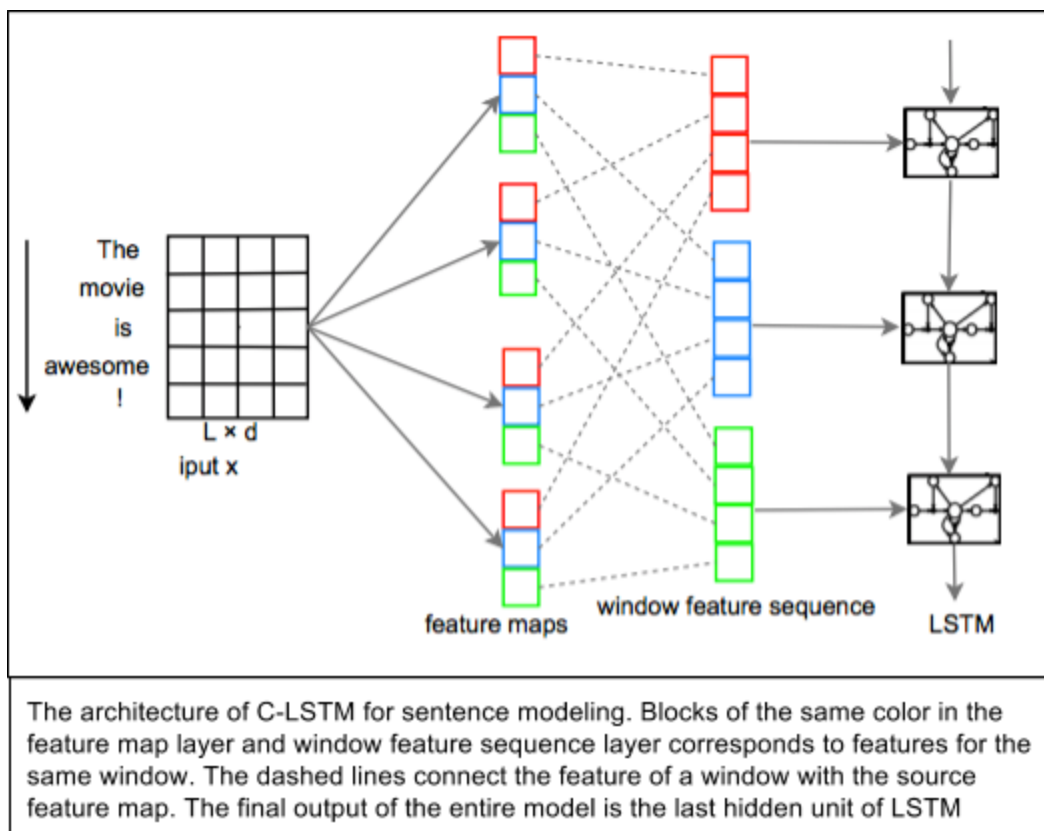
The model has bidirectional LSTM that captures context information from nearby words in both the directions. Then the max pool layer captures the key features that are fed as input into the softmax layer that classifies the text provided.

Keras model summary and code for RCNN can be found [here](#)

C-LSTM Neural Networks

Zhou et al (2015) [35] proposed C-LSTM Neural Networks, which uses CNN to capture local features of phrases and RNN to capture global and temporal sentence semantics.

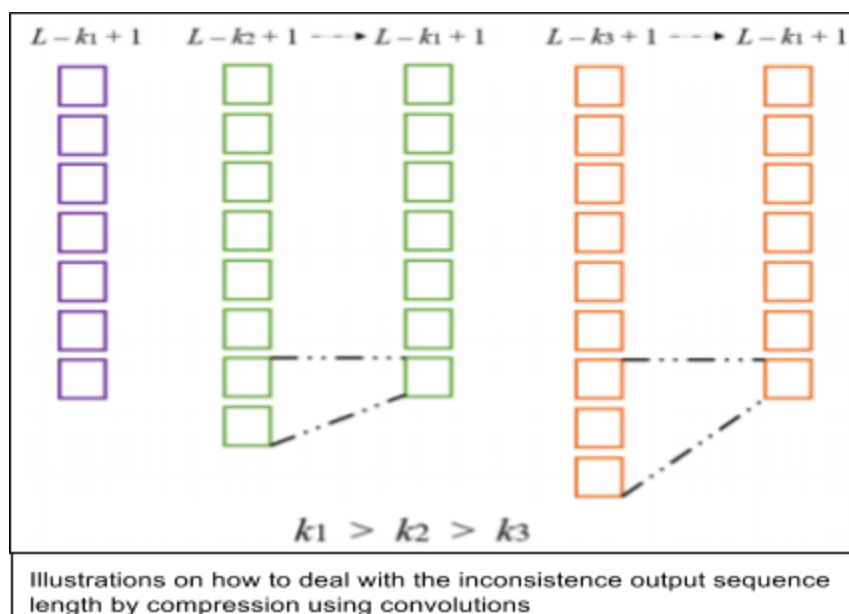
The model extracts high-level correlations from n-gram features with shorter windows in CNN models and are fed as a sequence to the following LSTM layer, which helps with creating an embedding for the document for classification. The output of each convolution (single feature map) has semantic information from the whole sentence as a sequence. From these multiple feature maps, without using a max-pool layer, the vectors representing positional information are grouped or concatenated which are fed to RNN layer as a sequence.



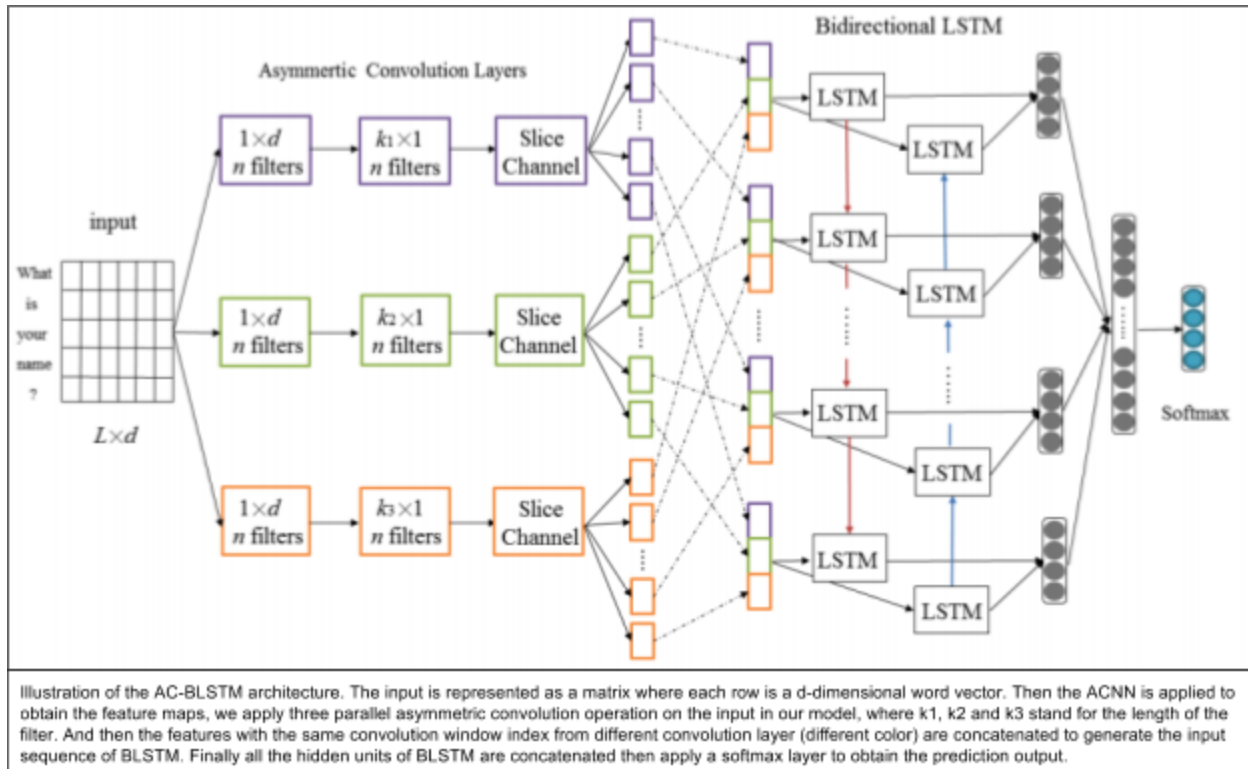
Keras model summary and code for the c-lstm model can be found [here](#)

AC-BLSTM Networks

Liang et al (2016) [37] proposed a model that combines asymmetric CNN (ACNN) and bidirectional LSTM (BLSTM). In order to make the models deep they factorized $n \times n$ convolutions to $1 \times n$ and $n \times 1$ convolutions. Another important feature of the model how the asymmetric convolutions are handled, instead of truncating lengths, smaller convolutions are performed on them. This model is similar to previous seen model Combination of CNN and RNN in handling how the feature are fed from CNN to RNN. RNN employed here is a bidirectional LSTM whose outputs are concatenated and fed as input to the final softmax layer for classification.



Keras model summary and code for the ac-blstm model can be found [here](#)



Refinement

I chose model "CNN for Sentence Classification" for refinement. The following are the refinement steps used to get better categorical accuracy as shown in the results section.

1. Increased initial number of filter which would capture not just phrases but also other structures. Since the whole document is considered for training and validation, filter with longer kernels captures longer sentence features.
2. After noting the high difference between the validation set and the test set accuracies, l2 kernel regularizers was used for the last convolution layer.
3. Performing 5-fold cross validation which improved accuracy

The above refinements provided 9% increase in the accuracy.

Results

The results are summarised in the below table. The kaggle competition compares the results against the predicted categories for the test data. Since this is a study, the information below compares the categorical accuracy obtained with the validation set. The validation set is same across all the models including the count models, controlled using random seed. Cross validation is performed on naive bayes model for various alpha parameter values.

Model Names	Categorical Accuracy
Yandex Technologies CatBoost	0.587453613072
SKlearn Multinomial Naive Bayes	0.663663663664
SKlearn Support Vector Machine	0.285285285285
SKlearn Softmax Regression	0.507507507508
SKlearn K Nearest Neighbour	0.612612612613
SKlearn Passive Aggressive Classifier	0.630630630631
SKlearn Decision Trees Classifier	0.597597597598
SKlearn AdaBoost Classifier	0.408408408408
SKlearn Random Forest Classifier	0.627627627628
Extreme Randomization Trees	0.630630630631

I decided to study all the models with keras default parameters for the layer arguments and hyperparameter tuning as suggested in the papers. Since there are multiple models available for refinement, I picked one of the foremost CNN model used for sentence classification “CNN for sentence classification” in the literature. Now, let's look at the results generated from these deep models.

Model Names	Categorical Accuracy
CNN for Sentence Classification (refined)	0.7606 (unrefined result 0.6276)
DCNN for Modelling Sentences	0.66366
VDNN for Text Classification	0.52600
Rationale Augmented CNN	0.65766
Multi Channel Variable size CNN	0.60164
Multiplicative LSTM	0.59331
Hierarchical Attention Networks	0.66967
Recurrent Convolutional Network	0.65465
C-LSTM Neural Networks	0.55856
AC-BLSTM Networks	0.59760

We can see all the non tuned models are fairly performing better than best count based models. Making refinements discussed above helped increasing the selected model accuracy by nearly 10%. It is clear hyperparameter tuning is important when it comes to deep neural models. The improvement can be attributed to the rich features being captured by increased number of different sized filters and cross validation.

Conclusion

The models discussed and implemented in this project are a collection of selected deep models that depend on state-of-the-art Dense Neural Network models CNN and RNN. In medical text data, I have seen only few papers making use of CNN based models to classify important medical tasks. Based on the results obtained for this one task from all the models trained, it is clear that there is much more room to improve and identify different deep neural architectures based on CNN and RNN that can be leveraged in the area of classifying medical text data.

More data always help reducing the variance in the model. There was a second stage for the kaggle competition from which the dataset was procured. In future project, I will train all the above models with more data provided to see how the training metric changes.

Levy et al (2015) [40] performed an extensive study on comparing neural models and count based n-gram models. This revealed that much of the performance gain of word embeddings are from design choices and hyperparameters rather than much revered word embeddings. This does leave more room for improvement and more research to be conducted to find better models for text classification. Similar surveys must be conducted on same data once a year to better assess the power of the deep neural models in the area of text classification for clinical evidence medical text data.

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