Learning Kernels over Strings using Gaussian Processes

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Abstract

Non-contiguous word sequences are widely known to be important in modelling natural language. However they are not explicitly encoded in common text representations. In this work we propose a model for text processing using string kernels, capable of flexibly representing non-contiguous sequences. Specifically, we derive a vectorised version of the string kernel algorithm and their gradients, allowing efficient hyperparameter optimisation as part of a Gaussian Process framework. Experiments on synthetic data and text regression for emotion analysis show the promise of this technique.

1 Introduction

Text representations are a key component in any Natural Language Processing (NLP) task. A common approach for this is to average word vectors over a piece of text. For instance, a bag-of-words (BOW) model uses one-hot encoding as vectors and it is still a strong baseline for many tasks. More recently, approaches based on dense word representations (Turian et al., 2010; Mikolov et al., 2013) also showed to perform well.

However, averaging vectors discards any word order information from the original text, which can be fundamental for more involved NLP problems. Convolutional and recurrent neural networks (CNNs/RNNs) can keep word order but still treat a text fragment as a contiguous sequence of words, encoding a bias towards short- over long-distance relations between words. Some RNN models like the celebrated LSTMs (Hochreiter and Schmidhuber, 1997) perform better in capturing these phenomena but still have limitations. Recent work (Tai et al., 2015; Eriguchi et al., 2016) showed evidence that LSTM-based models can be enhanced by adding syntactic information, which can encode relations between non-contiguous words. This line of work requires the employment of accurate syntactic parsers, restricting their applicability to specific languages and/or text domains.

In this work we propose to revisit an approach which goes beyond contiguous word representations: string kernels (SKs). Their main power comes from the ability to represent arbitrary non-contiguous word sequences through dynamic programming algorithms. Our main contribution is a model that combines SKs with Gaussian Processes (GPs) (Rasmussen and Williams, 2006), allowing us to leverage efficient gradient-based methods to learn kernel hyperparameters. The reasoning behind our approach is that by optimising hyperparameters in a fine-grained way we can guide the kernel to learn better task-specific text representations automatically.

To enable the learning procedure we redefine the SK algorithm in a vectorised form and derive its gradients. We perform experiments using synthetic data, giving evidence that the model can capture non-trivial representations. Finally, we also show how the approach fares in a real dataset and explain how the learned hyperparameters can be interpreted as text representations.

2 String Kernels

Here we give a brief intuition1 on string kernels, based on the formulation proposed by Cancedda

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1We give a thorough explanation of the original SK equations in the Supplementary Material, as well as a detailed derivation of our vectorised version with its hyperparameter gradients.
et al. (2003). Let $|s|$ be the length of a string $s$, $s_j$ the $j$-th symbol in $s$ and $s_{-1}$ the prefix containing the full string $s$ except for the last symbol. Define $\text{sim}(a, b)$ as a similarity measure between individual symbols $a$ and $b$. Given two strings $s$ and $t$ and a maximum n-gram length $n$, the string kernel $k(s, t)$ can be obtained using the recursion

$$k_0'(s, t) = 1, \text{ for all } s, t,$$

$$k_i'(sa, t) = \lambda_g k_i'(sa, t) + k_i''(sa, t),$$

$$k_n'(sa, t) = k_n(s, t) + \lambda_m \sum_j \text{sim}(a, t_j)k_{n-1}'(s, t_{-1}),$$

$$k(s, t) = \sum_{i=1}^n \mu_i k_i(s, t),$$

where $\lambda_g$ and $\lambda_m$ are decay hyperparameters for symbol gaps and matches, respectively, and $\mu_i$ is the weight for the kernel of n-gram order $i$. The decay hyperparameters smooth the kernel values when sequences are very similar to each other while the n-gram weights help to tune the signal coming from different subsequence lengths.

Our goal is to optimise the kernel hyperparameters in a fine-grained way using gradient-based methods. For this, we first redefine the kernel in a vectorised form. This not only eases gradient derivations but also allow our implementation to capitalise on recent advances in parallel processing and linear algebra libraries for better performance. Given two strings $s$ and $t$, the equations for our vectorised version are defined as

$$S = E_s E_t^T,$$

$$K'_0 = 1,$$

$$K'_i = D_{|s|} K''_i D_{|t|},$$

$$K''_i = \lambda_m^2 (S \circ K'_{i-1}),$$

$$k_i = \lambda_m^2 \sum_{j,k} (S \circ K'_i)_{j,k},$$

$$k(s, t) = \mu^T k,$$

where $E_s$ and $E_t$ are matrices of symbol embeddings for each string, $\circ$ is the Hadamard product

and $D_\ell \in \mathbb{R}^\ell \times \mathbb{R}^\ell$ is the matrix

$$D_\ell = \begin{bmatrix} 0 & \lambda_g^0 & \ldots & \lambda_g^{(|s| - 2)} \\ 0 & 0 & \ldots & \lambda_g^{(|t| - 3)} \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \lambda_g^0 \\ 0 & 0 & 0 & 0 \end{bmatrix}$$

with $\ell \in \{|s|, |t|\}$ being the corresponding string length for $s$ or $t$. The purpose of $D$ is to unroll the recursion from the original kernel equations. For the matrices $E$, we focus on dense word embeddings but they could also be one-hot vectors, simulating hard matching between symbols.

Given this formulation, the hyperparameter gradients can be easily derived. From the vectorised definition, we can see that gradients with respect to $\mu$ are simply $k$, the intermediate n-gram specific kernel values. For $\lambda_g$ and $\lambda_m$ the gradients simply follow the kernel equations in an analogous manner. Note that the gradient calculations do not affect the time or space complexity of the main kernel, and in practice they can be obtained together using a single algorithm since they share many common terms.

Finally, we incorporate the kernel into a Gaussian Process regression model (henceforth, GP-SK). We assume the label $y$ for an input string $s$ is sampled from a function $f(s) \sim \mathcal{GP}(0, K(s, t))$, with $t$ iterating over all other strings in a dataset. With this, we can define the marginal likelihood as

$$\log p(y|s, \theta) = \log \int_f p(y|s, \theta, f) p(f),$$

$$= - \frac{y^T G^{-1} y}{2} - \frac{\log |G|}{2} - n \log 2\pi,$$

where $G$ is the Gram matrix with respect to the training set $s$ and $\theta$ is the set of kernel hyperparameters. By taking its derivative and plugging in the kernel gradients, we can optimise its hyperparameters using gradient-based methods.

### 2.1 Complexity and Runtime Analysis

The original string kernel algorithm has complexity $O(n|s||s'|)$, i.e., quadratic in the size of the largest string. Our vectorised version is cubic, $O(n\ell^3)$, where $\ell = \max(|s|, |s'|)$, due to two matrix multiplications in the equations for $K'_i$.

Another way of reaching this result is to realise that

We refer the reader to Rasmussen and Williams (2006, Chap.5) for an in-depth explanation of this procedure.
K′′_i is actually not needed anymore: all calculations can be made by updating K′_i only. In fact, Lodhi et al. (2002) introduced the term k′′ as a way to reduce the complexity from O(n|s||s'|^2) to O(n|s|s'|). The complexity for the gradient calculations is also O(n|s|s'|).

However, even though our vectorised version has higher complexity, in practice we see large gains in runtime. We assess this empirically running the following experiment with synthetic strings:

- We employ characters as symbols with a one-hot encoding as the embedding, using all English ASCII letters, including uppercase (52 symbols in total);
- The maximum subsequence length is set to 5;
- 100 instances are generated randomly by uniformly sampling a character until reaching the desired string length.

We test our kernels with lengths ranging from 10 to 100.

Figure 1 shows wall-clock time measurements as the string lengths increase. It is clear that the vectorised version is vastly faster than the original one, with up to two orders of magnitude. Comparing the CPU and GPU vectorised implementations, we see that we can reap benefits using a GPU when dealing with long sentences. GPU processing can be further enhanced by allowing portions of the Gram matrix to be calculated in batches instead of one instance at a time.

These results are intriguing because we do not expect a quadratic complexity algorithm to be outperformed by a cubic one. It is important to note that while we made efforts to optimise the code, there is no guarantee that either of our implementations is making the most of the underlying hardware. We plan to investigate these matters in more detail in the future.

3 Experiments

We assess our approach empirically with two sets of experiments using natural language sentences as inputs in a regression setting. The first one uses synthetically generated response variables, providing a controlled environment to check the modelling capacities of our method. The second set uses labels from an emotion analysis dataset and serve as a proof of concept for our approach.

3.1 Synthetic Labels

Consider an ideal scenario where the data is distributed according to a GP-SK. Here we aim at answering two questions: 1) whether we can retrieve the original distribution through optimisation and 2) whether a simpler model can capture the same distribution. The first gives us evidence on how feasible is to learn such a model while the second justify the choice of a SK compared to simpler alternatives.

To answer these questions we employ the following protocol. First we define a GP-SK with the following hyperparameter values:

\[
\begin{align*}
\lambda_g &= 0.5 \\
\lambda_m &= 0.2 \\
\sigma^2 &= 0.1 \\
\mu_1 &= 1.0 \\
\mu_2 &= 0.5 \\
\mu_3 &= 0.25
\end{align*}
\]

where \(\sigma^2\) is the label GP noise. This choice of hyperparameter values is arbitrary: our goal is simply to check if we can retrieve these values through optimisation. The same procedure could be applied for different values.

After defining the GP-SK model we calculate the its corresponding Gram matrix using a set of sentences and their respective word embeddings. This matrix contains the covariances of a multivariate Gaussian distribution with mean vector \(\mathbf{0}\) and we can sample from this Gaussian to create synthetic labels. As inputs we use a random sample of sentences from the Penn Treebank (Marcus...
et al., 1993) and represent each word as a 100d GloVe embedding (Pennington et al., 2014). The data sampled from the procedure above is used to train another GP-SK with randomly initialised hyperparameters, which are then optimised. We run this procedure 20 times, using the same inputs but sampling new labels every time.

**Hyperparameter stability** Figure 2 shows the hyperparameter values retrieved after optimisation, for increasing training set sizes. The decay hyperparameters are the most stable ones, being retrieved with high confidence independent of the dataset size. The original noise value is also obtained but it needs more instances (1000) for that.

The n-gram coefficients are less stable compared to the other hyperparameters, although the uncertainty seems to diminish with larger datasets. A possible explanation is the presence of some level of overspecification, meaning that very different coefficient values may reach similar marginal likelihoods, which in turn corresponds to multiple plausible explanations of the data. Solutions for this include imposing bounds to the coefficients or fixing them, while freely optimising the more stable hyperparameters.

**Predictive performance** To evaluate if the GP-SK models can be subsumed by simpler ones, we assess the predictive performance on a disjoint test set containing 200 sentences. Test labels were sampled from the same GP-SK distribution used to generate the training labels, simulating a setting where all the data follows the same distribution.

Figure 3 shows results across different training set sizes, in Pearson’s $r$ correlation. We compare with GP baselines trained on averaged embeddings, using either a linear or a Squared Exponential (SE)$^7$ kernel. The SK model outperforms the baselines, showing that even a non-linear model can not capture the GP-SK distribution.

To investigate the influence of hyperparameter optimisation, we also show results in Figure 3 for a SK model with randomly initialised hyperparameter values. Clearly, optimisation helps to improve the model, even in the low data scenarios.

### 3.2 Emotion Analysis

As a first step towards working with real world data, we employ the proposed approach in an emotion analysis setting, where the goal is to model latent emotions in text. We use the “Affective Text” dataset from the SemEval2007 shared task (Strapparava and Mihalcea, 2007), composed of 1,250 News headlines annotated with 6 scores, one per emotion. Scores are in the $[0−100]$ range and were provided by human judges. The models, baselines and embeddings are the same used in Section 3.1. Instead of using a fixed split, we perform 10-fold

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6[nlp.stanford.edu/projects/glove](http://nlp.stanford.edu/projects/glove). We use the version trained on Wikipedia and Gigaword.

7Also known as RBF kernel.
Table 1: Emotion analysis results, averaged over all emotions and cross-validation folds.

|       | NLPD | MAE  | r    |
|-------|------|------|------|
| SK    | 4.06 | 10.53| 0.586|
| Linear| 4.09 | 11.03| 0.539|
| SE    | 4.03 | 10.09| 0.611|

Table 2: SK hyperparameter values for a single model predicting the emotion **surprise**.

| λ_g  | 7.36 × 10^{-7} | λ_m | 0.0918 |
|------|----------------|-----|--------|
| µ_1  | 12.37          | µ_2 | 33.73  |
| µ_3  | 154.51         | µ_4 | 2.58   |
| µ_5  | 8.54           |     |        |

cross validation and average the results.

Table 1 compares the performance of GP-SK with the baselines trained on averaged embeddings. Besides Pearson’s $r$ correlation, we also compare the models in terms of Mean Absolute Error (MAE) and Negative Log Predictive Density (NLPD), a metric that takes into account the full predictive distribution into account (Quiñonero-Candela et al., 2006). The main figure is that GP-SK outperforms the linear baseline but lags behind the SE one. This shows that non-linearities present in the data can not be captured by the GP-SK model. Since the string kernel is essentially a dot product over exponentially-sized vectors, it is not surprising that it is unable to capture non-linear behaviour. This gives us evidence that developing non-linear extensions of string kernels could be a promising avenue for future work.

### Inspecting hyperparameters

Probing the hyperparameters can give us insights about what kind of representation the kernel is learning. On Table 2 we show the values for one of the models that predict the emotion **surprise**. We can see that $\lambda_g$ has a very low value, while the $\mu$ values show a preference for subsequences up to 3 words. This lets us conclude that the kernel learned a text representation close to contiguous trigrams.

### 4 Related Work

String kernels were originally proposed for text classification (Lodhi et al., 2002; Cancedda et al., 2003) while recent work apply them for native language identification (Ionescu et al., 2014) and sentiment analysis (Giménez-Pérez et al., 2017), with promising results. Hyperparameter optimisation in these works is done via grid search and could potentially benefit from our proposed approach.

Gaussian Processes have been recently employed in a number of NLP tasks such as emotion analysis (Beck et al., 2014), detection of temporal patterns in microblogs (Preoiuc-Pietro and Cohn, 2013), rumour propagation in social media (Lukasik et al., 2015) and translation quality estimation (Cohn and Specia, 2013; Shah et al., 2013; Beck et al., 2016). These previous works encode text inputs as fixed-size vectors instead of working directly on the text inputs.

Among other recent work that aim at learning general structured kernels, the most similar to ours is Beck et al. (2015), who use GPs to learn tree kernels. Lei et al. (2017) unroll string kernel computations and derive equivalent neural network architectures. In contrast, our work put the learning procedure inside a GP model, inheriting the advantages of Bayesian model selection procedures. Nevertheless, many of their kernel ideas could be applied to a GP setting, which we leave for future work.

### 5 Conclusion

In this paper we provided the first steps in combining string kernels and Gaussian Processes for NLP tasks, allowing us to learn the text representations used by the kernels by optimising its hyperparameters in a fine-grained way. Experiments showed promising results in capturing text patterns that are not modelled by simpler baselines.

For future work, we plan to extend the model to account for non-linear representations, using approaches such as Arc-cosine kernels (Cho and Saul, 2009) and also applying the ideas from Lei et al. (2017). Another important avenue to pursue is to scale the model to larger datasets using recent advances in Sparse GPs (Titsias, 2009; Hensman et al., 2013). These in turn can enable richer kernel parameterisations not only for strings but other structures as well.

### Acknowledgements

Daniel Beck was supported by funding from CNPq/Brazil (No. 237999/2012-9) and from the Australian Research Council (DP160102686). Trevor Cohn is the recipient of an Australian Research Council Future Fellowship (project number FT130101105). The authors would also like to thank the anonymous reviewers for their comments.
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