COFFIN: A Computational Framework for Linear SVMs

Soeren Sonnenburg

SOEREN.SONNENBURG@TU-BERLIN.DE

Berlin Institute of Technology, Franklinstr. 28/29, 10587 Berlin, Germany Friedrich Miescher Laboratory, Max Planck Society Spemannstr. 39, 72076 Tübingen, Germany

Vojtěch Franc XFRANCV@CMP.FELK.CVUT.CZ

Czech Technical University in Prague, Technicka 2, 166 27 Praha 6, Czech Republic

Abstract

In a variety of applications, kernel machines such as Support Vector Machines (SVMs) have been used with great success often delivering stateof-the-art results. Using the kernel trick, they work on several domains and even enable heterogeneous data fusion by concatenating feature spaces or multiple kernel learning. Unfortunately, they are not suited for truly large-scale applications since they suffer from the curse of supporting vectors, i.e., the speed of applying SVMs decays linearly with the number of support vectors. In this paper we develop COFFIN — a new training strategy for linear SVMs that effectively allows the use of on demand computed kernel feature spaces and virtual examples in the primal. With *linear* training and prediction effort this framework leverages SVM applications to truly large-scale problems: As an example, we train SVMs for human splice site recognition involving 50 million examples and sophisticated string kernels. Additionally, we learn an SVM based gender detector on 5 million examples on low-tech hardware and achieve beyond the stateof-the-art accuracies on both tasks. Source code, data sets and scripts are freely available from http://sonnenburgs.de/soeren/coffin.

1. Introduction

Many applications in e.g. Bioinformatics, IT-Security and Text-Classification come with *huge* amounts (e.g. millions or billions) of training examples, which are indeed *needed* to obtain state-of-the-art results. At the same time predictions need to be made on billions of data points demanding

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for linear time algorithms that additionally can be effectively parallelized. Thus computationally highly effective methods are needed that can cope with ever growing data sizes. While classical kernel machines

$$f(\mathbf{x}) = \operatorname{sign}\left(\sum_{i=1}^{m} \alpha_i K(\mathbf{x}, \mathbf{x}_i) + b\right)$$
 (1)

often deliver state-of-the-art results, they are not suited for truly large scale applications since they suffer from the curse of supporting vectors, i.e. the number of non-zero coefficients α_i above. The total evaluation complexity to predict t elements, for m_{sv} support vectors and kernel complexity c is $\mathcal{O}(tm_{sv}c)$. Since $m_{sv} = \mathcal{O}(m)$, i.e., is linear in the training set size (Steinwart, 2004), kernel machines are in big-O notation (and for many practical applications) not at all faster than k-nearest neighbor: The number of training examples/support vectors m_{sv} becomes dominant, especially if kernel computations are already linear. Reduced set methods (Schölkopf & Smola, 2002) partially alleviate this problem by enforcing a low number of non-zero α_i in a post-processing step. Nevertheless, the computational complexity of determining a reduced support vector set and the potential performance degradation and the still prevailing prediction complexity render them infeasible for truly large-scale learning applications. Collobert et al. (2006) show that using non-convex loss function can largely reduce the number of support vectors, however, this is paid with more tricky optimization of a non-convex objective function. Keerthi et al. (2006) propose a greedy algorithm which simultaneously selects the set of support vectors and optimizes over the parameters α_i in (1). Again, this method optimizes non-convex cost and it is applicable to problems of moderate size only.

It is also operation (1) that slows down training in prominent SVM packages and potentially has caused a shift in interest from kernel SVMs back to linear SVMs for large-scale applications. Many of the recently proposed linear SVM solvers, are very efficient and guaranteed to converge to a ε -precise solution in $\mathcal{O}(m)$ (e.g., Joachims (2006)). In

this paper we develop a training strategy for linear SVMs that effectively allows the use of on-demand computed kernel like non-linearity and of virtual examples in the primal and thus leverages SVM applications to truly large-scale problems: As an example, we train a linear SVM on a gender classification dataset of almost 5 million images on a plain notebook with just 4GB of memory and on a bioinformatics splice site recognition task of 50 million examples using a 185 million dimensional string kernel feature space approximation to the traditionally spanned feature space of size $n > 10^{14}$. The paper is structured as follows: In Section 2 we introduce the COmputational Framework For IINear syms (COFFIN) which is a collection of simple ideas put into a coherent framework that can lead to dramatic performance improvements as we demonstrate in Section 3 where our proposed approach COFFIN is applied on two real-world data sets. Section 4 concludes the paper.

2. Leveraging Linear SVMs

Given labeled training examples $(\mathbf{x}_i, y_i)_{i=1}^m \in (\mathbb{R}^n \times \{-1, +1\})^m$ and a regularization constant C > 0, SVMs learn a linear classification rule $f(\mathbf{x}) = \operatorname{sign}(\langle \mathbf{w}, \mathbf{x} \rangle + b)$ by minimizing the following quadratic optimization problem

$$\min_{\mathbf{w},b} F(\mathbf{w}) := \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_{i=1}^m \max\{0, 1 - y_i(\langle \mathbf{w}, \mathbf{x}_i \rangle + b)\}.$$

Among the most prominent linear SVM solvers minimizing $F(\mathbf{w})$ are quasi-Newton methods (implemented in sub-BFGS; Yu et al. (2008)), (stochastic) subgradient descent algorithms (implemented in e.g. SGD Bottou & Bousquet (2008)), dual coordinate descent (implemented in Liblinear; Fan et al. (2008)) and cutting plane based algorithms (implemented in SVMPerf, BMRM and OCAS; Joachims (2006); Teo et al. (2007); Franc & Sonnenburg (2009)). For the latter four, convergence guarantees exist and they have been proven to have linear training time, i.e., achieve a ε precise solution in $\mathcal{O}(m)$ iterations.

Underneath, these algorithms are applied by supplying a set of sparse vectors \mathbf{x}_i and corresponding labels y_i as their input and optimize over a dense vector \mathbf{w} (and the bias b). In this work we strive to combine the flexibility of kernels with the computational efficiency of linear SVMs. In addition we aim at integrating virtually computed examples that were often shown to amend performance (in, e.g., digit recognition). This naturally requires code changes in the core of the participating SVM solvers and it is not obvious how to achieve this goal with the above considered linear SVM solvers. However, all of the above SVM solvers do not require direct access to elements of \mathbf{w} or examples \mathbf{x}_i , but merely require the following two operations:

(i) dot product between feature vector and the vector \mathbf{w} : $r \leftarrow \langle \mathbf{x}, \mathbf{w} \rangle$ **DOT**

(ii) multiplication with a scalar $\alpha \in \mathbb{R}$ and addition to the vector $\mathbf{v} \in \mathbb{R}^n$: $\mathbf{v} \leftarrow \alpha \mathbf{x} + \mathbf{v}$ ADD

To see this, recall that all SVM solvers mentioned above access the examples only to compute (i) outputs $f(\mathbf{x}_i) = \langle \mathbf{w}, \mathbf{x}_i \rangle$, $i = 1, \ldots, m$, and (ii) a sub-gradient $\mathbf{g} = \sum_{i \in I \subseteq \{1, \ldots, m\}} \pi_i \mathbf{x}_i$, $\pi_i \in \{-1, 0, +1\}$ of $F(\mathbf{w})$ (BMRM, SVM perf , OCAS) or its stochastic estimate (SGD). Note that the Hessian required by the quasi-Newton method (subBFGS) is also estimated via sub-gradients. In turn, **DOT** and **ADD** are sufficient operations, i.e., the only operations directly accessing the examples.

We propose a new training framework COFFIN whose main essence is the on demand computation of features and examples within the **DOT** and **ADD** operations. We show that by well organizing these computations the proposed strategy can significantly save both memory and computational demand which are the main hurdles in large-scale learning.

We describe two directions of the framework by introducing kernel like non-linearity (Section 2.1) and learning invariant classifiers using virtual examples (Section 2.2).

2.1. A Kernel Framework

To enable the use of non-vectorial data and kernels, we now consider $(\mathbf{z}_i, y_i)_{i=1}^m \in (\mathcal{Z} \times \{-1, +1\})^m$ as input and a feature extractor $\Phi: \mathcal{Z} \mapsto \mathbb{R}^n$ that maps objects **z** from the input space \mathcal{Z} to a real valued vectorial representation. While classical SVM optimizers operating in the dual can easily make use of the kernel trick $k(\mathbf{z}, \mathbf{z}') = \langle \Phi(\mathbf{z}), \Phi(\mathbf{z}') \rangle$, i.e., work without ever *explicitly* computing the mapping $\Phi(\mathbf{z})$, this is not straight-forward in the primal. Even though optimization over kernel expansion provides this trick also in the primal, it again leads to the curse of support vectors and hurts any large-scale learning applications. To endow kernel like non-linearity Φ is commonly applied in a preprocessing step. However, if $\dim(\mathcal{Z}) \ll n$ this quickly renders any kind of large-scale learning infeasible, since only few vectors x will fit in memory. In addition, it should be noted that such large objects will cause CPU cache misses whenever they are accessed slowing computations down significantly.

Computing Features on-demand We propose to use on-demand computed features, i.e., instead of applying the mapping $\Phi(\mathbf{z})$ in a preprocessing step we compute the non-zero elements of $\mathbf{x} := \Phi(\mathbf{z})$ on demand whenever \mathbf{x} is accessed. Formally, we define the non-zero elements

$$\Phi_{\neq 0}(\mathbf{z}) := \left\{ (\Phi_{\neq 0}(\mathbf{z}))_{v_1}, \dots, (\Phi_{\neq 0}(\mathbf{z}))_{v_\ell} \right\}$$

and their number as follows

$$|\Phi_{\neq 0}(\mathbf{z})| := \sum_{k=1}^{n} I(\Phi(\mathbf{x})_k \neq 0)$$

where I(.) is the indicator function that evaluates to 1 if its argument is true and to zero otherwise.

For the operations **ADD** and **DOT** to be efficient, it is required that (a) the individual features $\Phi_{\neq 0}(\mathbf{z})_{v_i}$ can be computed quickly, e.g. in $\mathcal{O}(1)$ (b) can be indexed by v_i , $i=1,\ldots,\ell$, (c) their subsequent access to $(\mathbf{w})_{v_i}$ is fast and (d) the number of non-zero features $|\Phi_{\neq 0}(\mathbf{z})|$ is low, i.e., optimally linear in the dimensions of the input

$$\mathcal{O}(|\Phi_{\neq 0}(\mathbf{z})|) = \mathcal{O}(\dim(\mathcal{Z})).$$

Examples for Φ are the construction of a (low-degree) polynomial kernel feature space on very sparse features (e.g., investigated for liblinear in Chang et al. (2010)), string kernel based features (n-gram counts), hashed feature values, decompression algorithms. They are described in more more detail in Section 2.1.2. We now discuss data structures that allow us to efficiently represent \mathbf{w} .

2.1.1. Data structures for representing w

Dealing with a potentially huge number of features, most of which potentially zero, requires an efficient representation of the SVM-w. In Sonnenburg et al. (2007a) we noted that there are three basic operations required when dealing with w, a clear operation to set all components of w to zero, an add operation that coincides with operation ADD in this paper and a lookup operation to access all non-zero elements efficiently and is required in the DOT operation. We can thus make use of their linadd trick to represent the SVM-w not necessarily as a dense vector, but if more appropriate in a sparse data structure like a tree or a sorted list. In addition, we will make use of hashing to lower the effective number of dimensions. Hashing has been first investigated in depth and successfully used in hash kernels (Shi et al., 2009).

We briefly review the data structures and their complexity:

Representing w as dense vector If n is not overly large then one should keep the whole vector w in memory. In this case each **ADD** and **DOT** operation can be done in $\mathcal{O}(|\Phi_{\neq 0}(\mathbf{z})|)$ time at a cost of a potentially huge, i.e., $\mathcal{O}(n)$ memory requirement. However, note that the dimensionality of w is independent of the number of examples m.

Sorted Array More memory efficient considering sparse data, but computationally more expensive are sorted arrays of index-value pairs $\{(v_1, w_{v_1}), \ldots, (v_\ell, w_{v_\ell})\}$. Assuming ordered tuples $(v'_k, (\Phi(\mathbf{x}))_{v'_k}), k = 1 \ldots, \ell'$ (indexed by \mathbf{v}'_k) **ADD** and **DOT** can be performed in $\mathcal{O}(\ell' + \ell)$.

Tree In particular when dealing with strings a way of organizing non-zero elements are trees, like binary trees, *tries* or suffix arrays (Knuth, 1973; Fredkin, 1960; Teo &

Table 1. Computational complexity of the **ADD** and **DOT** operations computed for a single **z** for the different data structures. In addition, the memory requirement of **w** is shown.

		Dense	Sorted Array	Tree
ĺ	Add	$\mathcal{O}(\Phi_{\neq 0}(\mathbf{z}))$	$\mathcal{O}(\mathbf{w} _{\neq 0}) + \Phi_{\neq 0}(\mathbf{z}))$	$O(\Phi_{\neq 0}(\mathbf{z}))$
				to $\mathcal{O}(d\Phi_{\neq 0}(\mathbf{z}))$
ĺ	Dot	$\mathcal{O}(\Phi_{\neq 0}(\mathbf{z}))$	$\mathcal{O}(\mathbf{w} _{\neq 0}) + \Phi_{\neq 0}(\mathbf{z}))$	$O(\Phi_{\neq 0}(\mathbf{z}))$
				to $\mathcal{O}(d\Phi_{\neq 0}(\mathbf{z}))$
	Mem	$\mathcal{O}(n)$	$\mathcal{O}(\sum_{i=1}^{m} \Phi_{\neq 0}(\mathbf{z}_i))$	$\mathcal{O}(\sum_{i=1}^{m} \Phi_{\neq 0}(\mathbf{z}_i))$

Vishwanathan, 2006). Depending on the tree used, the **ADD** operation needs $\mathcal{O}(d|\Phi_{\neq 0}(\mathbf{z})|)$ (trie; d is the depth of the tree) or $\mathcal{O}(|\Phi_{\neq 0}(\mathbf{z})|)$ (suffix array). Similar the complexity of **DOT** varies from $\mathcal{O}(d|\Phi_{\neq 0}(\mathbf{z})|)$ (trie) or $\mathcal{O}(|\Phi_{\neq 0}(\mathbf{z})|)$ (suffix array). Note that the computational complexity of both operations is independent of the number of d-mers/elements stored in the tree but comes at the cost of an additional storage overhead.

Hashing Table 1 summarizes computational complexity and memory requirements of the considered data structures in big-O notation. This unfortunately hides the large constants involved when dealing with the seemingly efficient trees. For example while $\mathcal{O}(\sum_{i=1}^m |\Phi_{\neq 0}(\mathbf{z}_i)|)$ seems like a low memory requirement (this quantity is linear in the amount of data), it is sufficient to already impose practical limits, e.g., Sonnenburg et al. (2007a) require 20 bytes per node for their already tuned DNA-tries; the highly memory-efficient suffix array algorithm of Teo & Vishwanathan (2006) still requires 19 bytes per character. The sorted array has an additional index attached to it increasing data size by at least factor 2.

On the other hand, **DOT** and **ADD** are very fast for dense w (no hidden large constants) but suffer from huge memory requirements (for some string kernels $n > 10^{14} \gg m$).

This is where hashing comes to the rescue: For an index set J, a number of bits γ , a hash function $h(J) \mapsto 1, \dots, 2^{\gamma}$ computes an approximation of $\Phi(\mathbf{z})_i$ via

$$(\widehat{\Phi}(\mathbf{z}))_j = \sum_{i \in J; h(i)=j} (\Phi(\mathbf{z}))_i$$

ignoring potential hash collisions, i.e., the new vector $\Phi(\mathbf{z})$ has only 2^{γ} dimensions. This trick and the resulting (minor) information loss has been extensively discussed in (Shi et al., 2009) w.r.t. both theory showing its influence on generalization bounds and empirically for dense w. It has the big advantage that we can use a fixed hash-table size of size $n=2^{\gamma}$ for w either in dense representation (Shi et al., 2009) or a sparse one (Sonnenburg et al., 2007a).

It should be noted that the use of such data structures is not exclusive either-or, for example for a string kernel one might want to use a dense representation for short string lengths and for the remaining use sorted arrays, suffix arrays or hashes. In this work we will exclusively focus on either explicit or hashed dense representations of \mathbf{w} since — for very large m — they have the lowest memory requirements and \mathbf{DOT} and \mathbf{ADD} can be computed fastest.

2.1.2. Computing Φ for a Variety of Kernels

In this section we give examples on how to efficiently compute Φ for a variety of kernels.

Polynomial Kernel of low degree The homogeneous¹ polynomial kernel of degree d is defined as $K(\mathbf{z}, \mathbf{z}') = (\langle \mathbf{z}, \mathbf{z}' \rangle)^d$, $\mathbf{z}, \mathbf{z}' \in \mathbb{R}^p$. The feature space of the polynomial kernel can be defined as $\Phi \colon \mathbb{R}^p \to \mathbb{R}^n$

$$\Phi(\mathbf{z}) = \left(\left(\frac{d}{u} \right)^{\frac{1}{2}} \mathbf{z}^{|\mathbf{u}|} \mid \mathbf{u} \in \mathbb{N}^p, |\mathbf{u}| = d \right),$$

where $\mathbf{u}=(u_1,\ldots,u_p)\in\mathbb{N}^p$ is a multi-index, $|\mathbf{u}|=\sum_{i=1}^p u_i, \begin{pmatrix} d\\u \end{pmatrix}=\frac{d!}{(d-|\mathbf{u}|)!\prod_{i=1}^p u_i!}$ and $\mathbf{z}^{|\mathbf{u}|}=\prod_{i=1}^p z_i^{u_i}$. The dimensionality of the feature space is $n(p,d)=\sum_{\mathbf{u}\in\mathbb{N}^p}I(|\mathbf{u}|=d)$. In turn, computing **ADD** and **DOT** operations require in general case $\mathcal{O}(n(p,d))$ operations and thus are feasible only if degree d is low and the input vectors \mathbf{z} are low-dimensional or very sparse. Let $J_{=0}(\mathbf{z})$ be a set of indices of zero components of \mathbf{z} and let $U_{\neq 0}(\mathbf{z})=\{\mathbf{u}\in\mathbb{N}^p\mid |\mathbf{u}|=d,u_i=0,i\in J_{=0}(\mathbf{z})\}$ be set of multi-indexes of non-zero monomials. Then, computing **ADD** and **DOT** require traversing only through the non-zero monomials $\mathbf{z}^{|\mathbf{u}|}$, $\mathbf{u}\in U_{\neq 0}(\mathbf{z})$. Hence the computation complexity of sparse **ADD** and **DOT** decreases to $\mathcal{O}(n(p-|J_{=0}(\mathbf{z})|,d))$. Note that one may save memory by using a hashed approximation of the multi-index $h(\mathbf{u})$.

Bag of Words, Spectrum and n-gram Kernel The spectrum kernel (e.g. Sonnenburg et al. (2007a)) implements the n-gram or bag-of-words kernel as originally defined for text classification in the context of biological sequence analysis. $\Phi_d(\mathbf{z})$ computes counts of all possible d-grams that are contained in the string \mathbf{z} , i.e., given an alphabet Σ and all possible d-grams $\mathbf{u} \in \Sigma^d$

$$\Phi_d(\mathbf{z}) = (\#\mathbf{u}_1(\mathbf{z}), \dots, \#\mathbf{u}_{|\Sigma^d|}(\mathbf{z})).$$

A flavor of this kernel considers all k-grams of length $1\dots d$, i.e. $\Phi_d^{wspec}(\mathbf{z}) = (\sqrt{\beta_1}\Phi_1(\mathbf{z}),\dots,\sqrt{\beta_d}\Phi_d(\mathbf{z})),$ where $\beta_k \in \mathbb{R}^+$ are some non-negative weights.

For small alphabets and d-gram lengths individual d-grams can be stored in fixed-size variables, e.g., DNA d-grams of length $d \le 8$ can be efficiently represented as 16-bit integer values. The ability to store d-grams in fixed-bit variables or even CPU registers greatly improves performance, as only a single CPU instruction is necessary to compare or index

a d-gram. The computational complexity of computing Φ is linear in the length of the sequences, i.e., $\mathcal{O}(|\mathbf{z}|_{\neq 0})$. Note that this representation allows efficient computation of the Weighted Spectrum kernel in $\mathcal{O}(d|\mathbf{z}|_{\neq 0})$ without requiring additional storage. Finally, note that for long strings \mathbf{z} or low d it can indeed be more efficient to store pre-processed tuples of $(\#\mathbf{u} \in \mathbf{z}, \mathbf{u})_{\forall \mathbf{u} \in \mathbf{z}}$ instead.

Depending on the alphabet size, the spectrum kernel is best represented explicitly, i.e., using a dense \mathbf{w} , index \mathbf{u} for small alphabets or a dense \mathbf{w} (Sonnenburg et al., 2007a), but hashed index $h(\mathbf{u})$ (Shi et al., 2009).

Weighted Degree Kernel The weighted degree kernel (Sonnenburg et al., 2007a) (WDK) can be conceived as a weighted spectrum kernel for each sequence position. This kernel has been excessively used to detect genomic signals (Sonnenburg et al., 2008) and its feature space can be expressed as

$$\Phi_d^{wd}(\mathbf{z}) = (\Phi_d^{wspec}(\mathbf{z}_{1,\dots,d}),\dots,\Phi_d^{wspec}(\mathbf{z}_{|\mathbf{z}|-d+1,\dots,|\mathbf{z}|})).$$

As a result the feature space of the WDK is $\mathcal{O}(l|\Sigma|^d)$ dimensional (Sonnenburg et al., 2007a) and thus for the usually considered d=20 even for relatively short DNA sequences too big for a dense representation. Previously, we used a sparse trie representation for Φ (Sonnenburg et al., 2007a). However, in this work we propose to use multiple dense hash tables instead, one hash table for each degree and position. For this to be efficient we require incremental or rolling hashes, i.e. hashes that can be seeded with the previous seed,

$$h(x_{1,\ldots,k+1},\sigma) = h(x_{1,\ldots,k+1},h(x_{1,\ldots,k},(\ldots,(h(x_1,\sigma)))),$$

where σ is some initial seed.

Similar to the spectrum kernel we can explicitly represent k-grams for small k to further speed up computations.

Other Examples Other examples for Φ include fast decompression algorithms like LZO² that can efficiently decompress a sequence at one third of the speed of a usual memcpy, but also other expert chosen general basis functions like sine waves, exponentials etc. While one could use the empirical kernel map or sparse kernel approximations to approximate general kernels the kernel evaluations with a subset of the training examples creates a huge speed penalty rendering on-the-fly computation of Φ hard.

2.2. Incorporating Invariance by Virtual Examples

In many applications, we know that there are transformations of the input measurements $z \in \mathcal{Z}$ which leave the class membership y invariant. A commonly used way to

¹The derivation for the inhomogeneous case is analogous.

²http://www.oberhumer.com/opensource/lzo/

incorporate prior knowledge into SVM classifiers is to augment the set of training examples with virtual examples (VE) that are created by applying a set of transformations (against which we want invariance) to the training examples (DeCoste & Schölkopf, 2002).

To put it formally, our prior knowledge is described by a set \mathcal{T} which contains a finite number of transformations $T \colon \mathcal{Z} \mapsto \mathcal{Z}$. We require that

$$f(\Phi(T\mathbf{z}_i)) = y_i$$
, $\forall T \in \mathcal{T}, i = 1, \dots, m$,

where $\{(\mathbf{z}_i, y_i)\}_{i=1}^m$ are given training examples. Training of f can be expressed as training of a standard SVM classifier from $|\mathcal{T}|m$ virtual training examples

$$\{(\mathbf{z}, y) \mid \mathbf{z} = T\mathbf{z}_i, y = y_i, T \in \mathcal{T}, i = 1, \dots, m, \}.$$

The VE method has two important advantages. First, it does not impose any constraints on the transformations \mathcal{T} . Second, existing SVM solvers can be used to train the invariant classifier. However, the cardinality of \mathcal{T} may increase exponentially when the transformation T is composed of s simpler ones, $T = T_1 \odot \cdots \odot T_s$ and thus $\mathcal{T} = \mathcal{T}_1 \times \cdots \times \mathcal{T}_s$. Thus, VE are computationally demanding because they (a) significantly increases the number of training examples and (b) pose huge memory requirements to store all $m|\mathcal{T}|$ virtual examples.

This is were COFFIN comes to the rescue: Instead of precomputing the VE in advance, we generate them on demand. Since only the original examples need to be stored in memory, this approach drastically reduces memory requirements. In case when transformations T can be computed quickly, the on demand generation of the virtual examples also speeds up the training. E.g., transformations of 2D images (needed in OCR and image recognition) can be computed on GPUs — a dedicated hardware for these transformations.

In Section 3.2, we demonstrate effectiveness of the proposed approach COFFIN on the problem of gender estimation from face images showing that COFFIN has by an order of magnitude less memory requirements compared to the standard approach. A practical outcome is that we can train the gender classifier from 4,808,250 example images on a notebook with only 4GB of memory instead of highend computing node with > 50GB of memory.

2.3. Implementation and Parallelization

We integrated COFFIN in the state-of-the-art cutting plane solver OCAS, the dual coordinate descent based LibLinear and SGD. We implemented a general framework that allows stacking of a variety of features that support **ADD** and **DOT** operations, namely dense and sparse real-valued features, weighted spectrum and WD kernel features for

specified k-mer length, once using an explicit representation and once using hashing. We implemented the virtual example method to OCAS solver as its API provides easy way to customize **ADD** and **DOT** operations.

Since the **DOT** operation is the most time consuming when performing predictions and when using batch-solvers we trivially parallelized this part of the code (based on shared memory parallelization, i.e., posix threads). However, an important detail here needs considerable attention: *memory locality*. CPUs are i/o bound, i.e. computation speed is drastically limited by memory speed and parallelized code cannot help this. To alleviate that bottleneck, off the shelf CPUs have rather large data and instruction caches. For example an AMD Opteron CPUs often have 64k level 1 data cache and 1MB level 2 data cache.³ Within the **DOT** operation, it is highly beneficial to split $\mathbf{w} = (\mathbf{w}_{B_1}, \dots, \mathbf{w}_{B_k})$ into smaller blocks, parallelizing within each block $\mathbf{r} = \sum_{j=1}^k \sum_{i=1}^t \langle \mathbf{x}_{i,B_j}, \mathbf{w}_{B_j} \rangle \mathbf{e}_i$ where the inner sum is distributed among cores.

3. Experiments

3.1. Human Acceptor Splice Site Recognition

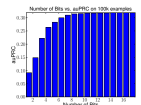
To demonstrate the effectiveness of our proposed method COFFIN, we apply it to the problem of human acceptor splice site detection. This problem can be formulated as a two-class classification problem discriminating true splice sites from fake ones. Due to the importance of this problem in computational gene finding, many different methods to detect splice sites have been proposed. They all predict splice sites based on the local context, i.e., a short window around the actual splice site. Currently, support vector machines are the most accurate splice site detectors (Degroeve et al., 2005; Sonnenburg et al., 2007b; Franc & Sonnenburg, 2009). In particular, in (Sonnenburg et al., 2007b) we have shown that prediction accuracy steadily increases with training sample size. However, even though we already used the linadd algorithm (Sonnenburg et al., 2007a) to speed up string kernel-based SVMs on a quad-core system, we could not use all available 50 million training points (but "only" 8 million).

Degroeve et al. (2005) trained a linear SVM based on a number of pre-selected and explicitly computed string kernel feature spaces that are subsets from the spectrum and WD kernel feature spaces: Left and right of the splice site spectrum kernels of order 3 up to order 6 were used. Over the whole window, a WD kernel of order 3 with weights equal to 1 was used. Even though this approach scales well, they used < 100,000 data points (potentially, since they relied on the unmodified SVM $^{\rm light}$ binary).

http://en.wikipedia.org/wiki/Opteron

Table 2. Training times and auPRC for human splice site detection for various data set sizes and w representations and d's of the weighted degree kernel. (first row) The previous state-of-the-art was an SVM employing linadd and a weighted degree shift kernel (WDS). (following rows) COFFIN employing OCAS (OC) and liblinear (LL) is compared to linadd and by far outperforms linadd in accuracy and speed when using hashing. Note that previous experiments were re-run on the same machine to aid a direct comparison.

Method / Sample Size	10^{4}		10^{5}		10^{6}		10^{7}		$5 \cdot 10^{7}$	
SVM ^{light} +linadd WDS	65s	7.14%	1114s	26.37%	24861s	43.80%	2112112s	53.01%	-	
SVM^{light} +linadd $d=8$	57s	10.37%	970s	28.62%	34110s	43.78%	-		-	
${ m SVM}^{light}$ +linadd $d=20$	56s	11.15%	1033s	31.80%	34586s	46.27%	-		-	
COFFIN OC $d = 8$ (dense)	167s	10.00%	948s	28.57%	9952s	43.84%	131202s	53.26%	909105s	57.78%
COFFIN OC $d = 20$, $\gamma = 12$	65s	10.81%	435s	31.80%	5349s	46.12%	76311s	54.31%	908654s	57.89%
COFFIN LL $d = 20$, $\gamma = 12$	61s	10.59%	360s	31.59%	3783s	45.97%	25902s	54.17%	132581s	57.75%
COFFIN LL $d = 20$, $\gamma = 16$	111s	10.52%	604s	31.69%	4590s	46.26%	44232s	54.56%	247907s	58.57%



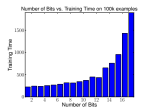


Figure 1. Performance in terms of auPRCand training times on the human acceptor splice site experiment using 100,000 examples and varying bit sizes for the hash of the central WD kernel. For this experiment OCAS was used. It can be seen that already starting with about 8-10 bits the auPRCreaches a plateau. In addition, training times start to drastically increase as soon as hashes of more than 12 bit are used. This drop in performance indicates that the whole hash-table does no longer fit in the CPU data cache for larger hash tables. For 12bits w is 11,725,480 dimensional.

Recently, we could demonstrated in a proof of concept study for OCAS (Franc & Sonnenburg, 2009) that training on all the available examples in richer feature spaces improves performance. However, we could not use the full potential of higher order string kernels and achieved inferior performance compared to (Sonnenburg et al., 2007b) for the same sample size (cf., COFFIN OC d=8 vs. Linadd d=20 WDS in Table 2).

Experimental Setup We reproduced the result of (Sonnenburg et al., 2007b) on the same computer node and following (Franc & Sonnenburg, 2009), we train COFFIN on all available 50 million strings of length 141 using the features corresponding to two weighted spectrum kernels (one left and one right of the splice site, i.e., positions 1-59 and 62-141) and a WD kernel. We fixed C=1 and used the weighted spectrum kernel order d=8, for the WD kernel order d=8 or d=20 respectively, which were found optimal in (Franc & Sonnenburg, 2009). We use a dense explicit 174, 760 dimensional representation for the spectrum kernels and dense or hashed representations for the WD kernel for hash sizes $\gamma=12$ and $\gamma=16$ (cf., Fig. 1 for a discussion about optimal hash sizes). The resulting spanned string kernel feature space has 12, 495, 340 (WD

d=8 explicit), 11,725,480 (WD d=12 hash $\gamma=16)$ and 184,986,280 (WD d=20 hash $\gamma=16)$ dimensions respectively.

As the raw string-based dataset is already of size $7.1 \cdot 10^9$ bytes and even a sparse representation of each string increases the dataset by a factor of more than 3,000 ((141 + 59+80)·12 bytes per feature vector, assuming a 4 byte integer and an 8 byte float), it is only by the means of COFFIN that we can solve such huge optimization problems.

To provide a fair comparison we measure training times and auPRC on a held out test set of 4,627,840 examples for various training set sizes and explicit representations or higher order ones with hashing. We consider SVM^{light} employing linadd and OCAS and liblinear employing COFFIN in this comparison.⁴

Using COFFIN within liblinear, training on 50 million examples using a single CPU-Core of a 16 core AMD Opteron Linux-based machine, leads to record area under the precision recall curve (auPRC) of 58.57% in less than 3 days. For comparison, the previous best dual method already using the linadd speedups (Sonnenburg et al., 2007a) achieved an auPRC of 53.01% on 10^7 examples in about 24 days. Using OCAS (Franc & Sonnenburg, 2009) we achieved an auPRC of 57.77%. With COFFIN we obtain the same precision in *one third* of the time (cf., Table 2, row 1 (linadd), 4 (COFFIN OC) and 7 (COFFIN LL)). We could not train OCAS on the $\gamma=16$ hashed data set since a single cutting plane already requires about 1.4GBof memory. However, since OCAS is a batch method, we could train it using 16 CPU cores on 50 million examples employing parallelized **DOT** operation resulting in 19hours spend in computing outputs instead of 252 (speedup factor 13) demonstrating the effectiveness of our memory access pattern. Since liblinear is an online style solver, it cannot easily be parallelized but could benefit from the recent work of M. Zinkevich (2009) on delayed gradient updates. Liblinear with d=20, $\gamma=16$ involves a feature space of

 $^{^4}$ Preliminary results showed that SGD failed miserably — potentially requiring step-size tuning for the different d.

size 184,986,281. Training times for COFFIN employing liblinear and a L2 loss were slightly lower at the cost of slightly decreased performance (results not shown).

3.2. Gender Estimation From Face Images

The task of this binary classification problem is to discriminate digital images into two classes – males and females. A robust classifier should be invariant against common image transformations: translation, rotation, scale and illumination changes. Currently, there exist feature representations invariant only against one of the transformations. For

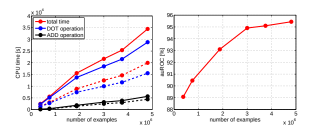


Figure 2. Performance of the gender classifier trained from virtual examples. The left figure shows CPU time needed by **DOT** operation (blue), **ADD** operation (black) and the total training time (red). The solid lines correspond to COFFIN's on demand computation strategy while the curves for standard approach are dashed. Because all precomputed examples do not fit into 4GB RAM, the times for the standard approach (dashed) are estimated. The right figure shows auROC w.r.t. training set size.

example Local Binary Patterns (LBP) (Ojala et al., 2002) are the current state-of-the-art image descriptors thare are invariant against any monotonic change in intensity values. In order to gain robustness against translation, rotation and scale, we apply the method of virtual examples (c.f. Sec. 2.2) in training. As mentioned, the bottle neck of the method is that the set of virtual examples quickly grows very large, imposing artificial memory limits. We show that using COFFIN computing virtual examples on demand significantly alleviates the memory problem at the price of only minor increase in training time. In particular, we train the gender classifier from 4,808,250 virtual examples on a notebook with Intel 2.66 GHz CPU and only 4GB of memory. Storing all the precomputed examples in memory would require more than 50GB.

We collected a dataset of 18,504 images with human faces downloaded from the Internet. We split the images into 12,822 training and 5,682 test examples. The faces were manually segmented and labeled with the gender. The segmentation is given by a position and size of a window containing a face. When applying the classifier in "real world", position and size of the window are taken from a pre-trained face detector and are thus often imprecise. To cope with this, we cropped the testing faces using the an-

notated segmentation applying translational and rotational distortions: The position distortion was in range of ± 2 pixels in both axis and the distortion in scale was $\pm 5\%$ of the base window size. The parameters of the distortions were estimated from outputs of a real AdaBoost face detector.

We generate the virtual examples by applying the mentioned transforms to the annotated images. The classifier input is a quadruple $\mathbf{x}=(\mathbf{I},\mathbf{p},s,\varphi)$, where $\mathbf{I}\in\mathbb{N}^{90\times60}$ is gray-scale image, $\mathbf{p}\in\mathbb{N}^2$ is position of the 60×40 pixels base window, $s\in\mathbb{R}$ is scale and $\varphi\in\mathbb{R}$ is the rotation of the base window. We define the transformation $T(\Delta\mathbf{p},\Delta s,\Delta\varphi)(\mathbf{I},\mathbf{p},s,\varphi)=(\mathbf{I},\mathbf{p}+\Delta\mathbf{p},s+\Delta s,\varphi+\Delta\varphi)$ parametrized by the triplet $(\Delta\mathbf{p},\Delta s,\Delta\varphi)$ which defines the change in translation, scale and rotation, respectively. Then we construct the set $\mathcal{T}=\{T(\Delta\mathbf{p},\Delta s,\Delta\varphi)\mid \Delta p\in\mathcal{T}_{\varphi}\}$ where

$$\begin{array}{rcl} \mathcal{T}_p & = & \{\Delta \mathbf{p} \mid \Delta \mathbf{p} = (u; v), u, v \in \{-2, -1, 0, 1, 2\}\} \\ \mathcal{T}_s & = & \{\Delta s \mid s \in \{-0.05, 0, 0.05\}\} \\ \mathcal{T}_\varphi & = & \{\Delta \varphi \mid \Delta \varphi \in \{-6, -3, 0, 3, 6\}\} \end{array}$$

i.e., for each training image we generate $|\mathcal{T}| = 5^2 \cdot 5 \cdot 3 = 357$ virtual examples.

The feature representation $\Phi(\mathbf{I},\mathbf{p},s,\varphi)\in\mathbb{R}^n$ is computed from responses of the LPB filter on pyramidal representation of the base window that is cropped from image \mathbf{I} according to (\mathbf{p},s,φ) . The pyramid is composed of images $60\times 40,\ 30\times 20,\ 15\times 10,\ 7\times 5$ pixels, obtained from the base window which was then three times downscaled by factor 2. It results in n=723,712-dimensional sparse feature vector composed of all zeros and 2,827 coordinates equal to one. The feature vector is most efficiently represented by storing 2,827 indexes (4 bytes each) of non-zero coordinates, i.e., we need $2,827\cdot 4=11,308$ bytes per example. Hence pre-computing all $357\cdot 12,822=4,808,250$ virtual examples requires ≈ 51 GB.

We alleviate the memory problem by computing the virtual examples on demand. For each training image, we pre-compute only their rotated and scaled versions because these are the most expensive operations and they still fit to memory; to store $12,822\cdot 5$ (rotation) $\cdot 3$ (scale) = 192,330 images we need 1.3 GB. The image translations and the pyramid of LBP features are computed on demand. Note that translating and downscaling an image by 2 can be computed very efficiently. By using COFFIN we need 25 times less memory to store the training examples compared to the standard approach when the features are pre-computed. In particular we need 1.3GB instead of 51GB.

For different training set sizes, $m \in 375 \cdot \{1000, 2000, \dots, 10000, 12822\}$, we trained SVMs using OCAS and measured computation time and accuracy. The results presented further are obtained for C=0.001 which was found by tuning on the original

training examples. Figure 2 (left) shows the time for computing outputs (dominated by **DOT** operation), the time for computing cutting planes (**ADD** operation) and the total time. Figure 2 (right) shows the accuracy, measured in terms of the area under ROC (auROC, Fawcett (2003)), w.r.t. number of examples. It is seen that generating more virtual examples significantly helps the performance at a marginal increase of training time. Without COFFIN , we could only fit $\approx 400,000$ precomputed examples into 4GB of memory. COFFIN enables training on over *four million* examples increasing auROC from 89.57% (on the original 12, 822 examples; $\approx 90\%$ auROC with 400,000 examples) to an auROC of 95.44% obtained with all virtual examples.

4. Conclusion

We have presented COFFIN — a very efficient computational framework for on-demand creation of features and virtual examples. COFFIN combines the computational efficiency of linear SVMs with the flexibility of kernel based learning. In the experimental section we have demonstrated (a) that our approach allows efficient computations even on low-budget hardware and (b) leads to state-of-the-art results solely due to the fact that it enables the use of all available training data. For example we could train a linear SVM on about 5 million example images for the task of gender recognition on a standard notebook with just 4GB of memory and a linear SVM for human acceptor splice site recognition on 50 million examples in a more than 184 million dimensional feature space in less than 3 days achieving new record performance. Still, we see further potential in improving our approach, by considering the computational costs to create virtual vectors $T(\mathbf{x})$ or features $\Phi(\mathbf{z})$ respectively in the core optimization process of the linear SVM solver. For example computed elements could be cached and solvers could put focus on optimizing for the few cached elements before tuning the rest.

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- ⁵ A principled caching strategy might bring speed of both approaches up to par.

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