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Sequential Data Selection for Predicting the Pathogenic Effects of Sequence Variation

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Abstract—Recent improvements in sequencing technologies provide unprecedented opportunities to investigate the role of genetic variation in human disease. In previous work we have proposed a machine learning approach to predicting whether single nucleotide variants (SNVs) are functional or neutral in human disease. Many data sources from the Encyclopaedia of DNA Elements (ENCODE) may be relevant to this problem. To integrate these data sources, we applied integrative multiple kernel learning (MKL) that weights each source according to its relevance. Using an MKL optimization that yields sparse weights, we were able to eliminate the least informative data sources from our model. However, when selecting from a wide assortment of data sources, we have found that MKL may not be an efficient method for eliminating uninformative sources. Many data sources related to the human genome are incomplete: this can reduce dramatically the data available for training and the proportion of novel predictions that exploit all relevant sources. Here we introduce a greedy sequential selection method that assesses data sources in a structured fashion prior to MKL weight optimization. This method allows us to eliminate a majority of uninformative data sources prior to assigning kernel weights. When we use this method with our coding-region predictor, we select just five kernels for our final model, yielding increased accuracy over our previous model. In addition, by reducing the amount of data required for novel predictions, we are able to increase by five fold our model’s coverage for new predictions.

I. INTRODUCTION

The introduction of fast and inexpensive sequencing technologies is providing many new insights into the role of genetic variation in human disease. In this work we consider single nucleotide variants (SNVs) in the human genome. Predicting which of these are functional, as against neutral, promises to improve our understanding of the molecular mechanisms underpinning human disease. In a recent study we proposed a novel algorithmic approach to predicting the functional consequences of both coding and non-coding SNVs (FATHMM-MKL) [12]. Our approach uses integrative multiple kernel learning (MKL), a method that learns to weight different types of data according to their relative informativeness. In our work we use SimpleMKL [9], an MKL implementation that uses an $L_1$ norm to yield sparse solutions that implicitly exclude data sources by assigning them zero weights. In our previous study, our predictor for non-coding SNVs outperformed competing methods using just 4 out of 10 data sources [12]. Our coding-region predictor, using all 10 data sources, matched competitors’ performance when all data were available, but its performance suffered when data were missing from some sources.

With MKL, different types of input data are encoded into kernel matrices that quantify the similarity of data objects. A number of different methods have been proposed for deriving kernel matrices for different types of data objects, including data with discrete and continuous values, sequence data and graph data [10]. With MKL, each constituent data type is encoded into a corresponding base kernel $K_\ell$ (where $\ell = 1,\ldots,p$ if there are $p$ feature groups), from which we can derive a composite kernel matrix $K = \sum_{\ell=1}^{p} \lambda_\ell K_\ell$, where $\sum_{\ell=1}^{p} \lambda_\ell = 1$ and $\lambda_\ell \geq 0$. The $\lambda_\ell$ are kernel weights. This aggregate kernel can then be used with a kernel-based classifier, such as a Support Vector Machine (SVM) [1], which was the classifier used here.

Suppose the training set for a Support Vector Machine consists of vectors $x_i$ with associated labels $y_i = \pm 1$. The index $i$ labels the training example $(x_i, y_i)$ and we will assume there are $m$ such training examples in the training set. During the training process for the SVM, the learning parameters $\alpha_i$ are found by maximizing the following convex (quadratic) objective function in $\alpha_i$ [1]:

$$W(\alpha) = \sum_{i=1}^{m} \alpha_i - \frac{1}{2} \sum_{i,j=1}^{m} \alpha_i \alpha_j K(x_i, x_j)$$  \hspace{1cm} (1)

subject to the linear constraints:

$$\alpha_i \geq 0 \quad , \sum_{i=1}^{m} \alpha_i y_i = 0$$  \hspace{1cm} (2)

Suppose $\alpha_i^*$ are the values of the training parameters at the optimum of the objective function stated in (1). From the $\alpha_i^*$ it is then straightforward to find the bias, $b^*$, or offset in the decision function via:
For the binary decision function of an SVM, the predicted label of a novel input $z$ is then decided by the sign of $\phi(z) = \sum_{i=1}^{m} \alpha_i^* y_i K(x_i, z) + b^*$. With a composite kernel of the form $K(x_i, x_j) = \sum_{\ell=1}^{P} \lambda_{\ell} K_{\ell}(x_i, x_j)$ substituted into (1) we see that the learning process now involves optimization of a linearly constrained linear program in the kernel weights $\lambda_{\ell}$ and a linearly constrained quadratic program in the learning parameters $\alpha_i$. This is a tractable problem in optimisation theory and can be approached via a wide variety of methods [5], including semi-definite programming or quadratically constrained linear programming (QCLP), for example.

A variety of methods have been proposed for MKL [5] and this approach has been successfully demonstrated with various classification problems in bioinformatics, which use different types of input data e.g. [15]. By using all available data encoded into a set of kernels, MKL classifiers most frequently outperform a single kernel classifier constructed for one type of data. In addition, the kernel weights are adjusted according to the relative informative-ness of the different types of data: this enhances overall performance and interpretation of the model. In its simplest form, all weights in a composite kernel are the same ($\lambda_{\ell} = \frac{1}{P}$), a form we call an unweighted aggregate. When all constituent data sources are at least somewhat informative, an unweighted aggregate may perform as well as one with fully-trained kernel weights. However, when there is disagreement between kernels, performance may decline substantially if uninformative kernels outnumber informative ones. This behavior allows us to evaluate the potential impact of each set of kernels prior to optimizing kernel weights.

MKL optimization methods rely on the assumption that data are available for every kernel for every training example. However, as ENCODE is an ongoing project to annotate the human genome, many data sources are incomplete. As we add data sources it can become increasingly difficult to use MKL to select the most informative ones, as MKL requires training examples common to all of them (see Figure 1). This same restriction can impact novel predictions when data are missing for some sources. In previous work we addressed this by re-weighting the remaining kernels [12] but this may yield lower accuracy than a full-featured prediction (Figure 2). Accordingly, we have developed a novel, greedy approach that pre-selects data sources according to the accuracy of their corresponding kernels. We assess each kernel using cross-validation (CV) and rank the data sources according to their accuracy on a validation set. We then build unweighted aggregate models starting with the two most accurate kernels and sequentially adding the next-lowest-ranked kernel until the aggregate model’s performance on the validation set reaches a plateau or declines. Our results suggest that this approach can yield state-of-the-art predictors that dramatically increase the proportion of full-featured predictions.
must integrate data from diverse sources and identify the most informative of them. Our FATHMM-MKL method used integrative MKL to weight kernels constructed from different data sources, where zero weights implicitly exclude the least informative sources. However, this may not be a practical way to handle a large number of datasets: as we add new datasets, training an MKL model becomes difficult because many examples are not represented in all data sources. For example, in our previous work only 2,146 out of 87,518 training examples (2.5%) were represented in all 10 of the datasets we used [12]. In addition, a parsimonious model that uses a few well-chosen data sources may generalize better than a model constructed from many sources.

To assess each data source, we train a single-kernel support vector machine (SVM) and compute its average accuracy in 5-fold CV. By isolating each data source in this manner, we can assess its performance across the full spectrum of SNVs represented in the data. To make results comparable between data sources, we use balanced sets of 1,000 positive (pathogenic) and 1,000 negative (neutral) examples in each case. We then rank the data sources according to their average CV accuracy on a validation set.

To identify the data sources we will include in our final model, we construct an unweighted aggregate model using the two top-ranked data sources and record its accuracy on a validation set. We build subsequent models by adding data sources in descending order of accuracy, constructing an aggregate for each combination of data sources. For each combination we establish separate training, validation and test sets: a training set to train the model; a validation set to determine optimal parameters and to record accuracy, and a test set we leave aside to test the final MKL model. We select as our final combination the data sources associated with the unweighted aggregate where accuracy on validation data reaches a plateau or declines. At this point we optimize MKL weights and evaluate the final model on the test set.

### III. Results

#### A. Tests on original data

To assess our proposed method, we compared our original FATHMM-MKL coding-region classifier with models composed of two to ten component kernels, using the same datasets as the original model. Two of these datasets, 100-way conservation and 46-way conservation, were constructed from FATHMM [11], PhastCons [13] and Phylop [7] scores, while the remaining data were downloaded from ENCODE [3] (for more details on the datasets and how they were selected, see [12]). These data consist of ten feature groups, with an intersection set of 7,597 examples (2,218 positive and 5,379 negative). To evaluate each component kernel, we performed nested 5-fold CV on balanced sets of 1,774 positive/1,774 negative training examples and 443 positive/443 negative test examples. We used the same training and test examples for all kernels to ensure that we could compare CV statistics between them. The test sets were put aside to test MKL models from the same training data, while the training data were used to evaluate individual kernels. Within each fold, we randomly split the training data into training and validation subsets (80% training and 20% validation). We used the validation set to establish optimal kernel parameters and to determine the maximum accuracy (per fold) for each kernel. We then averaged each kernel’s accuracy across folds to yield a kernel ranking (Table I).

![Table I](image)

<table>
<thead>
<tr>
<th>Rank</th>
<th>Feature group</th>
<th>Source</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>100-way conservation</td>
<td>FATHMM</td>
<td>0.812</td>
</tr>
<tr>
<td>2</td>
<td>46-way conservation</td>
<td>FATHMM</td>
<td>0.805</td>
</tr>
<tr>
<td>3</td>
<td>TFBS (Peak-Seq)</td>
<td>ENCODE</td>
<td>0.698</td>
</tr>
<tr>
<td>4</td>
<td>Histone (ChIP-Seq)</td>
<td>ENCODE</td>
<td>0.685</td>
</tr>
<tr>
<td>5</td>
<td>TFBS (SPP)</td>
<td>ENCODE</td>
<td>0.655</td>
</tr>
<tr>
<td>6</td>
<td>Open chromatin (DNase-Seq)</td>
<td>ENCODE</td>
<td>0.623</td>
</tr>
<tr>
<td>7</td>
<td>Open chromatin (FAIRE)</td>
<td>ENCODE</td>
<td>0.575</td>
</tr>
<tr>
<td>8</td>
<td>Genome segmentation</td>
<td>ENCODE</td>
<td>0.562</td>
</tr>
<tr>
<td>9</td>
<td>DNA footprints</td>
<td>ENCODE</td>
<td>0.555</td>
</tr>
<tr>
<td>10</td>
<td>GC content</td>
<td>ENCODE</td>
<td>0.552</td>
</tr>
</tbody>
</table>

Next we constructed unweighted aggregate models from two or more component kernels and compared their performance to the original FATHMM-MKL on the same test examples. For $k = 2, \ldots, 10$ kernels, we constructed $k$-kernel aggregates using the top $k$ component kernels according to the rankings shown in Table I. All of the unweighted aggregate models performed better than any of their component kernels, and at $k = 5$ we observed a nominal peak in performance, where the unweighted aggregate performed nearly as well as the original FATHMM-MKL (Table II and Figure 3). The $k$-kernel aggregate models were trained on 80% of the data and tested on the remaining 20% for five folds, hence each was trained using less data than FATHMM-MKL and none had prior exposure to the test examples. Despite this arguably unfair comparison, the strong performance of these models and the consistent accuracy for models with $k = 5, \ldots, 10$ suggests that we can use fewer datasets than the original model without sacrificing performance.

#### B. Constructing a new model

For this study we obtained 57,276 pathogenic SNVs from HGMD and identified 109,667 neutral (presumed benign) SNVs from the 1,000 Genomes database for a total of 166,843 SNV examples. We were able to generate FATHMM scores (100-way conservation and 46-way conservation) for all of these examples. Within the 12 additional datasets we selected from ENCODE, we found just one database (Mappability) with all of these examples, while DNA footprints had just 15,399 (Table III). For our greedy aggregation procedure, we used the data common to all 14 datasets. When we combine all 14 datasets, our training data consists of 4,849 SNVs common to all of them, including 1,300 pathogenic and 3,549 neutral
**TABLE II**

<table>
<thead>
<tr>
<th>Model</th>
<th>Acc.</th>
<th>ROC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Original</td>
<td>0.846</td>
<td>0.912</td>
</tr>
<tr>
<td>2-kernel</td>
<td>0.820</td>
<td>0.899</td>
</tr>
<tr>
<td>3-kernel</td>
<td>0.831</td>
<td>0.892</td>
</tr>
<tr>
<td>4-kernel</td>
<td>0.828</td>
<td>0.882</td>
</tr>
<tr>
<td>5-kernel</td>
<td><strong>0.833</strong></td>
<td><strong>0.907</strong></td>
</tr>
<tr>
<td>6-kernel</td>
<td>0.829</td>
<td>0.889</td>
</tr>
<tr>
<td>7-kernel</td>
<td>0.831</td>
<td>0.903</td>
</tr>
<tr>
<td>8-kernel</td>
<td>0.831</td>
<td>0.901</td>
</tr>
<tr>
<td>9-kernel</td>
<td>0.832</td>
<td>0.901</td>
</tr>
<tr>
<td>10-kernel</td>
<td>0.832</td>
<td>0.901</td>
</tr>
</tbody>
</table>

Performance of unweighted aggregate kernels on original training data. Shown are the prediction accuracy and ROC score for the original version of FATHMM-MKL (Original) and for aggregates consisting of up to 10 kernels on the same test data.

**TABLE III**

<table>
<thead>
<tr>
<th>Rank</th>
<th>Feature group</th>
<th>Examples</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>100-way conservation</td>
<td>166,843</td>
<td><strong>0.819</strong></td>
</tr>
<tr>
<td>2</td>
<td>46-way conservation</td>
<td>166,843</td>
<td><strong>0.785</strong></td>
</tr>
<tr>
<td>3</td>
<td>TFBS (Peak-Seq)</td>
<td>63,106</td>
<td><strong>0.706</strong></td>
</tr>
<tr>
<td>4</td>
<td>Histone (ChIP-Seq)</td>
<td>150,882</td>
<td><strong>0.681</strong></td>
</tr>
<tr>
<td>5</td>
<td>TFBS (uniform)</td>
<td>48,882</td>
<td><strong>0.673</strong></td>
</tr>
<tr>
<td>6</td>
<td>TFBS SPP</td>
<td>32,494</td>
<td><strong>0.652</strong></td>
</tr>
<tr>
<td>7</td>
<td>Genome segmentation</td>
<td>166,703</td>
<td>0.601</td>
</tr>
<tr>
<td>8</td>
<td>Open chromatin (DNase-Seq)</td>
<td>79,540</td>
<td>0.600</td>
</tr>
<tr>
<td>9</td>
<td>DNA uniform</td>
<td>79,219</td>
<td>0.586</td>
</tr>
<tr>
<td>10</td>
<td>Open chromatin (FAIRE)</td>
<td>48,505</td>
<td>0.573</td>
</tr>
<tr>
<td>11</td>
<td>Riken CAGE</td>
<td>81,004</td>
<td>0.566</td>
</tr>
<tr>
<td>12</td>
<td>GC content</td>
<td>164,656</td>
<td>0.553</td>
</tr>
<tr>
<td>13</td>
<td>Mappability</td>
<td>166,843</td>
<td>0.496</td>
</tr>
</tbody>
</table>

The full set of feature groups considered for our new model, shown in descending order of accuracy. Accuracy was determined by selecting from the data available for each data set, as opposed to the cross-section used in our original model. The resulting performance, along with new data sources (italics) changed the rankings for some feature groups. Using our greedy selection method, we identified six data sets (bold) as likely to be the most informative.

For each of these datasets we performed nested 5-fold CV using balanced sets of 1,000 pathogenic and 1,000 neutral SNV examples and recorded the average accuracy. For each kernel we used the same examples in each fold, to ensure that the accuracy results were comparable.

As described in Section III-A, we optimised kernel parameters for each fold by splitting the training data into training and validation subsets. We did not find that any of the kernels were linearly separable, so we searched C-values from $10^{-3}$ to $10^3$ and selected the value that yielded the highest accuracy and still permitted convergence. We then computed average 5-fold CV accuracy for the validation sets to establish kernel rankings. These new rankings are similar to those in our first test (Table III): the two datasets based on FATHMM conservation scores yielded the strongest performance, with accuracy up to 84%. For the ENCODE datasets accuracy ranged from 50% for Riken CAGE to 68% for TFBS (Peak-Seq). The new ENCODE datasets altered the rankings slightly, but only TFBS (Uniform), performed well enough to appear in the top half of the rankings.
for $k = 2$ to 82% for $k = 6$, but declined after that. We also see a sharp decline in individual kernel performance between the sixth and seventh ranked kernels (Table III, TFBS SPP and Genome segmentation), so we selected the top six kernels for MKL optimization. Note that the unweighted aggregates may not perform as well as their best constituent kernels, as conflicting scores from constituent kernels may cancel each other out. However, the model can realize substantial gains once optimum weights have been learned.

Two of these kernels are associated with FATHMM conservation scores and represent all available training data. Missing data in the remaining sets reduced our overall training set to 28,998 examples—still vastly more than we had for our original study. We used SimpleMKL to establish kernel weights for each fold, using only the training and validation sets. For each kernel we then used the average weight as the weight for the final model (Figure 5). Note that the optimized weights do not track the relative rankings very closely. This is likely due to redundancies between the data sets: the 100-way conservation and 46-way conservation scores are closely related, so the 46-way conservation weight is considerably lower than we might expect given the relatively small difference in their kernel accuracies (Table III). Similarly, one of the three TFBS kernels, TFBS (SPP) received zero weight, allowing us to eliminate that source. In turn, this increased the number of examples available to train the final model, to 41,476 examples. This also improved coverage for novel predictions, from under 5% for the original classifier to 24.9% for the new model, a five-fold improvement.

To test our final model, we ran 5-fold CV using a balanced set of 2,000 pathogenic and 2,000 neutral examples. We compared these predictions with those of the original FATHMM-MKL and two other state-of-the-art methods, CADD [6] and DANN [8] (Figure 6). Our new model yields substantial improvements over our previous model (Figure 6, top), likely due to the additional training data now available. All of the top competitors yield similar performance (Figure 6, top): the newest version of CADD (v1.3) is the best of these competitors (AUC 0.90) while our new model yields the top AUC score of 0.91. While these results do not suggest a clear winner, they demonstrate that our new model provides accuracy that is competitive with an ever-improving state of the art. In addition, we found that we could obtain scores for all five of our datasets across 24.9% of coding regions in the human genome, a dramatic improvement over the severely restricted coverage we obtained when using a 10-kernel model.

Fig. 5. We used SimpleMKL to optimize kernel weights for the final model. Weights were highly consistent across the five folds, as indicated by the error bars. This result helped us eliminate an additional data set, TFBS (SPP). Note that although the 46-way conservation data are ranked higher than TFBS (Peak-Seq), the associated weight is lower. This is likely due to redundancy within the two FATHMM score datasets.

IV. CONCLUSIONS

Motivated by observations with our previous development of MKL methods [16], in this paper we propose a greedy CADD version 1.3 was released as this draft was in preparation, so we present results for versions 1.2 and 1.3.
approach to pre-selecting data sources. Our new model gives
greater test accuracy than our original model [12]. This greedy
approach suggests that certain types of data can be ignored
because the information they contain is implicit in an already
learned data source (encoded into its respective kernel), or
because a new data source contains little new information and
may also contribute a substantive extent of noise.

These results suggest further promising directions for future
work, to improve the data integration procedure. We intend
exploring other MKL methods such as those surveyed in
[5]. Indeed, the difference of convex approach we proposed in
[16], achieved up to 6% greater test accuracy over the
SimpleMKL method used here, for benchmarking studies with
some datasets (though it has an adjustable parameter which
must be found via a validation study). The method in [16]
also has the advantage that the kernel weights \( \lambda_i \) are found
separately from the learning parameters \( \alpha_i \). After deriving the
kernel weights, and hence a composite kernel, this means that
the composite kernel can be used in any kernel-based learning
method. For example, the Core Vector Machine [14] can
handle larger datasets than an SVM (computational complexity
scales as \( m \) rather than the \( m^3 \) of the SVM) and, by using
more training data, we could improve performance. Rather
than integrating component feature groups at the level of the
data, via a composite kernel, it would also be possible to
integrate classifiers via ensemble learning. In future projects,
we shall investigate these potential improvements in addition
to devising bespoke predictors for labelling variants in specific
disease contexts, such as cancer.

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