Inferring latent task structure for Multitask Learning by Multiple Kernel Learning

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Abstract

Background: The lack of sufficient training data is the limiting factor for many Machine Learning applications in Computational Biology. If data is available for several different but related problem domains, Multitask Learning algorithms can be used to learn a model based on all available information. In Bioinformatics, many problems can be cast into the Multitask Learning scenario by incorporating data from several organisms. However, combining information from several tasks requires careful consideration of the degree of similarity between tasks. Our proposed method simultaneously learns or refines the similarity between tasks along with the Multitask Learning classifier. This is done by formulating the Multitask Learning problem as Multiple Kernel Learning, using the recently published q-Norm MKL algorithm.

Results: We demonstrate the performance of our method on two problems from Computational Biology. First, we show that our method is able to improve performance on a splice site dataset with given hierarchical task structure by refining the task relationships. Second, we consider an MHC-I dataset, for which we assume no knowledge about the degree of task relatedness. Here, we are able to learn the task similarities ab initio along with the Multitask classifiers. In both cases, we outperform baseline methods that we compare against.

Conclusions: We present a novel approach to Multitask Learning that is capable of learning task similarity along with the classifiers. The framework is very general as it allows to incorporate prior knowledge about tasks relationships if available, but is also able to identify task similarities in absence of such prior information. Both variants show promising results in applications from Computational Biology.
in Computational Biology and highly structured relation across organisms (tasks), we apply our method to two important Computational Biology problems, namely MHC-I binding prediction and splice site prediction. The competitiveness of our results shows the validity of our approach.

Preliminaries
In a single-task supervised learning scenario, a sample of example-label pairs \( D = \{(x_i, y_i)\}_{i=1}^n \) is given, where the \( x_i \) live in an input space \( X \) and \( y_i \in \{-1, 1\} \) (for binary classification). The goal is to learn a function \( f \) such that \( f(x_i) = y_i \) that generalizes well to unseen data.

Before we describe our formulation of MTL as MKL approach, we briefly review the formulations of MTL and MKL that lay the foundations for our approach.

Multitask Learning
In MTL [1], we are given one labeled sample \( D_t \) for each of \( T \) tasks. Similar to the single-task supervised learning scenario, we are now interested in obtaining \( T \) hypotheses \( f_t \) one for each task.

We will formulate our method based on the Support Vector Machine (SVM), which has proven to generalize well [11], scales to large amounts of training data [12,13] and is able to incorporate arbitrary data sources by means of kernels (e.g., [14]). The generalization to other learning approaches appears straightforward as we mainly consider the extension of kernels to reflect task similarity, although details regarding the learning of their linear combination may differ.

Therefore, we start out with a regularization-based Multitask Learning method that was similarly proposed in the context of SVMs [2,10,15]. The basic idea is that models \( w_t \) are learned simultaneously for all tasks. Information transfer between tasks is achieved by sharing a general component \( w_0 = \sum_{t=1}^{T} \alpha_t \) and enforcing similarity of each \( w_t \) to \( w_0 \) in the joint optimization problem via regularization. We use the following formulation, leaving out some constants for readability

\[
\min_{w_1, \ldots, w_T} \frac{1}{2} \sum_{t=1}^{T} \|w_t\|_2^2 + \sum_{t=1}^{T} \|w_t - w_0\|_2^2 + C \sum_{t=1}^{T} \sum_{(x_i, y_i) \in D_t} \ell(x_i, w_t y_i).
\]

where \( \ell \) is the hinge loss, \( \ell(z,y) = \max\{1 - yz, 0\} \).

It was shown in [15], that the dual formulation of the above corresponds to the standard SVM using a modified kernel function:

\[
\max_{\alpha} -\frac{1}{2} \sum_{i=1}^{n} \sum_{j=1}^{n} \alpha_i \alpha_j y_i y_j \tilde{K}(x_i, x_j) + \sum_{i=1}^{n} \alpha_i
\]

s.t. \( \alpha^T y = 0 \), \( 0 \leq \alpha_i \leq C \forall i \in \{1, n\} \),
where \( K_B \) denotes the base kernel that captures the interactions between examples from all tasks and

\[
\tilde{K}(x_i, x_j) = K_B(x_i, x_j) + \delta_{t(i),t(j)} K_B(x_i, x_j).
\]

Here, \( t(i) \) denotes the task of example \( x_i \). In the above formulation, \( \tilde{K} \) is composed of the general kernel \( K_B \) and the kernel \( \delta_{t(i),t(j)} K_B(x_i, x_j) \) that captures only inter-domain interactions. In [9], the latter kernel is referred to as Dirac kernel. A slightly more general formulation of \( \tilde{K} \) is the following, which allows to adjust the trade-off between the general kernel and the task-specific kernel:

\[
\tilde{K}(x_i, x_j) = \beta_1 K_B(x_i, x_j) + \beta_2 \delta_{t(i),t(j)} K_B(x_i, x_j),
\]

where \( \beta_1, \beta_2 \geq 0 \) and \( \beta_1 + \beta_2 = 1 \). Clearly, \( \tilde{K} \) is a convex combination of base kernels and thus a valid kernel. MKL is a technique to learn the domain interactions. In [9], the latter kernel is referred to as \( \delta \) kernel. A slightly more general formulation of \( \tilde{K} \) is the following, which allows to adjust the trade-off between the general kernel and the task-specific kernel:

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\]
yield a sparse solution. Most subset weights will be set to zero, yielding only a few relevant subsets with weights greater than zero. We expect that the examples in these subsets come from similar distributions and that it is therefore beneficial to consider interactions between them, when obtaining a multitask predictor.

While L1-regularization of MKL results in a sparse combination of kernels, it does not address the computational complexity of the optimization problem over this exponential search space. With the current implementation, the method is limited to approximately 10 tasks depending on the number of training examples and available resources. However, there are techniques to handle the case where the number of tasks may become prohibitive, for instance, as proposed in [20]. The idea is to iteratively generate new kernels based on the current solution ($\beta$, w). These methods are known to converge to the optimal solution, if one can identify appropriate kernels in a larger set. In the current case, this could be done by solving an integer linear program.

Hierarchical MT-MKL

In the second scenario, we assume that we are given a tree structure $G$ that relates our tasks at hand (see Figure 1). In this context, a task $t_i$ corresponds to a leaf in $G$. Assuming hierarchical relations between tasks is particularly relevant to Computational Biology where often different tasks correspond to different organisms. In this context, we expect that the longer the common evolutionary history between two organisms, the more beneficial it is to share information between these organisms in a MTL setting. We can exploit the hierarchical structure $G$ to determine which subsets might play a role for Multitask Learning. In other words, we use the hierarchy to restrict the space of task sets. Let $\text{leaves}(n) = \{|l| l \text{ is descendant of } n\}$ be the set of leaves below the sub-tree rooted at node $n$. Then, we can give the following definition for the hierarchically decomposed kernel function

$$\hat{K}(x, y) = \sum_{\tau_i \in G} \beta_i K_{\text{leaves}(\tau_i)}$$

As an example, consider the kernel defined by a hierarchical decomposition according to Figure 1. Clearly, the number of $\beta_i$ corresponds to the number of nodes. For a perfect binary tree this leads to $2^m - 1$ nodes, where $m$ is the number of leaves/tasks. We expect that learning the contributions of the individual levels of the taxonomy makes sense for cases, where the edge lengths of $G$ are unequal.

Relation to task similarity

The learned weights $\beta_i$ reflect the importance of the subset $S_i$. Clearly, if two tasks $t_k$ and $t_l$ are often jointly present in subsets with high weights, we expect those tasks to be similar to each other. One can infer a measure of pairwise similarities between tasks $\gamma_{k,l}$ from the weights $\beta_i$ of the subsets $S_i$. We define the collection of task sets containing task $t_k$ as $T_{t_k} = \{S| t_k \in S \land S \in T\}$. Using this definition, we can define the similarity $\gamma_{k,l}$ between two tasks by summing up the weights of the shared task sets $S_i$

$$\gamma_{k,l} = \sum_{S_i \in T_{t_k} \cap T_{t_l}} \beta_i.$$  \hspace{1cm} (3)

This similarity measure can be used for downstream analyses, as it provides insight about the task relationships. A high $\gamma_{k,l}$ between tasks suggests a considerable resemblance between tasks and could help to generate domain knowledge (e.g., evidence that two cell-receptors bind to similar class of proteins, or the molecular mechanisms of the splicing machinery particularly similar).

Results and discussion

We performed experiments in two settings. In the first setting, we considered a set of MHC-I (major histocompatibility complex) proteins. Here, we assume we are not given any prior information to relate them. In the second setting, we used splice site data from 15 organisms and assumed that the task relationship is given by a hierarchical structure according to their evolutionary history. The examples are string data over an alphabet.
\{A,T,G,C\} (DNA) in the splicing case and the alphabet of 20 amino acids in the MHC-I case. To incorporate string features, we used the Weighted Degree String Kernel \cite{21}, which amongst other kernels such as the Spectrum Kernel \cite{22}, has been successfully employed in problems from Computational Biology.

In addition to the two MKL-based methods, we evaluated the following baseline methods:

- **Union** - One global model is obtained on the union of examples from all tasks.
- **Plain** - For each task, a model is trained independently, not taking into account any out-of-domain information.
- **Vanilla MTL** - Our algorithms consists of two components - the MTL formulation and the adjustment of weights \(\beta_i\) with MKL. In the vanilla approach, we fix all weights at \(\beta_i = 1\).

Experiments were performed by using cross-validation for model-selection on the training splits. We only tuned one hyper-parameter \(C\), for which we considered values between 0. 01 and 1000 on a logarithmic scale in 8 steps. After having obtained an optimal regularization parameter, a classifier is retrained on all training splits and final performance is obtained on a dedicated test set, that was not involved in hyper-parameter selection.

**MHC-I binding prediction using Powerset MT-MKL**

In this experiment, the task is to predict whether a peptide binds to a certain MHC molecule (binder) or not (non-binder). It has been previously shown that sharing information between related molecules (alleles) and thus cast- ing the problem in a Multitask Learning scenario, can be beneficial \cite{9}. In the MHC setting, different tasks correspond to different MHC proteins. The data consists of peptide sequences of length \(l = 9\) for 7 tasks. In total, we are given 7367 examples (\(A_{2403}=254\), \(A_{6901}=833\), \(A_{0201}=3089\), \(A_{0202}=1447\), \(A_{0203}=1443\), \(A_{2402}=197\), \(A_{2301}=104\)). For cross-validation, the data was split randomly into 5 splits of the same size. Unlike the setting of splice site prediction, we do not have a hierarchical structure relating our tasks at hand. To demonstrate that meaningful groups of tasks can be identified by Powerset MT-MKL, we do not assume any prior knowledge of task relationships. Please note, however, that we do have access to the sequences of the MHC-I proteins. We use these sequences to evaluate the task similarities obtained by our approach.

We report the area under the precision recall curve (auPRC) for the individual tasks in Figure 2 and the summary of performances in Table 1.

From Figure 2, we observe that the MKL-based approach outperforms the baseline methods. Furthermore, simply combining the data for different tasks to obtain a single model (**Union**) does not outperform the naïve method of obtaining an individual classifier for each task (**Plain**). This hints at rather large differences between the tasks. Learning the weights with MKL, improves performance compared to the **Vanilla MTL** approach, which already outperforms the other two baselines.

Figure 3 shows the distribution of weights obtained by the L1-regularized MKL approach. As expected, we observe that most task sets are assigned a weight of zero (or close to zero). Only a few get assigned a higher weight, so it is worthwhile to investigate the list of tasks that get assigned a weight \(\beta_i > 0.05\). From Table 2, we observe that the tasks \(A_{0201}, A_{0202}, A_{0203}\) are often grouped in the same task set, which is in agreement with domain knowledge. Based on the assigned weights, we compute the task similarity as defined in Equation (3). For evaluation of the learned similarities, we compare them to the hamming distance (or similarity) between the amino acids in the binding pocket \cite{23} of the MHC-I molecules (Figure 4). By visual inspection, we find a good agreement between the inferred task similarity and the molecule-based similarity.

Using MKL, we successfully identify groups of tasks among which information sharing is sensible, thus allowing for a smart combination of information from different tasks in the absence of prior knowledge.

The improvement in performance over the **Vanilla MTL** method is relatively small (a property most likely inherited from MKL). However, we are compensated for this by simultaneously obtaining a sensible task structure.

**Splice-site prediction using hierarchical MT-MKL**

In this setting, we take into account a given hierarchy (see Figure 5) relating the 15 organisms in our data set. The data set consists of 6000 examples for 15 tasks, each at a positive to negative ratio of 1:100, similar to the one used in \cite{3}. The data is split into 4 splits, three splits with 333 examples each and a large test split with 5000 examples. The dataset was created that way to establish a scenario where positive training examples are extremely rare.

We report the area under the precision recall curve (auPRC), which is well suited for unbalanced data sets. For the **Vanilla MTL** method, we use the given hierarchy \(G\) to define the initial task sets, but not further optimize their individual influence.

From Figure 6, we can make a few very interesting observations. First, in accordance with the results from the MHC-I experiment (see Table 3), the non-sparse Hierarchical MT-MKL methods outperform the baselines **Union** and **Plain**.

The second observation is that we get different results for different \(q\)-norms. In particular, we see a degraded
performance for $q = 1$, which complies with our expectation that weights for this approach (assuming the hierarchy is correct) should not be sparse. For the $q$-norms that we considered, $q = 2$ performs best. Lastly, we can show that we are able to outperform the Vanilla MTL method (all $b_i = 1$) by refining the task relations given by the structure $G$ with MKL. Intuitively, using Hierarchical MT-MKL corresponds to estimating the edge lengths of $G$, whereas the other method is restricted to directly using the similarities encoded into the taxonomy.

Conclusions

We have presented a principle way of formulating Multitask Learning as a Multiple Kernel Learning approach.

Table 1 Results for the MHC experiment in auPRC for the model selection step and the final prediction on the test set. Reported is the average performance over all tasks

| auPRC   | Plain | Union | Vanilla MTL | Powerset MT-MKL |
|---------|-------|-------|--------------|-----------------|
| cross-validation | 0.668 | 0.657 | 0.676 | 0.692 |
| test set      | 0.671 | 0.576 | 0.679 | 0.699 |

Figure 2 Result for the MHC experiments. Performance is shown for each of the 7 tasks. The performance averaged over all organisms is shown in the rightmost column mean.

Figure 3 Histogram of weights. Shows the distribution of weights $b_i$ that are learned for the elements of the power set by MKL. As expected, most are (close to) 0.

Table 2 List of task sets and their respective weights $b_i$ that were assigned by 1-norm MKL

| Task Set       | weight |
|----------------|--------|
| A_0201, A_0202, A_0203, A_6901 | 0.186 |
| A_0201, A_0202, A_0203, A_2301 | 0.178 |
| A_0202, A_0203, A_2301, A_2402, A_2403 | 0.110 |
| A_0201, A_0202, A_2301, A_2402, A_2403 | 0.091 |
| A_0201, A_0202, A_2301, A_2402, A_6901 | 0.074 |
| A_0201, A_0202, A_2301, A_2402, A_2403 | 0.066 |
Figure 4 Comparison between learned similarities and similarities based on the comparison of allele sequences. The learned similarity of A-2301 with A-0203, A-0201 and A-0202 in (b) can be attributed to structural features that cannot be easily inferred from the allele sequence.

Figure 5 Hierarchical structure that defines the relationship between the organisms.

Figure 6 Performance on test set for individual tasks. The performance averaged over all organisms is shown in the rightmost column mean.
Table 3 Results for the splice site experiment in auPRC for the model selection step and the final prediction on the test set. Reported is the average performance over all tasks. This table shows only the performance for the best-performing variant of Hierarchical MT-MKL with norm q = 2.

| auPRC         | Plain | Union | Vanilla MTL | Hierarchical MT-MKL |
|---------------|-------|-------|-------------|---------------------|
| cross-validation | 0.043 | 0.092 | 0.087       | 0.010               |
| test set      | 0.059 | 0.153 | 0.169       | 0.190               |

Following the basic idea of task-set-wise decomposition of the kernel matrix, we present a hierarchical decomposition and a power set based approach.

These two methods allow us to elegantly identify or refine structure relating the tasks at hand in one global optimization problem. We expect our methods to work particularly well in cases, where edge weights differ within the hierarchical structure or where the task structure is unknown.

Our experiments illustrate that the MT-MKL approach on the power set of all tasks works well for the MHC binding problem: First it increases the accuracy of the predictors compared to the baseline methods and second, the inferred task similarity reflects the prior knowledge that is available for this problem. Also for the splice site prediction problem where the task hierarchy is given by the organisms’ phylogeny, our approach manages to achieve an improvement over standard approaches. Using MKL on top of regular Multitask Learning methods may uncover latent task structure and thereby provide insight into the problem domain, which might be relevant to downstream analyses. In conclusion, this work constitutes a valuable proof-of-concept outlining a principle way of using MKL to improve Multitask Learning.

List of abbreviations used
MKL: Multiple Kernel Learning; MTL: Multi Task Learning; MHC: Major Histocompatibility Complex; SVM: Support Vector Machine; auPRC: area under the Precision Recall Curve.

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Authors’ contributions
Christian Widmer worked out the idea and implementation, performed the experiments and prepared part of the manuscript. Yasemin Altun was involved in the discussions, the development of methods on which this paper is based and the preparation of the manuscript. Nora C. Toussaint contributed to the discussions, provided the data for the MHC-I experiments and contributed to the preparation of the manuscript. Gunnar Rätsch came up with the original idea for the project and supervised the project at each step.

Competing interests
The authors declare that they have no competing interests.

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