Automatic Representation and Classifier Optimization for Image-based Object Recognition

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Abstract: The development of image-based object recognition systems with the desired performance is – still – a challenging task even for experts. The properties of the object feature representation have a great impact on the performance of any machine learning algorithm. Manifold learning algorithms like e.g. PCA, Isomap or Autoencoders have the potential to automatically learn lower dimensional and more useful features. However, the interplay of features, classifiers and hyperparameters is complex and needs to be carefully tuned for each learning task which is very time-consuming, if it is done manually. This paper uses a holistic optimization framework with feature selection, multiple manifold learning algorithms, multiple classifier concepts and hyperparameter optimization to automatically generate pipelines for image-based object classification. An evolutionary algorithm is used to efficiently find suitable pipeline configurations for each learning task. Experiments show the effectiveness of the proposed representation and classifier tuning on several high-dimensional object recognition datasets. The proposed system outperforms other state-of-the-art optimization frameworks.

1 INTRODUCTION

The object recognition problem widely occurs in many relevant real-world applications like optical character recognition (OCR), robotic vision or driver assistance systems. The human visual system has extraordinary pattern recognition capabilities and can perform many of the aforementioned tasks effortlessly. There is currently no general purpose computational model for object recognition with similar capabilities. Generally, object recognition is subdivided into object segmentation, feature extraction and classification. This paper aims at an automatic optimization of the last two aspects in a holistic manner.

The input data is low-level and noisy pixel data which is usually not invariant to e.g. scale, rotation or illumination changes. The properties of the feature representation have a tremendous effect on the classifier performance. Some popular standard features exist, e.g. SIFT (Lowe, 2004) or Local Binary Patterns (Ojala et al., 2002), which provide a reasonable degree of invariance. However, in many cases task-specific features have to be developed manually to fulfill the performance requirements.

The field of representation learning is focused on the analysis and automatic construction of good features for machine learning (Bengio et al., 2013). There are multiple approaches to learn new features; deep neural networks (Ngiam et al., 2011) have shown a great success in fields of object and speech recognition. Manifold learning is a promising approach to generate lower-dimensional features that potentially circumvent the curse of dimensionality. There are numerous different algorithms based on e.g. statistical analyses, neural networks or neighborhood graphs, and their performance heavily depends on the learning task. The interplay of selected features, manifold learning algorithms, classifiers and hyperparameters is very complex and needs to be carefully adapted for each learning task.

In order to tackle the burden of manual optimization this paper uses a holistic optimization pipeline with all the aforementioned components recently proposed in (Bürger and Pauli, 2015). The framework contains portfolios for manifold learning algorithms and multiple classifier concepts. The extremely large search space of this optimization problem is handled with evolutionary algorithms so that optimized processing pipelines can be obtained within a few hours on a normal workstation computer. The huge degree of adaptability bears the risk of overfitting to the train-

1Hyperparameters influence a learning algorithm itself, like the regularization parameter $C$ in a support vector machine (SVM).
A manifold learning algorithm is incorporated in the cross-validation process. Additionally, the variation of the best solutions of the evolutionary algorithm is exploited to build a multi-pipeline classifier.

The proposed framework already showed promising results on rather low-dimensional (4–60 dimensions) classification datasets from the UCI database (Bache and Lichman, 2013). This paper presents improvements on the framework and experiments on high-dimensional (256–1370 dimensions) real-world, multiclass object recognition tasks. The performance of automatically learned features out of low-level features is compared with standard high-level features. Also, the impact of the portfolio of manifold learning algorithms is evaluated. Finally, comparisons to the state-of-the-art optimization framework Auto-WEKA (Thornton et al., 2013) are made.

2 RELATED WORK

The field of object recognition is too large to give an extensive overview of the topic. This section focuses on feature construction methods and automatic optimization of machine learning systems.

2.1 Feature Construction and Manifold Learning

Manifold learning and dimension reduction feature transforms are one aspect of representation learning. The basic idea is to feature distributions typically do not fill all dimensions equally, but contain areas of lower-dimensional structures that are embedded in the high dimensional space. Manifold learning makes use of these geometrical structures and correlations to learn a model to transform high-dimensional feature vectors into the intrinsic lower dimensional space. There are numerous different linear and non-linear manifold learning algorithms; a list of common methods and references can be found in the appendix. The work of (Van der Maaten et al., 2009) and (Ma and Fu, 2011) provide an overview of these methods.

These algorithms base on completely different approaches – e.g. neural networks, statistical analyses, kernel methods, neighborhood graphs –, but they fit to a generalized feature transform interface \( f_{\text{FeatTrans}} \). A training set \( T \) is given with \( 1 \leq i \leq m \) feature vectors \( \mathbf{x}_i \in \mathbb{R}^D \) and class labels \( y_i \in \{\omega_1, \omega_2, \ldots, \omega_c\} \). Note that most manifold learning algorithms are unsupervised and thus ignore the labels. A manifold learning algorithm to reduce the dimensionality from \( D \) to \( d \) can expressed with a learning function

\[
M = f_{\text{learn}}(T, d)
\]

that derives the model variables \( M \). The transform function

\[
\mathbf{x} = f_{\text{trans}}(\mathbf{x}, M) \in \mathbb{R}^d
\]

embeds a vector \( \mathbf{x} \in \mathbb{R}^D \) into the new subspace using the model \( M \).

In real-world applications, there are several problems with manifold learning algorithms. First, the feature transform function must be capable of embedding previously unseen feature vectors to allow an out-of-sample extension. Linear methods use a transformation matrix and can extend any vector, but many non-linear methods lack a direct extension method. The Nyström theorem (Bengio et al., 2003) can be used to estimate this extension for methods that rely on spectral decompositions like LLE, Isomap or Laplacian Eigenmaps. Furthermore, many methods work well for artificial datasets, but fail to produce useful features on noisy real-world datasets as experiments in (Van der Maaten et al., 2009) show.

2.2 Automatic Machine Learning Optimization

Automatic optimization frameworks aid developers to select suitable features, classifier concepts and hyperparameters which is often referred to as model selection problem. The more components contain degrees of freedom, the more complex the optimization gets and only few publications deal with holistic approaches. Usually, search-based meta heuristics are used to find solutions within a reasonable time. In (Huang and Wang, 2006) and (Huang and Chang, 2007) “classical” feature selection is combined with hyperparameter optimization of a single classifier using evolutionary algorithms (see section 4.2). The Auto-WEKA framework (Thornton et al., 2013) aims to solve the combined feature selection, classifier selection and hyperparameter optimization problem using a Bayesian approach. It provides the most comprehensive amount of optimized components as it relies on the numerous algorithms contained in the WEKA machine learning framework (Hall et al., 2009). Therefore, Auto-WEKA will be compared in the experiment section in section 5.

However, Auto-WEKA does not contain the selection of manifold learning or feature transform methods. In (Bürger and Pauli, 2015), we introduced a holistic optimization framework for feature selection, multiple feature transforms, multiple classifiers and hyperparameters that uses evolutionary algorithms. This framework is briefly described in the following.
3 HOLISTIC CLASSIFICATION PIPELINE

The framework proposed in (Bürger and Pauli, 2015) provides a connection between classifier with hyperparameter selection, feature selection and automatic feature construction methods while the last two aspects are considered as representation optimization. The central processing algorithm is based on a classification pipeline framework which is described in the following subsections.

3.1 Pipeline Structure

The classification pipeline structure contains four pipeline elements that are depicted in figure 1. It has basically two modes – first, the training mode in which the training dataset $T$ is used to adapt and train all pipeline elements. In the classification mode new feature vectors are classified with the trained pipeline. The general processing concept of the pipeline is consecutive dimension reduction after each pipeline element:

$$d_{in} \geq d_{FeatSel} \geq d_{FeatTrans} \geq d_{Label} = 1.$$  \hspace{1cm} (3)

3.1.1 Feature Scaling Element

The first pipeline element performs a rather simple feature scaling to a value range of $[0,1]$ based on the minimum and maximum values in $T$. This preprocessing step usually leads to a performance improvement for all machine learning algorithms that rely on distance metrics.

3.1.2 Feature Selection Element

The feature selection element is the first dimension reduction in the pipeline and selects a subset of single features $S_{FeatSel} \in \mathcal{P}(\{1,2,\ldots,d_{in}\}) \setminus \emptyset$. The idea is that this element removes any irrelevant features that could possibly disturb any following algorithm. The dimensionality of the remaining feature space is denoted as $d_{FeatSel}$.

3.1.3 Feature Transform Element

The feature transform element is used to apply a manifold learning algorithm $f_{FeatTrans}$ and finally transform the data into a new feature space with dimensionality $d_{FeatTrans}$. During training, one algorithm is chosen out of a portfolio $S_{FeatTrans}$ (see appendix) and trained with the selected feature subset $S_{FeatSet}$ of $T$. The portfolio $S_{FeatTrans}$ also contains the identity function (or simply no transform) because all other transformations might fail to produce more useful features than the original ones.

3.1.4 Classifier Element

The last pipeline element selects a classifier $f_{Classifier}$ out of a portfolio $S_{Classifiers}$ of concepts listed in table 1. The no-free-lunch theorem states that no single classifier concept performs best for all learning tasks and the potential of a good feature transform might be lost if a suboptimal classifier concept is chosen. In training mode, the transformed data from the previous element is used to train the classifier. Note that each classifier has its own set of independent hyperparameters $S_{Params}(f_{Classifier})$ that can be of arbitrary type (also see table 1).

3.2 Pipeline Configuration

The proposed pipeline has numerous degrees of freedom to adapt to each learning task. The most important parameters are summarized in the pipeline configuration

$$\theta = (S_{FeatSet}, f_{FeatTrans}, d_{FeatTrans},$$

$$f_{Classifier}, S_{Params}(f_{Classifier})), \hspace{1cm} (4)$$

namely the selected feature subset $S_{FeatSet}$, the manifold learning algorithm $f_{FeatTrans}$ and its target dimensionality $d_{FeatTrans}$ as well as the classifier concept $f_{Classifier}$ and its corresponding hyperparameters $S_{Params}(f_{Classifier})$. 
Table 1: Popular classifier concepts and corresponding hyperparameters and ranges. References to these classifier concepts can be found e.g. in (Bishop and Nasrabadi, 2006) and (Huang et al., 2006).

<table>
<thead>
<tr>
<th>Classifiers</th>
<th>parameter ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>Naive Bayes</td>
<td>C: [10^{-2}, 10^4]</td>
</tr>
<tr>
<td>C-SVM linear kernel</td>
<td>C: [10^{-2}, 10^4], α: [10^{-2}, 10^4]</td>
</tr>
<tr>
<td>C-SVM Gaussian kernel</td>
<td>k: [3, 20], metric: [Euclidean, Mahalanobis, Cityblock, Chebychev]</td>
</tr>
<tr>
<td>k nearest neighbors (kNN)</td>
<td>hidden layers: [0, 10], neurons per layer: [1, 200]</td>
</tr>
<tr>
<td>Multilayer Perceptron (MLP)</td>
<td>number trees: [1, 100]</td>
</tr>
<tr>
<td>Extreme Learning Machine (ELM)</td>
<td></td>
</tr>
<tr>
<td>Random Forest</td>
<td></td>
</tr>
</tbody>
</table>

4 OPTIMIZATION ALGORITHM

The search space of all possible configurations is huge as it contains feature selection with exponential complexity $O(2^N)$ and the combinations of all manifold learning algorithms, classifiers and hyperparameters. The optimization approaches presented in (Bürger and Pauli, 2015) and improvements are described in the following.

4.1 Target Function

In order to prevent overfitting, a suitable optimization target metric based on a wrapper\(^2\) approach combined with $k$-fold cross-validation (Jain et al., 2000) is employed. The training dataset is divided into $k$ subsets and $k$ validation rounds are made. In each round $k - 1$ subsets are used for training and one is left out for validation.

The manifold learning algorithm has a great impact on the performance of the whole pipeline as a non-linear feature transform may inherit parts of the classifier’s “intelligence” as a linear classifier model might be sufficient. However, if the manifold learning is only used as a preprocessing step and is not involved into the cross-validation, the generalization of the out-of-sample function is never measured. Therefore, the manifold learning algorithm has to be considered in each validation round. The training set is separated into $k = 5$ training and validation tuples $\{(\mathcal{T}_{\text{train},1}, \mathcal{T}_{\text{valid},1})\}$. In each cross-validation round $1 \leq l \leq k$, the selected manifold learning algorithm is trained with $\mathcal{T}_{\text{train},l}$ and the out-of-sample extension transforms the test and validation dataset – separately – into the new feature space denoted as $(\mathcal{T}_{\text{train},l}, \mathcal{T}_{\text{valid},l})$. The selected classifier is trained with the transformed training dataset $\mathcal{T}_{\text{train},l}$ and the accuracy is measured with the predictions on the transformed validation dataset $\mathcal{T}_{\text{valid},l}$. The average accuracy of all cross-validation rounds is used as target metric.

4.2 Optimization with Evolutionary Strategies

Evolutionary algorithms (EA) are especially suitable to solve complex optimization problems with high dimensional search spaces. The basic idea of EA is the imitation of the biological evolution of species that adapt to their environment over time while Evolutionary Strategies (ES) are one variant of EA that are especially capable to optimize sets of heterogeneous parameters (Beyer and Schwefel, 2002). The objective function $f(y)$ needs to be optimized with respect to the object parameter set $y$. These parameter sets can be any combination of different data types:

- numeric data types $\mathbb{R}^N$ and $\mathbb{Z}^N$ with length $N$ and corresponding minimum and maximum values,
- bit string data types $\mathbb{B}^N$ with length $N$,
- categorical data types $\mathcal{S} = \{s_1, s_2, ..., s_N\}$ with $N_\mathcal{S}$ items without any order.

ES uses populations of individuals containing parameter sets $y$. The optimization starts with random individuals which are evolved over time with the help of the evolutionary operators selection, recombination and mutation. The fitness function $f(y)$ evaluates each individual and only the fittest survive and generate offspring.

Figure 2 depicts a strategy to code the pipeline configuration $\vartheta$ completely as parameter set $y$. The feature subset $S_{\text{FeatSet}}$ is coded as bit string $\mathbb{B}^{n_{\text{feat}}}$. The manifold learning algorithm $f_{\text{FeatTrans}}$ and the classifier concept $f_{\text{Classifier}}$ are both coded as categorical type $\mathcal{S}$. The target dimensionality $d_{\text{FeatTrans}}$ for the manifold learner cannot be coded directly as it depends on the number of selected features in $S_{\text{FeatSet}}$. Instead, a dimension fraction factor $\alpha \in [0, 1]$ is used for the coding and the target dimensionality is obtained with

$$d_{\text{FeatTrans}} = \lfloor \alpha \cdot d_{\text{FeatSet}} \rfloor, \quad d_{\text{FeatTrans}} \geq 1. \quad (5)$$

The initial random values for $\alpha$ are sampled from a smaller range of $[0, 0.1]$ which is a prior for lower dimensional feature spaces.

The structure of the classifier hyperparameter optimization problem $S_{\text{Params}}(f_{\text{Classifier}})$ is hierarchical as the selection of a classifier concept $f_{\text{Classifier}}$ selects the classifier specific set of hyperparameters (see Table 1).
In order to optimize all hyperparameters in a single evolutionary way, the hierarchical problem is linearized in the following way: All hyperparameters of all classifiers are simply appended to the object parameter set and evolved together. Exponentially ranged hyperparameters (C and γ values for the SVM) are optimized using the exponents log10(x). For the evaluation of the fitness (see section 4.1), the pipeline configuration uses those hyperparameters that belong to the selected classifier f_{classifier} and simply neglects the rest.

The parametrization of the ES optimization is summarized in the (µ/ρ + λ) notation. The initial population contains 500 random individuals and in each generation λ = 200 individuals from ρ = 2 parents are generated while the best µ = 50 individuals survive. The mutation operator contains several probability parameters:

- the probability of bit flips for the feature selection is \( p_{\text{bitflip}} = 0.3 \).
- the probability of choosing a random item for all parameters of the categorial type \( S \) is \( p_{\text{cat}} = 0.3 \).
- for all numerical values, namely the α factor and the classifiers’ hyperparameters, the mutation is performed using an additive normally distributed noise \( \mathcal{N}(0, \sigma^2) \). The standard deviation is adaptive to the value range of the corresponding hyperparameter with \( \sigma = 0.2 \cdot (v_{\text{max}} - v_{\text{min}}) \).

The ES terminates if the improvement of the fitness of the best individuals is less than \( \varepsilon = 10^{-2} \) after three consecutive generations.

### 4.3 Multi-pipeline Classifier

The result of an ES optimization is a set of configurations with corresponding fitness values \( \{ (\theta_j, f_j) \} \). This set can be sorted by the fitness values to obtain a top list of configurations, such that \( \theta_1 \) is the best solution. Of course, this top configuration can be used to set up a classification pipeline, but a single pipeline can be problematic in the sense of overfitting. A simple solution is the fusion of multiple pipelines to a multi classifier which generally leads to a better generalization when the diversity among the classifiers is large enough (Ranawana and Paalde, 2006). A multi-pipeline classifier can easily be built with setting up the top-\( n \) pipelines with the corresponding configurations \( \theta_j \) with \( 1 \leq j \leq n \). New instances are classified by all pipelines and a majority voting is performed to obtain the final label.

### 5 EXPERIMENTS

The proposed framework is tested on two real-world image-based object recognition tasks with different scenarios. The datasets contain low-level pixel features and relatively few training instances compared to the number of dimensions. This leads to the curse of dimensionality which is a typical issue for image-based classification tasks.

The focus of the experiments is the question what role manifold learning plays and how different types of features influence the performance. Three variants of the proposed framework are evaluated:

- **Manifold Learning Deactivated (ML: none):** The set \( S_{\text{FeatTrans}} \) just contains the identity function.
- **Only PCA (ML: PCA):** The set \( S_{\text{FeatTrans}} \) contains the “popular” linear PCA and the identity.
- **All Manifold Learning Methods (ML: All):** The set \( S_{\text{FeatTrans}} \) contains all linear and non-linear manifold learning algorithms listed in the appendix.

All datasets are separated randomly into 70% training and 30% test data. The framework is implemented in Matlab using the parallel computing toolbox. A baseline SVM classifier with Gaussian kernel is used for comparison as well as the Auto-WEKA framework with 24 hours time budget.

#### 5.1 Classification of Coins

For this task, 5 different classes of Euro coins (1, 2, 5, 10 and 20 cent coins) need to be distinguished by using RGB-color images. For each class, 16 different coins are photographed from four different angles leading to a total amount of 320 images. The coins are segmented by simple thresholding (see figure 3) and used to generate three datasets with different feature sets:
Figure 3: Examples of the coins dataset.

Table 2: Training cross-validation accuracy values for the coins datasets.

<table>
<thead>
<tr>
<th>Feature Set</th>
<th>Low</th>
<th>High</th>
<th>Low+high</th>
</tr>
</thead>
<tbody>
<tr>
<td>ML: none</td>
<td>69.78</td>
<td>85.78</td>
<td>83.56</td>
</tr>
<tr>
<td>ML: PCA</td>
<td>71.56</td>
<td>85.78</td>
<td>82.22</td>
</tr>
<tr>
<td>ML: All</td>
<td>68.44</td>
<td>93.78</td>
<td>92.00</td>
</tr>
<tr>
<td>Baseline</td>
<td>58.67</td>
<td>70.22</td>
<td>70.22</td>
</tr>
</tbody>
</table>

- **Low-level Features.** Normalized pixel values (zero mean and standard deviation of 1) of 30 × 30 pixel images; total dimensionality: 900.
- **High-level Features.** Area in pixels, statistical features of gray and hue values in the HSV\(^3\) space (mean, standard deviation, skewness and kurtosis), histograms with 50 bins of gray and hue values, Local Binary Patterns in four variants: basic (256 dimensions), uniform (59 dimensions), rotation invariant (36 dimensions), rotation invariant and uniform (10 dimensions); total dimensionality: 470.
- **Low+high-level Features.** Low- and high-level features together; total dimensionality: 1370.

5.1.1 Train and Test Performance

Table 2 lists the best cross-validation accuracy values during training for all three framework variants (rows) and the three feature sets (columns). When all manifold learning algorithms are used (variant ML: All) a significant accuracy gain is achieved for the high- and low-high-level features. Only for the low-level features, the ML: PCA variant achieves the best result, likely due to local minima during the optimization of the full manifold learning algorithm portfolio. The baseline SVM classifier is clearly outperformed in all cases.

The optimization times on an Intel Xeon workstation with 6 × 2.5 Ghz are listed in table 3. Due to the larger search space and the high computation times of some of the manifold learning algorithms (caused by the Matlab implementations), the total optimization times are much larger for the ML: All variant.

Table 3: Optimization times in minutes for the coins datasets.

<table>
<thead>
<tr>
<th>Feature Set</th>
<th>Low</th>
<th>High</th>
<th>Low+high</th>
</tr>
</thead>
<tbody>
<tr>
<td>ML: none</td>
<td>19.2</td>
<td>29.7</td>
<td>46.9</td>
</tr>
<tr>
<td>ML: PCA</td>
<td>30.1</td>
<td>33.6</td>
<td>47.8</td>
</tr>
<tr>
<td>ML: All</td>
<td>314.7</td>
<td>224.4</td>
<td>911.7</td>
</tr>
</tbody>
</table>

Table 4: Generalization accuracy values on the test dataset for the coins datasets (single top-1 configuration).

<table>
<thead>
<tr>
<th>Feature Set</th>
<th>Low</th>
<th>High</th>
<th>Low+high</th>
</tr>
</thead>
<tbody>
<tr>
<td>ML: none</td>
<td>78.95</td>
<td>83.16</td>
<td>83.16</td>
</tr>
<tr>
<td>ML: PCA</td>
<td>78.95</td>
<td>82.11</td>
<td>80.00</td>
</tr>
<tr>
<td>ML: All</td>
<td>75.79</td>
<td>84.21</td>
<td>92.63</td>
</tr>
<tr>
<td>Baseline</td>
<td>66.32</td>
<td>68.42</td>
<td>77.89</td>
</tr>
<tr>
<td>Auto-WEKA</td>
<td>72.63</td>
<td>89.47</td>
<td>92.63</td>
</tr>
</tbody>
</table>

Table 4 shows the generalization accuracy values on the test dataset when the overall best configuration is used. The proposed framework achieves a large performance boost on the low-level feature set compared to the training accuracy. The performance on the high-level feature set is significantly lower. When the low+high-level features are used, the accuracy on the test dataset is in the same range than during training. Clearly, overfitting effects are responsible for the partly relatively low generalization performance of the proposed framework. The Auto-WEKA framework performs best or equal on two of the three feature sets while the baseline SVM classifier shows a poor performance on all feature sets.

5.1.2 Multi-pipeline Classifier

The generalization performance values of the multi-pipeline classifiers can be found in figure 4. A general trend is visible: The performance is increasing when more pipelines are used. Usually, a strong boost is already achieved for less than 10 pipelines.

The distribution of the top configurations can contain useful information about the classification problem. Figure 5 shows the top-50 configurations for the ML: All framework variant on the low+high-level feature set as a graph. The frequencies of the components in the top solutions are denoted with different shadings. In this case, only one manifold learning algorithm is under the best solutions, namely the Large-Margin Nearest Neighbor (LMNN) method. It is interesting that the problem becomes linear as a linear SVM showed the best performance. Furthermore, the importance of the different features is easily visible – in this case the area is the most important feature.

\(^3\)The Hue Saturation Value (HSV) color space is used to analyze the color independently from brightness.
number pipelines
1 5 10 15 20 25 30 35 40 45 50
accuracy
75
80
85
ML: none
ML: PCA
ML: All
(a) Low-level features

number pipelines
1 5 10 15 20 25 30 35 40 45 50
accuracy
80
85
90
95
ML: none
ML: PCA
ML: All
(b) High-level features

number pipelines
1 5 10 15 20 25 30 35 40 45 50
accuracy
75
80
85
90
95
ML: none
ML: PCA
ML: All
(c) Low+high-level features

Figure 4: Generalization accuracy values on the test dataset of the multi-pipeline classifiers for the coins datasets depending on the number of pipelines.

Figure 5: Visualization of the top-50 configurations for the coins dataset with low+high level features using the ML: All variant. The overall best solution contains the items marked with an asterisk.

which is not surprising as the coins have a different size in reality. Also the contrast (gray value standard deviation feature) is important.

5.2 Handwritten Digits

For the second experiment the semeion-digits dataset (Buscema, 1998) from the public UCI database (Bache and Lichman, 2013) is used. The dataset contains 1593 samples of handwritten digits (0–9) in form of binary images with a size of 16 × 16 pixels (see figure 6). This leads to a 10-class problem with a feature dimensionality of 256. In this dataset, only the low-level pixel features are used.

5.2.1 Train and Test Performance

The training and test results are listed in table 5. It is remarkable that the baseline SVM classifier already performs very well on this dataset – the performance gain of the proposed framework and Auto-WEKA is marginal. The optimization times can be found in table 6 and are higher because there are more training samples compared to the coins datasets.

Table 5: Accuracy on training and test dataset for the semeion-digits dataset.

<table>
<thead>
<tr>
<th></th>
<th>training dataset</th>
<th>test dataset</th>
</tr>
</thead>
<tbody>
<tr>
<td>ML: none</td>
<td>95.16</td>
<td>92.66</td>
</tr>
<tr>
<td>ML: PCA</td>
<td>95.16</td>
<td>93.50</td>
</tr>
<tr>
<td>ML: All</td>
<td>93.28</td>
<td>92.87</td>
</tr>
<tr>
<td>Baseline</td>
<td>93.46</td>
<td>92.03</td>
</tr>
<tr>
<td>Auto-WEKA</td>
<td>-</td>
<td>94.13</td>
</tr>
</tbody>
</table>

Table 6: Optimization times in minutes for the semeion-digits dataset.

<table>
<thead>
<tr>
<th></th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>ML: none</td>
<td>28.8</td>
</tr>
<tr>
<td>ML: PCA</td>
<td>99.0</td>
</tr>
<tr>
<td>ML: All</td>
<td>1167.1</td>
</tr>
</tbody>
</table>

5.2.2 Multi-pipeline Classifier

The generalization performance of the multi-pipeline classifiers (see figure 7) is slightly increasing with the number of pipelines, and the best achievable performance is better or equal to Auto-WEKA. However, the performance gain is marginal for this dataset.

Figure 8 shows the distribution of the best configurations for the ML: All variant. The variety of well performing manifold learning algorithms and classifiers is large for this dataset.

6 CONCLUSIONS

This work presented improvements and extended
evaluations of a holistic optimization framework for feature selection, manifold learning, classifier and hyperparameter selection. The evaluations were performed on image-based classification problems with evident curse of dimensionality and different types of feature sets. The results show that the proposed optimization methods find a reasonable pipeline configuration within a few hours. The benefit of the manifold learning algorithms depends on the task and the chosen features, but can be potentially large (see the LMNN transform on the coins dataset). The type features (low- vs. high-level) does not play a big role to predict the success of manifold learning.

Overfitting effects are still noticeable even though cross-validation is used. However, detailed studies of the multi-pipeline classifier showed a reasonable boost in generalization performance, so that the state-of-the-art optimization framework Auto-Weka is beaten for all datasets. However, the relatively long optimization times are not always justifiable, if a baseline SVM already performs very well.

Future work will focus on the improvement of the optimization process in terms of speed and generalization. One approach can be the use of a fast estimation algorithm at the beginning to generate a better initial population. Furthermore, early rejection of inferior individuals during cross-validation is a promising approach. A remedy against overfitting is an increased diversity of configurations which can be achieve with e.g. the extension of the evolutionary selection operator with a maximum age of individuals. Furthermore, alternative target metrics can be applied, e.g. bootstrapping (Jain et al., 2000).

ACKNOWLEDGEMENTS

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REFERENCES


APPENDIX

List of Linear and Non-linear Dimension Reduction and Manifold Learning Methods in the Framework

Linear

Principal Component Analysis (PCA) (Pearson, 1901), Linear Local Tangent Space Alignment algorithm (LLTSA) (Zhang et al., 2007), Locality Preserving Projection (LPP) (Niyogi, 2004), Neighborhood Preserving Embedding (NPE) (He et al., 2005), Factor Analysis (Spearman, 1904), Linear Discriminant Analysis (LDA) (Fisher, 1936), Neighborhood Components Analysis (NCA) (Goldberger et al., 2004), Large-Margin Nearest Neighbor (LMNN) (Weinberger and Saul, 2009).

Non-linear

Kernel-PCA with polynomial and Gaussian kernel (Schölkopf et al., 1998), Denoising Autoencoder (Hinton and Salakhutdinov, 2006), Local Linear Embedding (LLE) (Donoho and Grimes, 2003), Isomap (Tenenbaum et al., 2000), Manifold Charting (Brand, 2002), Laplacian Eigenmaps (Belkin and Niyogi, 2001), parametric t-distributed Stochastic Neighborhood Embedding (t-SNE) (Van der Maaten, 2009).