Seminar aus Computer Vision und Mustererkennung

Dynamic Feature Selection for Media Classification

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Abstract

TODO Bla bla bla...

1 Introduction

Thanks to the technological development of the recent years, nowadays it is possible to collect and store a vast amount of data. High-dimensional data is omnipresent in a broad range of real-life applications, such as text analysis in social media or the analysis of genomes in bio informatics. But these possibilities also led to new challenges when evaluating those datasets. Not only the sample size, also the dimensionality of data has increased, which is the cause for the so called "curse of dimensionality": The complexity of the analysis (regarding memory consumption and computation time) grows exponentially with increasing dimensionality of the samples. Furthermore, many algorithms tend to overfit when too many sample points are given, which is another drawback of big feature sets. [?; ?]

To conquer this problem, various techniques to reduce dimensionality were developed in the field of machine learning. Two broad categories can be distinguished: feature extraction and feature selection. In feature extraction, features are combined to create "new" features in a lower dimensional space. Feature selection on the other hand tries to select a few, important features out of the given set. The features produced or chosen by either one of those methods are then used for classification and analysis of the sampled data.

The goal of this paper is to set a focus on feature selection and present some of the most common techniques and state-of-the art.

The main strategy of feature selection is to chose only a small subset of the available features for analyzing the data, which leads to higher accuracy, lower computational cost and better interpretability of the results.[] Such a subset would optimally consist only of relevant features, whereas redundant or useless features should be removed. What properties define a relevant feature depends on the application, but generally speaking, the optimal feature should be discriminant when performing classification (so when looking at a certain feature, a clear classification of the current data point should be possible) and should not be redundant (two different features should not provide the same kind of information). Possible criteria for evaluating the relevance of features will be covered in more detail in section FOO.

2 Methods

TODO Bla bla...es gibt flat und structured features...

2.1 Methods for Flat Features

If we look at flat features, three different types for feature selection methods can be distinguished:

- Filter methods
- · Wrapper methods
- · Embedded methods

In this section, a short introduction on those three categories will be given, including a discussion of their benefits and disadvantages.

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2.1.1 Filter Methods

As the name suggests, filter methods try to filter out relevant features from not-relevant ones. This is done by considering only the actual dataset, the properties of the classifier which will be used afterwards are ignored.

Thus, filter methods are independent of the classification algorithm: Any classifier can be used on the filtered features, which allows to try out different algorithms on the feature set.

This can be seen as an advantage, but on the other side, complete independency can also have its drawbacks. As characteristics of the specific classifiers are not considered, it is not clear which classifier works best with the selected subset. Thus the selected subset does not guarantee optimal performance of the classifier.

Quadratic programming Feature Selction: http://jmlr.org/papers/volume11/rodriguez-lujan10a/rodriguez-lujan10a.pdf

Fisher Score TODO Bla bla bla...

Information Gain TODO Bla bla bla...

Min. Redundancy - Max. Relevance TODO Bla bla bla...

Relief The original algorithm (Relief) was presented by Kira and Rendell [?], and can be used for binary classification problems. The algorithm is efficient (it runs in polynomial time), robust and tolerant to noisy data, but the quality of the results depends on the number of iterations. The basic idea is to use a weight vector, which has as many entries as there are features. In each iteration, an instance/variable x is chosen at random. Then the near-hit and near-miss instance of the remaining set in relation to x are selected: This is the most similar instance from the same class as x, and respectively the most similar instance of the different class. The euclidean distance between x and the near-hit and near-miss is calculated to determine if a feature is relevant or not.

Intuitively, if a feature is relevant, the near-hits should be closer than the near-misses on average, while for irrelevant features, near-hits and near-misses are independent from each other. For each feature f_i , the corresponding weight w_i in the weight-vector is updated dependent on those distances by applying equation 1:

$$w_i = -(x_i - nearhit)^2 + (x_i - nearmiss)^2 \tag{1}$$

Relevant features thus score values larger than zero, while irrelevant features become zero or negative. After a desired number of iterations, the features whose weights have a value above a certain threshold are chosen for the classification or further processing.

The original algorithm was further developed by [Kononenko et al. 1997]. ReliefF is extension so that the algorithm is able to handle incomplete data and multiple-class problems. RRELIEFF (Regressional ReliefF) [Robnik-Sikonja and Kononenko 1997] adapts the algorithm to handle also linear regression problems.

The family of Relief-algorithms and eventual adaptations have successfully been used on feature selection problems in the recent years. Moore used an adaptation called Tuned ReliefF (TuRF) for genetic analysis [Moore and White 2007], where the worst features are removed systematically in each iteration. By removing a fixed number of features in each iteration, the accuracy of weight-estimation is increasing compared to the results of the original ReliefF algorithm. Eppstein and Haake [Eppstein and Haake 2008] criticize that for truly genome-wide association analysis, where a huge number of features is used (up to hundreds of thousands), all the proposed methods don't scale well and the estimated weights become basically random values. Very large scale ReliefF (VLSReliefF) tries to overcome this limitation by simply applying Relief on subsets of the features, and then combining the results to gain the weights for all features. This process can be paralellized on the GPU to speed it up [] (Bibtex Quote? http://worldcomp-proceedings.com/proc/p2015/PDP2993.pdf)

(Evtl. other application: One application was automated classification of websites [Jin et al. 2007].)

2.1.2 Wrapper Methods

In contrast to filter methods, wrapper methods are dependent on the classifier used after the feature selection. The feature-set is selected so that it fits the biases and heuristics of the classifier as good as possible.

This is achieved by sequentially applying two steps:

- A search routine selects a set of features
- The selected subset is evaluated with the desired classifier

When choosing a search routine, one should take the structure of the search space into account. Especially its size plays a crucial role, as it grows exponentially with increasing the number of features. (The more features there are, the more possible subsets can be created.) As real-life applications can involve thousands of features, the search can not be done randomly, but has to follow a certain strategy. Typical algorithms used are hill-climbing, best-first, or greedy-search, to name a few.

When a potential subset is found, its suitability for the desired classifier has to be evaluated. This can be done for example by simply using a validation set, or by performing cross-validation.

The search and evaluation steps are repeated until a desired quality of the subset is reached. The first step will produce a set of features, which then are used by the classifier. The feature set with the best results (e.g. highest estimation values) will then be used for the actual classification task.

The major drawback of this approach is the computation time needed, as the subsequent search- and evaluation steps are computationally expensive. Compared to filter methods, better performance and more accurate estimates are achieved for the chosen classifier. Also, as the selected subset is dependent on the classifier, it cannot be used with any classifier, as the results will be biased.

This chapter will go more in-depth about different search-techniques, before discussing validation-strategies shortly.

Hill climbing Hill climbing, sometimes also referred to as "greedy search" or "steepest ascend", is probably the simplest search technique that can be applied. Starting with no features at all, an evaluation function rates the available features and adds the most relevant one to the subset. Then the procedure is repeated, each time adding the most relevant feature of the remaining set to the subset. As soon as the influence of an added feature does not improve the overall quality of the set significantly, the search is stopped. The problem with hill climbing is that it gets stuck in local maxima easily, and fails in finding a globally optimal solution. [Kohavi and John 1997]

Best First Best-first search outperforms hill-climbing, as it is a more robust technique. [Kohavi and John 1997]

Branch and Bound Branch-and-bound (BB) is a general design principle for solving discrete and combinatorial optimization problems. BB-algorithms work by building up a search tree with possible candidate solutions (branch phase). Further, a criterion function has to be designed, which evaluates the "quality" of possible solutions represented by nodes in the tree. Then, the branches of the tree are explored to find a global optimum for the given function. To avoid that the search is exploring the whole possible space, and thus degenerating to a brute-force-approach, it is limited by so-called bounds, which prune the tree when it becomes unlikely to find an optimum in the current branch.

For feature selection, Narendra et al. [Narendra and Fukunaga 1977] first proposed a BB-algorithm based on efficient subset selection. The root of the tree represents the full set of features, whereas leaf-nodes represent single features (or subsets of the smallest desired size). Monotonicity is required regarding the criterion function and the subsets. That means, if a current subset is below a bound, then its following child nodes are assumed to be also lower than that bound. Thus, the search in this particular branch can be aborted, and continued at another branch. Additionally, the sorting of the nodes is done with an efficient enumeration scheme to augment finding an optimal solution in a short amount of time.

The problem with BB-methods is that it is not guaranteed to perform better (faster) than exhaustive search methods. It is not granted that enough sub-trees will be pruned to speed up the search sufficiently, in worst-case, the criterion-function is evaluated for every node in the tree. Additionally, evaluating the function close to the root takes longer than evaluating it at the leaves. [Somol et al. 2004] proposed a fast BB approach, which tries to keep the evaluations of the criterion function at a minimum to speed up the search. The algorithm is "learning" how the removal of features influences the functions value, and tries to "predict" changes without doing the actual computation. Only if the predicted value comes close to a bound, the criterion function is evaluated, otherwise the algorithm continues descending in the tree.

Adaptive branch and bound for feature selection [Nakariyakul and Casasent 2007] also tries to reduce the number of evaluations of the criterion function, by applying several improvements. Already when building up the tree, the order of the nodes corresponds to the importance of the features. Then, instead of trying to descend sequentially from node to node, an adaptive, "jumping" search method is used to avoid more-or less redundant computations of the evaluation function. Additionally, the initial bound used in the search and also the initial search level are chosen in an optimal way. (Comment: only tested on 30 features?! Still check this out...)

Genetic Algorithms Before explaining how GA can be used for for feature selection, a short introduction into the very basic concept of those algorithms has to be made. The theoretical groundwork for genetic algorithms was laid by [Holland 1992]. As the name implies, genetic algorithms try to imitate the process of natural evolution: parents carry specific genetic information, and a (re-)combination of this genes is passed to their kids in order to create "better" offspring from generation to generation.

GAs imitate this behavior by encoding the data they should work on or optimize in so-called chromosomes, which thus represent a possible solution to the underlying problem. A common choice for the data structure of a chromosome is to use a vector of numerical values, which can be processed easily by the typical routines in a GA.

A desired number of chromosomes is initialized at the beginning of the algorithm, and two different operations are applied to create new "offspring"-chromosomes: crossover (a new chromosome can be obtained by combining the values of two other chromosomes) or mutation (changing random values in an already existing chromosome).

The resulting new chromosomes are then evaluated according to a fitness function, which evaluates the "quality" of the solution a chromosome actually represents. The chromosomes then can be ranked, and the best ones are taken again to create new offspring. This procedure is repeated as long until a desired quality/result is achieved.

For feature selection, chromosomes could represent subsets of the total feature set. The values in a chromosome would encode if a feature is selected or not. This can be achieved simply by using binary values: 1 indicates that a feature is selected for a possible subset, and 0 indicates that it is not selected.

The strength of GA is that they do not tend to get stuck at local optima, but instead are likely to find global optima, even in large datasets. Making an implementation efficient is the key for a successful use of GA for feature selection, as otherwise running time has to be fairly long

to gain acceptable results. Another problem is the choice of parameters for the algorithm (f.ex, when and how often to perform mutations over crossovers), which can influence the outcome to the better or the worse.

[Oh et al. 2004] criticize that GA are weak in fine-tuning local optima, and thus propose a hybrid algorithm for feature selection. While still performing a global search, classical search operations are applied on a local level, which led to significant performance improvements. Still, sequential search approaches tended to be faster regarding convergence. Parallel computing was suggested to overcome this problem, as the evaluation of single chromosomes is independent and could be done simultaneously.

[Fro] designed a very specific algorithm suited exclusively for feature-selection for SVMs. Computation time is reduced by considering already existing theoretical error bounds for this kind of classifier, instead of performing cross-validation to estimate the performance of a potential feature-subset. The genetic encoding of data is further used to simultaneously optimize parameters needed for the SVM itself. This method proved to be faster then RFE (Recursive feature elimination) when the number of features selected for a subset was not known beforehand. Furthermore, using error bounds tended less to overfitting of data compared to cross-validation.

Forward Selection/backward elimination In practice, two methods which are often used for large data sets are sequential forward selection (SFS) and sequential backward elimination (SBE).

Both technique work iteratively. Forward selection starts with an empty set, where no feature is selected in the beginning. Sequentially, one feature after the other is added to the subset, so that the new subset maximizes the quality of the subset, which is measured by some criterion function J. This is done until the set has reached a desired size. SBE works exactly the other way round, starting with the full set of features first and sequentially deleting features, until a smaller subset of sufficient quality is gained.

Both methods have a major drawback: Either features cannot be eliminated once they have been selected (SFS), or they cannot be selected again if they have been discarded once (SBE). The assumption, that the best five selected features must contain a subset of the best four selected features does not hold in practice. [Nakariyakul and Casasent 2008]

[Pudil et al. 1994] proposes Sequential forward floating selection as improvement (and, respectively, Selective backward floating elimination). A backtracking step is implemented after each sequential addition or deletion, which tries to find eventually better subsets. Both methods show admissible computation-time for small- or medium scaled problems, and perform better than other sequential methods on a variety of problems. However, it should be empathized that they do not always perform better than other methods, but at least "good enough" on the majority of problems. For very big datasets, they are outperformed by genetic algorithms. [Kudo and Sklansky 2000]

[Mao 2004] proposes *orthogonal* forward selection and backward elimination to overcome problems that occur with SFS and SBE. Instead of simply selecting features in a sequential way, they are first mapped to an orthogonal space. This mapping decorrelates the features, so they can be evaluated and selected individually. After the selection, the features are linked back to the same number of variables in the original measurement space. Using a mapping to orthogonal space proved to be very effective for features with high correlation. If the correlation between candidate features is only trivial, orthogonal transforms don't improve the results compared to existing sequential methods.

Greedy TODO Bla bla bla...

2.1.3 Embedded Methods

TODO Bla bla bla...

http://crpit.com/confpapers/CRPITV113Huda.pdf

As both filter and wrapper methods have different strengths and weaknesses, it seems only natural that research tried to combine both ideas to create so-called embedded methods, which are basically a hybrid of the former two. The strategy is to involve the classifier and its properties (like wrapper methods) while selection the features, and being computationally efficient at the same time (like filter methods).

Three types:

- Linear Classifiers (Pruning methods, SVM, logistic regression)
- ID3 / C4.5 with build-in mechanism (WTF?)
- Regularisation models

Regularisation models idea: fit the moddel, but try to have as many weightsas possible small or close to 0, so we can eliminate features with small/zero weights

TODO: more on embedded methods

2.2 Methods for Structured Features

TODO Bla Bla...es gibt die und die methods...

- Graph methods
- · Tree methods
- Group methods

2.3 Graph Methods

TODO Bla bla bla...

2.4 Tree Methods

TODO Bla bla bla...

2.5 Group Methods

TODO Bla bla bla...

2.6 Methods for Streaming Features

TODO Bla Bla...

Gender Prediction on Twitter TODO Bla bla...

NOTES AND LINKS:

Dont mix up FS with the classifiers (KNN, Naive bayes, etc)

PCA, Linear discriminant = feature EXTRACTION, brauch ma ned

Feature selection for document processing

3 Discussion

TODO Bla bla bla...

4 Conclusion

TODO Bla bla bla...

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