

Multi-Objective Approach for Support Vector Machine Parameter Optimization and Variable Selection in Cardiovascular Predictive Modeling

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Abstract: We present a heuristic-based approach for Support Vector Machine (SVM) parameter optimization and variable selection using a real-valued cooperative Multi-Objective Evolutionary Algorithm (MOEA). Due to the possibility to optimize several criteria simultaneously, we aim to maximize the SVM performance as well as minimize the number of input variables. The second criterion is important especially if obtaining new observations for the training data is expensive. In the field of epidemiology, additional model inputs mean more clinical tests and higher costs. Moreover, variable selection should lead to performance improvement of the model used. Therefore, to train an accurate model predicting cardiovascular diseases, we decided to take a SVM model, optimize its meta and kernel function parameters on a true population cohort variable set. The proposed approach was tested on the Kuopio Ischemic Heart Disease database, which is one of the most extensively characterized epidemiological databases. In our experiment, we made predictions on incidents of cardiovascular diseases with the prediction horizon of 7–9 years and found that use of MOEA improved model performance from 66.8% to 70.5% and reduced the number of inputs from 81 to about 58, as compared to the SVM model with default parameter values on the full set of variables.

1 INTRODUCTION

Cardiovascular diseases (CVDs) are one of the most frequent causes of people's deaths around the world for today. Stress, unhealthy diet, physical inactivity, harmful use of tobacco and alcohol increase the risk of CVDs significantly. Nowadays, even young people suffer from CVDs. According to the report of the World Health Organization, in 2015 about 17.7 million people died from CVDs (it was 31% of all global deaths) (World Health Organization, 2017). Early detection of a high CVD risk for a patient is considered to be the main issue for doctors to undertake appropriate measures in time and prevent non-fatal or fatal incidents of CVDs.

In this paper, we develop a predictive system based on a Support Vector Machine (SVM) and a

Multi-Objective Evolutionary Algorithm (MOEA), which is applied to tune SVM meta and kernel function parameters and select a proper set of input variables. There are many comparative studies in which SVMs outperform other predictive models on different problems, including medical diagnostics (Bellazzi and Zupan, 2008; Yu *et al.*, 2005). Moreover, this model copes successfully with a high-dimensional set of input variables (Ghaddar and Naoum-Sawaya, 2018), which is important for our study because the data used contains 81 variables.

A number of successful studies are devoted to optimizing SVM parameters by various heuristic approaches. In most cases, however, only one-criterion optimization algorithms are used (Ren and Bai, 2010; Liao *et al.*, 2015). In our study, we

employ a real-valued cooperative MOEA to optimize two criteria at once: the model predictive ability and the number of input variables (Cho and Hoang, 2017; Zhao *et al.*, 2011). In epidemiology, reducing the number of input variables is quite important because it leads to fewer clinical tests and lower costs for patients.

In medical data mining studies, researches often compare their proposals with conventional models on several test problems from repositories (Brameier and Banzhaf, 2001; Cheng *et al.*, 2006; Tu *et al.*, 2009). However, our goal is not to compare different models but to take the first step in building a predictive model based on real “raw” high-dimensional data. The presented model has been trained and tested on the Kuopio Ischemic Heart Disease (KIHD) dataset, which is one of the most properly characterized epidemiological study populations with a huge variety of variables: biomedical, psychosocial, behavioral, clinical and other. Previously, some of these variables have been pre-selected and used in risk factor analysis. In our approach, we do not involve experts to pre-select explanatory variables but perform variable selection algorithmically.

The next sections describe the approach proposed in detail, experimental results, conclusions, and future plans.

2 PREDICTIVE MODELING

SVM models are widely used in machine learning to solve classification and regression problems from various practical areas (Boser *et al.*, 1992). Generally, training SVM models is performed by minimizing the error function (1):

$$\frac{1}{2} \mathbf{w}^T \cdot \mathbf{w} + C \cdot \sum_{i=1}^N \xi_i \rightarrow \min, \quad (1)$$

which is subject to the constraints: $y_i(\mathbf{w}^T \cdot \mathbf{x}_i + b) \geq 1 - \xi_i$ and $\xi_i \geq 0$, $i = 1, \dots, N$, where C is an adjustable parameter of regularization, ξ_i expresses an error $\max(0, 1 - y_i \cdot (\mathbf{w}^T \cdot \mathbf{x}_i + b))$ on training examples (\mathbf{x}_i, y_i) , $y_i \in \pm 1$, $i = 1, \dots, N$, N is the number of training examples.

These models are more complex in comparison with a linear regression and allow reflecting non-linear dependencies due to the ‘kernel trick’, i.e. kernel functions map points into a higher dimensional space, where a linear separation is

applicable. In this work, we use a Radial Basis Function as a kernel (2):

$$K(\mathbf{x}_{i1}, \mathbf{x}_{i2}) = \exp(-\sigma \cdot \|\mathbf{x}_{i1} - \mathbf{x}_{i2}\|^2). \quad (2)$$

SVMs have a strong theoretical background and, in general, training these models is reduced to solving a dual quadratic programming problem with one global optimum.

By this time, a number of effective approaches to solve this quadratic programming problem have been proposed, for example, Sequential Minimal Optimization (SMO) (Platt, 1998). However, there are still some parameters which require proper tuning. They are meta (C) and kernel function parameters (σ) and, as can be found in other studies (Liao *et al.*, 2015; Syarif *et al.*, 2016), an arbitrary choice of their values may lead to a significant deterioration in the solution quality. The easiest way to tune these parameters is to use a grid search, but it requires quite a lot of computational time to check a good amount of values. As an alternative, heuristic optimization methods might be applied to find a pseudo-optimal combination of parameter values.

Evolutionary Algorithms (EA) operate with a set of candidate-solutions, which allows investigating a search space in a parallel way. One of important benefits of using EAs in optimizing SVM parameters is a possibility to incorporate variable selection into parameter tuning. This leads not only to finding proper values of SVM meta and kernel function parameters but also to determination of an effective input variable set corresponding to these particular SVM parameters. In our study, we propose to apply a MOEA to optimize two criteria simultaneously (3). The first criterion reflects the model predictive ability and the second one expresses the number of selected variables $N_{selected}$:

$$\begin{aligned} f_1 &= 1 - F_score \rightarrow \min; \\ f_2 &= N_{selected} \rightarrow \min. \end{aligned} \quad (3)$$

In the first criterion, we estimate the F-score metric (Goutte and Gaussier, 2005), specifically the F_1 -measure with equally weighted precision and recall.

To solve this two-criterion optimization problem (3), we have developed a real-valued cooperative modification of the Strength Pareto Evolutionary Algorithm 2 (SPEA2) (Zitzler *et al.*, 2002). In this algorithm, a chromosome consists of real-valued genes which code SVM parameters C , σ , and input

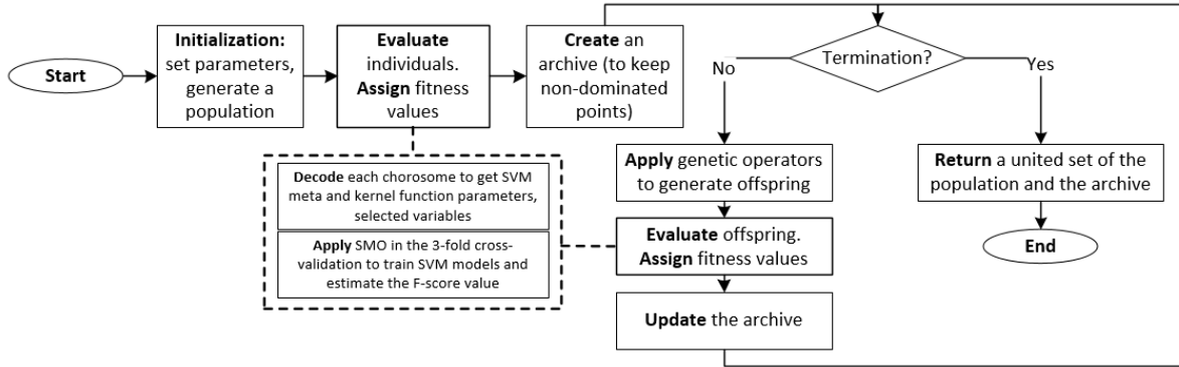


Figure 1: SVM training with a cooperative SPEA2.

variables. To evaluate criteria f_1 and f_2 for each chromosome, it should be decoded in the following way:

$$C = gene_1; \sigma = gene_2; \\ x_j = \begin{cases} \text{selected, if } gene_{j+2} \geq 0.5, \\ \text{not selected, if } gene_{j+2} < 0.5. \end{cases} \quad (4)$$

Then, for each pair (C, σ) with a certain set of selected variables, the SVM model is trained using the SMO algorithm. The F-score metric is estimated in the 3-fold cross-validation procedure on the training data.

The modified SPEA2 is based on an island model cooperation (Whitley *et al.*, 1997), which allows us to reduce computational time because a number of populations evolve in a parallel way. In addition to that, cooperative modifications of EAs demonstrate higher performance in comparison with their original versions (Brester *et al.*, 2018).

Available resources are divided among subpopulations (islands) equally: $M_{isl} = \frac{M}{L}$, where

M is the total number of individuals, L is the number of subpopulations. One crucial concern in the island cooperation is a migration process, which implies that in some m_r iterations (generations) islands exchange the best m_c individuals. In the proposed modification, the migration set replaces individuals with the worst fitness values in each island population. Thus, we have implemented a fully connected topology: each island sends its best solutions to all the other ones and, as a response, it receives solutions from other islands. The final solution is obtained by merging all the populations and archives, and, then, selecting the best solution based on the f_1 criterion (Figure 1).

Originally, SPEA2 operates with binary strings, however, for real-valued optimization problems a number of genetic operators have been developed.

To select effective solutions for the offspring generation, we apply binary tournament selection. As a crossover operator, we use intermediate recombination. In a mutation operator, we implement the next scheme (Liu *et al.*, 2009):

$$x_j' = \begin{cases} x_j + \gamma_j \cdot (b_j - a_j), & \text{with probability } p_m \\ x_j, & \text{with probability } 1 - p_m \end{cases}, \quad (5)$$

with

$$\gamma_j = \begin{cases} (2 \cdot rand)^{\frac{1}{\eta+1}} - 1, & \text{if } rand < 0.5 \\ 1 - (2 - 2 \cdot rand)^{\frac{1}{\eta+1}}, & \text{otherwise} \end{cases}, \quad (6)$$

where $rand$ is a uniformly random number $[0, 1]$. There are two control parameters: the mutation rate $p_m = \frac{1}{n}$, where n is the chromosome length, and the distribution index η is equal to 1.0 (6). a_j and b_j are the lower and the upper bounds of the j -th variable in the chromosome (5).

3 DATABASE DESCRIPTION

The epidemiologic ongoing cohort study, KIH (Kuopio Ischemic Heart Disease), was started in 1984 to investigate risk factors of CVDs and some other diseases in the population of Eastern Finland, where one of the highest rate of coronary heart disease (CHD) was recorded (Salonen, 1988).

Table 1: The KIHD data description.

| Examinations | Period | Participants | Variables |
|--------------|-----------|--------------|-----------|
| Baseline | 1984-1989 | Men | 8000 |
| 4-year | 1991-1993 | Men | 5000 |
| 11-year | 1998-1999 | Men, women | 3000 |
| 20-year | 2006-2008 | Men, women | 750 |

In Table 1, we present a timeline of the KIHD study, which currently consists of four examination periods. At the baseline time point (1984–1989), 2,682 middle aged (42, 48, 54, 60 years) recruited men from the city of Kuopio and its surrounding communities of Eastern Finland were randomly chosen for participation in the KIHD study. Later, in 1998-2001, 920 ageing recruited women joined this follow-up study.

This dataset is one of the most thoroughly characterized epidemiologic study populations in the world, with thousands of biomedical, psychosocial, behavioural, clinical, and other variables in its dataset. The KIHD study, as a valuable source for epidemiologic research, has attracted many scientists, which has yielded in more than 500 original peer reviewed articles in international scientific journals over the past 30 years. However, there are no studies yet that would try to use this unique database for predicting the appearance of CVDs without any variable pre-selection.

In spite of the fact that, originally, the main focus in the KIHD study was on CVDs, and especially on ischemic heart disease, other health outcomes such as cancer, diabetes, and dementia, have been also investigated (Kurl *et al.*, 2015; Tolmunen *et al.*, 2014, Virtanen *et al.*, 2016; Brester *et al.*, 2016).

In our predictive modeling, we engage the 20-year examination data with a representative subset of variables preselected by an experienced epidemiologist. This data contains information about incidents and fatal events of various diseases by 2015. In this work, we consider only CVD-related outcomes (stroke, CHD, acute myocardial infarction (AMI), etc.) All the subjects were labelled as ‘healthy’ (none of CVDs occurred) and ‘unhealthy’ (any incident of CVDs or fatal CVD event occurred).

Moreover, we applied the following pre-processing:

- 1) Subjects who had any CVD in their anamnesis at the 20-year examination were excluded so that we got state vectors (vectors of variables) for people who were healthy at the beginning of the observation period;
- 2) Subjects with more than 50% of missing values in the state vector and, then, variables

with more than 30% of gaps were removed. The rest of missing values were filled with a nearest neighbour method (Beretta and Santaniello, 2016).

Thus, all these pre-processing steps led to the dataset with 778 subjects and 81 variables: 360 sick subjects and 418 healthy subjects. In the next section, we present the experimental results of our modeling aimed at making accurate predictions for the subjects about CVDs by 2015 bases on their state vectors recorded in 2006–2008.

4 EXPERIMENTS AND RESULTS

Before training predictive models on the KIHD data, we checked if there were outliers or not. Data was collected from different sources, some variables were measured, others were recorded from questionnaires. It is clear that there might be mistakes. Moreover, there might be even wrong labels because people do not always go to the hospital if they have some CVD symptoms. Therefore, firstly, we decided to estimate Cook’s distance, which is used in Regression Analysis to find the most influential points in the sample (Cook, 1977). Cook’s distance for the *i*-th sample point is calculated based on the difference between a linear regression trained on the whole dataset and a regression model trained on the dataset after removing the *i*-th sample point. The higher Cook’s distance is, the more influential point we have.

In Figure 2 and 3, we demonstrate Cook’s distance values for KIHD subjects and Cook’s distance distribution in a form of a histogram.

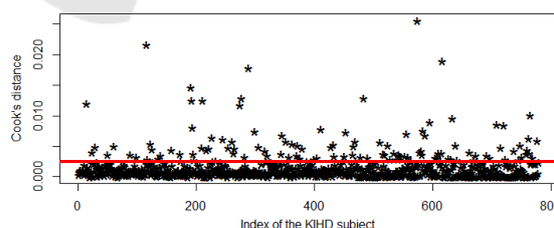


Figure 2: Cook’s distances for KIHD subjects.

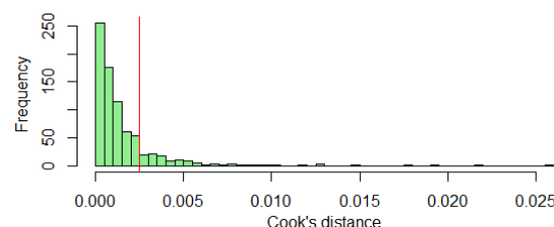


Figure 3: Cook’s distance distribution estimation.

In theory, there is no strict rule how to choose a threshold to determine outliers. After a number of experiments, we decided to remove KIHD subjects with Cook's distance which is higher than 0.0025, so that to keep 85% of the sample.

The SVM model performance was assessed in the 5-fold cross-validation procedure. In Figure 4, we compare F-score values before and after removing the most influential points. In this experiment, default values of C and σ were 1.0 and 0.01, correspondingly (these values are used in WEKA package (Hall *et al.*, 2009)).

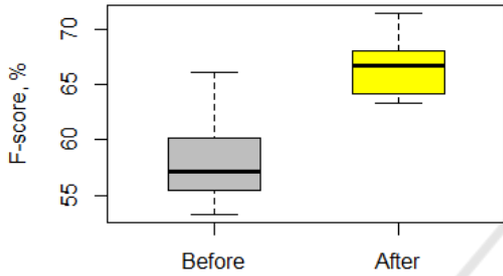


Figure 4: SVM performance with default parameters before and after removing outliers.

In the next experiment, we applied the modified SPEA2 to optimize SVM parameters and perform variable selection. The algorithm parameters were assigned as follows: $M_{isl} = 18$, $T = 20$ (the number of generations), $L = 3$, $m_r = 4$, $m_c = 2$, $\bar{M}_{isl} = 5$ (the archive size), $a_1 = 0.1$, $b_1 = 10$ (bounds of possible C values), $a_2 = 0.001$, $b_2 = 0.1$ (bounds of possible σ values), $a_j = 0$, $b_j = 1$, $j = \overline{3, n}$. To start the search from larger variable sets and prevent the algorithm from premature convergence to solutions with very few selected variables, in the initial population we generated the j -th gene ($j = \overline{3, n}$) based on the rule (7):

$$gene_j = \begin{cases} rand(0;0.5), & \text{with probability of } 0.2 \\ rand(0.5;1), & \text{with probability of } 0.8 \end{cases} \quad (7)$$

For each fold in the cross-validation procedure, we obtained the following optimized parameter (C, σ) values: 1 – (5.6876, 0.0541), 2 – (6.0277, 0.0536), 3 – (7.5469, 0.0387), 4 – (6.2696, 0.0380), 5 – (4.8698, 0.0771). The number of selected variables was equal to 57.8 averaged over 5 folds. In Figure 5, we compare F-score values achieved by

SVM models with default and optimized parameters, on the full and reduced variable sets.

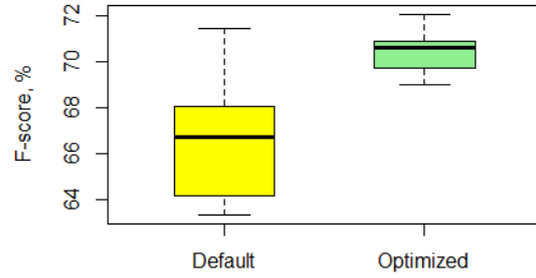


Figure 5: SVM performance with default parameters on the full dataset and optimized parameters on the selected variable set.

Thus, in this experiment we increased the average F-score value from 66.8 % up to 70.5% with the MOEA use.

At this point, we chose one best solution from all variants returned by MOEA islands. Indeed, we offer to return populations and archives with non-dominated individuals, as can be seen in Figure 1, because this allows us to choose not only one but also a number of good solutions which might be included in the ensemble of SVM models. This idea should be taken into account as a possible way to improve the achieved performance.

5 CONCLUSION

In this study, we introduced the effective approach aimed at training SVM models with optimized parameters and performing variable selection at once. Simultaneous optimization of two criteria became possible owing to the MOEA use, namely the real-valued cooperative SPEA2. The island model cooperation allowed us not only to reduce computational time but also to increase the performance of the original SPEA2.

Practically, our proposal was applied to cardiovascular predictive modeling: we investigated its effectiveness on the thoroughly characterized epidemiological data containing many different subjects and variables. Typically, CVDs are predicted by using a few pre-selected, already known, explanatory variables. Our approach diminishes the need for pre-selection and, simultaneously, may reveal novel previously unknown explanatory variables. On average, we managed to achieve 70.5% of F-score, which, definitely, should be improved later. However, applying the MOEA we could obtain 5.5% of the

relative improvement in F-score and accomplish variable selection. All in all, this study combines for the first time the real world epidemiological data, the advanced EA-based optimization and database pre-processing using Cook's distance.

Currently, we have predicted CVDs for the next 7-9 years with one time point predictors, whereas, in the future we plan to test the presented approach in multiple time point predictor data (baseline, 4-year, 11-year examinations) and make predictions for longer periods.

Moreover, at the next step, we should take advantage of the MOEA distinctive feature to return a number of non-dominated points, which might be involved in the ensemble of SVM models.

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