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NOVEL SUPPORT VECTOR MACHINES FOR DIVERSE LEARNING PARADIGMS

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of Philosophy at Virginia Commonwealth University.

by

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Abstract

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This dissertation introduces novel support vector machines (SVM) for the following traditional and non-traditional learning paradigms: *Online Learning*, *Multi-Target Regression*, *Multiple-Instance* classification, and *Data Stream* classification.

Three multi-target support vector regression (SVR) models are first presented. The first involves building independent, single-target SVR models for each target. The second builds an ensemble of randomly chained models using the first single-target method as a base model. The third calculates the targets' correlations and forms a maximum correlation chain, which is used to build a single chained SVR model, improving the model's prediction performance, while reducing computational complexity.

Under the multi-instance paradigm, a novel SVM multiple-instance formulation and an algorithm with a bag-representative selector, named MIRSVM, are presented. The contribution trains the SVM based on bag-level information and is able to identify instances that highly impact classification, i.e. bag-representatives, for both positive and negative bags, while finding the optimal class separation hyperplane. Unlike other multi-instance SVM methods, this approach eliminates possible class imbalance issues by allowing both positive and negative bags to have at most one representative, which constitute as the most contributing instances to the model.

Add OLLAWV and DS Application

Rigorous experimental studies and statistical analyses over various metrics and datasets were conducted in order to comprehensively compare the proposed solutions against modern, widely-used methods from all paradigms. The experimental studies and analyses confirm that the proposals achieve better performances and more scalable solutions than the methods compared, making them competitive in their respected fields.

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CHAPTER 1

INTRODUCTION

In traditional classification and regression problems, learning algorithms uncover dependencies and patterns that exist between given inputs (samples) and their outputs (categorical or continuous), using training data. Identifying these patterns is a non-trivial task due to many factors, such as the high dimensionality of the data, as well as dataset size. Over the past decade, dataset sizes have grown disproportionately to the speed of processors and memory capacity, limiting machine learning methods to computational time. Many real-world applications, such as human activity recognition, operations research, and video/signal processing, require algorithms that are scalable and accurate, while being able to provide insightful information in a timely fashion.

More recently, these traditional methods have been extended to accommodate various types of data paradigms. Examples include *Multiple Target* (MT) learning, *Multiple Instance* (MI) learning, and *Data Stream* learning. These emerging paradigms require algorithms to be robust and accommodate non-traditional data representations.

Multi-target learning is a challenging task that consists of creating predictive models for problems with multiple outputs [8, 23, 82]. MT learning includes *multi-target regression* (MTR), which addresses the prediction of continuous targets, *multi-label classification* [98] which focuses on binary targets, and *multi-dimensional classification* which describes the prediction of discrete targets [14]. This dissertation will be tackling the multi-target regression problem. Multi-target prediction has the capacity to generate models representing a wide variety of real-world applications, ranging from natural language processing [51] to bioinformatics [63]. A characteristic of the MT datasets used in these applications is that they are generated by a single system, indicating that the nature of the outputs captured has some structure. Even though modeling the multi-variate nature and possible complex rela-

tionships between the target variables is challenging, they are more accurately represented by a multi-target model [23, 38].

Several methods have been proposed for solving such multi-target tasks and can be categorized into two groups. The first being *problem transformation* methods in which the multi-target problem is transformed into multiple single-target (ST) problems, each solved separately using standard classification and regression algorithms. The second being *algorithm adaptation* methods which adapt existing traditional algorithms to predict all the target variables simultaneously [14]. It is known that *algorithm adaptation* methods outperform *problem transformation* methods. The most valuable advantage of using multi-target techniques is that, not only are the relationships between the sample variables and the targets exploited, but the relationships between the targets amongst themselves are as well [8, 23]. Single-target techniques, on the other hand, eliminate any possibility of learning from the possible relationships between the target variables because a single, independent model is trained for each target separately. Another advantage of MT techniques is model interpretability [9]. A single multi-target model is highly more interpretable than a series of single-target models. In addition, MT models could also be considerably more computationally efficient to train, rather than training multiple single-target models individually [39].

Multi-instance learning (MIL) is a generalization of supervised learning that has been recently been gaining interest because of its applicability to many real-world problems such as image classification and annotation [47], human action recognition [96], and drug activity prediction [32]. The difference between MIL and traditional learning is the nature of the data. In the multi-instance setting, a sample is considered a *bag* that contains multiple *instances* and is associated with a single label. The individual instance labels within a bag are unknown and bag labels are assigned based on a multi-instance assumption, or hypothesis. Introduced by Dietterich et. al. [32], the standard MI assumption states that a bag is labeled positive if and only if it contains at least one positive instance. Other hypotheses and frameworks have been proposed by Foulds and Frank [40] to encompass a wider range of applications with MI

data, but for the scope of this thesis, the focus will be on the standard MI assumption.

One of the major complexities associated with MIL is the ambiguity of the relationship between a bag label and the instances within the bag. This stems from the standard MI assumption, where the underlying distribution among instances within positive bags is unknown. There have been different attempts to overcome this complexity, such as “flattening” the MIL datasets, meaning instances contained in positive bags each adopt a positive label, allowing the use of classical supervised learning techniques [72]. This approach assumes that positive bags contain a significant number of positive instances, which may not be the case, causing the classifier to mislabel negative instances within the bag, decreasing the power of the MI model. To overcome this issue, a different MIL approach was proposed, where subsets of instances are selected from positive bags for classifier training [64]. One drawback of this type of method is that the resulting training datasets become imbalanced towards positive instances. Model performance further deteriorates when more instances are selected as subsets than needed [22].

The contributions of this thesis aim to deal with the drawbacks that exist within these non-traditional learning paradigms, using traditional and novel solvers for support vector machines. Support vector machines (SVM), proposed by Cortes and Vapnik [30], represent popular linear and non-linear (kernelized) learning algorithms based on the idea of a large-margin classifier. They have been shown to improve generalization performance for binary classification problems. SVMs are similar to other machine learning techniques, but literature shows that they usually outperform them in terms of scalability, computational efficiency, and robustness against outliers. They are known for creating sparse and non-linear classifiers, making them suitable for handling large datasets. A traditional approach for training SVMs is the *Sequential Minimal Optimization* (SMO) algorithm [71], a method for solving the L1-SVM’s Quadratic Programming (QP) task. Although SMO provides an exact solution to the SVM QP problem, its performance is highly dependent on the SVM hyperparameters. More recent approaches, which have been shown to surpass SMO in terms of scalability while

remaining competitive in accuracy, include the *Minimal Norm SVM* (MNSVM) [81] and the *Non-Negative Iterative Single Data Algorithm* (NNISDA) [57, 102]. With all the various efforts aimed at solving the SVM task efficiently, this area is still requires investigation.

To deal with issues of scalability, this thesis introduces a different approach which focuses on the minimization of the regularized L1-SVM through Stochastic Gradient Descent (SGD), a well-known simple, yet efficient technique for learning classifiers under convex loss functions. Recently, SGD algorithms have been shown to have considerable performance and generalization capabilities in the context of large-scale learning [16], and have been used to solve the SVM problem, such as *NORMA* [60] and *PEGASOS* [78, 100]. Although stochastic and iterative algorithms are very simple to implement and efficient, they also have their limitations. One of these limitations is the lack of meaningful stopping criteria for the algorithm; without a pre-specified number of iterations to train, the algorithms continue running [70]. Another limitation stems from the superlinear increase in training time as the number of samples increases. Incremental algorithms attempt to alleviate this issue, but they cannot guarantee a bound on the number of updates per iteration.

In this thesis, various approaches are devised for solving machine learning problems for various types of learning paradigms using traditional support vector machine solvers. Specifically, novel algorithms are proposed for solving multi-target regression (a subset of multi-target learning) and multi-instance classification problems.

1.1 Contributions of the Dissertation

The current leading MT models are based on ensembles of regressor chains, where random, differently ordered chains of the target variables are created and used to build separate regression models, using the previous target predictions in the chain. The challenges of building MT models stem from trying to capture and exploit possible correlations among the target variables during training, at the expense of increasing the computational complexity of model training. One of the contributions of this proposal aims to investigate the

performance changes when building a regression model using two distinct *algorithm adaptation* chaining methods versus building independent single-target models for each target variable using a novel framework. Specifically, this MTR contribution includes:

- Evaluating the performance of a Support Vector Regressor (SVR) as a multi-target to single-target *problem transformation* method to determine whether it outperforms current popular ST algorithms. Its performance is analyzed as a base-line model for MT chaining methods due to the fact that ST methods do not account for any correlation among the target variables.
- Building an MT ensemble of randomly chained SVR models (SVRRC), an *algorithm adaptation* approach, inspired by the chaining classification method, Ensemble of Random Chains Corrected (ERCC) [79], to investigate the effects and advantages of exploiting correlations among target variables during model training, in the context of regression problems. The main issues to be investigated with this approach are the *randomness* of the created chains because they might not capture of correlations between the targets, as well as the time taken to build all the regressors in the ensemble.
- Proposing an MT *algorithm adaptation* model of SVRs that builds a unique chain, capturing the maximum correlation among target outputs, named SVR Correlation Chains (SVRCC). The advantages of using this approach include exploiting the correlations among the targets which leads to an improvement in model prediction performance, and a reduction in computational complexity because a single SVR-chain model is trained, rather than building an ensemble of 10 base regressors.

To address the limitations presented by MIL algorithms, this thesis proposes a novel SVM formulation with a bag-representative selector, called Multiple-Instance Representative Support Vector Machine (MIRSVM). The algorithm does not assume any distribution of the instances and is not affected by the number of instances within a bag, making it applicable to a variety of contexts. The key contributions of this work include:

- Reformulating the traditional primal L1-SVM problem to optimize over bags, rather than instances, ensuring all the information contained within each bag is utilized during training, while defining bag representative selector criteria.
- Deriving the dual multi-instance SVM problem, with the Karush-Kuhn-Tucker necessary and sufficient conditions for optimality. The dual is maximized with respect to the Lagrange multipliers and provides insightful information about the resulting sparse model. The dual formulation is kernelized with a Gaussian radial basis function, which calculates the distances between bag representatives.
- Devising a unique bag-representative selection method that makes no presumptions about the underlying distributions of the instances within each bag, while maintaining the default MI assumption. This approach eliminates the issue of class imbalance caused by techniques such as flattening or subsetting positive instances from each bag. The key feature of MIRSVM is its ability to identify instances (support vectors) within positive and negative bags that highly impact the model.

To address the limitations presented by current popular SVM solvers, this paper proposes a novel *OnLine Learning Algorithm using Worst-Violators* (OLLAWV). This unique method iterates over samples, updates the model, and utilizes a novel stopping criterion. The model is updated by iteratively selecting (without replacement) the worst violating sample, i.e. the sample with the largest error according to the current decision function, and stops training when there are no more violating samples left to update. In other words, the algorithm is implicitly identifying support vectors and stopping when it has found them all. Because samples are selected and updated without replacement, coupled with the fact that the maximum number of iterations never exceeds the size of the dataset, OLLAWV does not use the regularization updating term. Instead, the regularization is achieved by early stopping. In [29], it has been shown that the smaller the number of updates (determined by here presented stopping criterion) is, the larger will be the margin. On the other hand,

the larger the margin, the better generalization of the model is. The experimental results presented here confirm both the theoretical statements in [29] and the validity of the approach proposed and taken in OLLAWV algorithm. Combining this method of updating the model and stopping criteria with the fact that SVMs are known for creating sparse kernel classifiers, the contribution aims to speed up the model training time without sacrificing the model’s accuracy. The key contributions of this work include:

- Devising a unique iterative procedure for solving the L1-SVM problem, as well as a novel method for identifying support vectors, or *worst-violators*. Rather than randomly iterating over the data samples, OLLAWV aims to reduce training time by selecting and updating the samples that are most incorrectly classified with respect to the current decision hyper-plane.
- Designing a novel stopping criteria by utilizing the worst-violator identification method. This aims to eliminate the added parameterization that is included with most online methods, where the number of iterations of the algorithm needs to be set in advance. Once there are no incorrectly classified samples left, the algorithm terminates.

CHAPTER 2

BACKGROUND

Support vector machines represent a popular set of learning techniques that have been introduced under Vapnik-Chervonenkis theory of *structured risk minimization* (SRM) [15, 30, 54, 75, 76]. SRM is an inductive principle for the purpose of model selection. It minimizes the expected probability of error, resulting in a generalized model, without making assumptions about the data distribution [76, 88]. This is the basis for developing the maximal margin classifier [88]. Based on the work of Aizerman et. al. [1], Boser et. al. [15] generalized the linear algorithm to the non-linear case. Then, Cortes and Vapnik [30] proposed the soft margin SVM; a modification that not only allowed maximal margin classifiers to be applied to non-linearly separable data, but also introduced a regularization parameter to prevent overfitting and gauge generalizability. That same year, the algorithm was extended by Vapnik et. al. [89] to the regression case.

This chapter presents a theoretical background of support vector machines. First, the SVM paradigm is discussed in the context of classification, introducing the concepts of the maximal margin, the linear *Soft Margin SVM* for overlapping classes, and its kernelized version. Next, Vapnik's ϵ -insensitivity loss function and the concept of Support Vector Regression (SVR) are introduced. Afterwards, popular methods for solving the SVM problem are presented and their advantages and problems are discussed. Finally, insights into the benefits of using SVMs are presented, along with a short comparison of SVMs to the classical statistical learning paradigm.

2.1 Support Vector Machine Classification

Supervised learning is the process of determining a relationship $f(\mathbf{x})$ by using a training dataset $\mathcal{S} = \{(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_i, y_i), \dots, (\mathbf{x}_n, y_n)\}$, which contains n inputs of d -dimensionality,

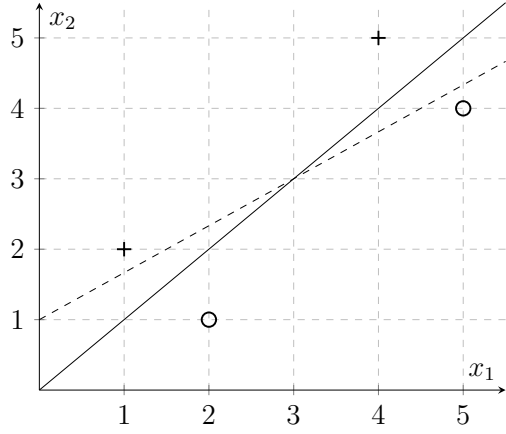


Fig. 2.1.: A 2-dimensional example of different possible separating hyperplanes that correctly classify all the toy data points.

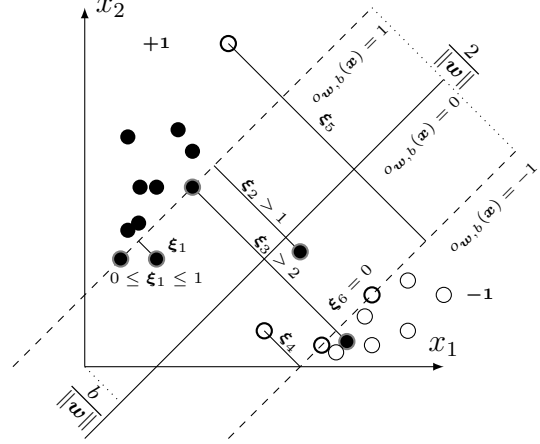


Fig. 2.2.: An illustration of the soft margin SVM solution on an example 2-dimensional non-linearly separable dataset.

$\mathbf{x}_i \in \mathbb{R}^d$, and their class labels y_i . In the case of binary classification, $y_i \in \{+1, -1\}$, where $+1$ and -1 are the two class labels.

The goal of the soft margin SVM classifier is to find a classification function

$$f(\mathbf{x}) = \text{SIGN } o_{\mathbf{w},b}(\mathbf{x}), \quad (2.1)$$

where $o_{\mathbf{w},b}(\mathbf{x}) = \mathbf{w} \cdot \mathbf{x}_i + b$ is a linear decision (output) function representing an affine mapping function $o : \mathbb{R}^d \rightarrow \mathbb{R}$ and is parameterized by $\mathbf{w} \in \mathbb{R}^d$, the weight vector, and $b \in \mathbb{R}$, the bias term. In addition, \mathbf{w} and b must satisfy the following,

$$y_i (\mathbf{w} \cdot \mathbf{x}_i + b) \geq 1 - \xi_i, \forall i \in \{1, \dots, n\}, \quad (2.2)$$

where $\boldsymbol{\xi} \in \mathbb{R}^n$ are the non-negative slack variables that allow for some classification error to account for overlapping datasets. The minimal distance between points belonging to opposite classes and the hyperplane is defined as the *margin* and has a width equal to $\frac{2}{\|\mathbf{w}\|}$, which is why the $\|\mathbf{w}\|$ must be minimal in order to maximize the margin.

In the example shown in Figure 2.1, if the training data points are slightly moved, the

solid line (with the larger margin) will still correctly classify all the instances, whereas the dotted line (with a much smaller margin, comparatively) will not. This illustrates that the location of the hyperplane has a direct impact on the classifiers generalization capabilities. The hyperplane with the largest margin is called the *optimal separating hyperplane*. Figure 2.2 shows the optimal separating hyperplane for overlapping training data points, where the filled data points are from the $+1$ class and the non-filled data points are from the -1 class. The training data points on the separating hyperplane (the circled data points), whose decision function value equals $+1$ or -1 , are called the *support vectors*.

The soft margin SVM problem is defined as follows,

$$\begin{aligned} \min_{(\mathbf{w}, b)} \quad & \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_{i=1}^n \xi_i \\ \text{s.t.} \quad & y_i (\mathbf{w} \cdot \mathbf{x}_i + b) \geq 1 - \xi_i, \quad \forall i \in \{1, \dots, n\} \\ & \xi_i \geq 0, \quad \forall i \in \{1, \dots, n\}, \end{aligned} \tag{2.3}$$

where the penalty parameter $C \in \mathbb{R}$ controls the trade-off between margin maximization and classification error minimization, penalizing large norms and errors.

Equation 2.3 can be rewritten as a regularized loss minimization problem by representing the constraints as the Hinge loss, given by:

$$L(y_i, o_{(w,b)}(\mathbf{x}_i)) = \max \{0, 1 - y_i o_{(w,b)}(\mathbf{x}_i)\}, \tag{2.4}$$

which penalizes errors satisfying the following: $y_i o_{(w,b)}(\mathbf{x}_i) < 1$ and is a crucial element that facilitates the SVM model's sparseness. The soft margin SVM represented as a regularized loss minimization problem becomes:

$$\min_{(\mathbf{w}, b) \in \mathcal{H}_o \times \mathbb{R}} R = \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_{i=1}^n L(y_i, o_{(w,b)}(\mathbf{x}_i)), \tag{2.5}$$

where \mathcal{H}_o is a general Hilbert space. To handle cases when the data are non-linearly separable, while enhancing the classifier's generalization capabilities, a kernel function can be

used [1], as shown in Equation 2.6:

$$\mathcal{K}(\mathbf{x}_i, \mathbf{x}_j) = \langle \phi(\mathbf{x}_i), \phi(\mathbf{x}_j) \rangle, \quad (2.6)$$

where $\phi(\cdot)$ represents a mapping function from the original feature space to a higher dimensional space. The advantage of utilizing kernels is being able to calculate the inner product in the input space rather than in the very high feature dimensional space (including the infinite dimensional ones). The SVM model output, o shown in Equation 2.7, for a given input vector \mathbf{x} is defined by the kernel as given below:

$$o(\mathbf{x}) = \sum_{i=1}^n \alpha_i \mathcal{K}(\mathbf{x}, \mathbf{x}_i) + b, \quad (2.7)$$

where $\alpha_i \in \mathbb{R}$ are the coefficients, or weights, of the expansion in feature space, and $b \in \mathbb{R}$ is the so-called bias term. Note that if a positive definite kernel is used, there is no need for a bias term b , but b can nevertheless be used. The two terms, $\boldsymbol{\alpha}$ and b , parametrize the SVM model. A model is called *dense* if the absolute value of all its weights are greater than 0, while a *sparse* model would be one that contains some $\alpha_i = 0$. The level of sparseness may vary, but the sparser the model, the more scalable the applications.

2.2 Support Vector Regression

The support vector machine was applied to the regression case [35, 87], maintaining all the maximal margin algorithmic features. Unlike pattern recognition problems where the desired outputs y_i are discrete, for the regression case they are continuous, real-valued, function outputs. Given training dataset $\mathcal{S} = \{(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_n, y_n) \in \mathbb{R}^d \times \mathbb{R}\}$, where $y_i \in \mathbb{R}$ is the continuous output of input $\mathbf{x}_i \in \mathbb{R}^d$, the goal is to learn a function $f(\mathbf{x})$ with at most ϵ deviation from the true targets y_i for all the training data, while being as flat as possible. This was introduced by Vapnik's linear loss function with ϵ -insensitivity zone, illustrated in

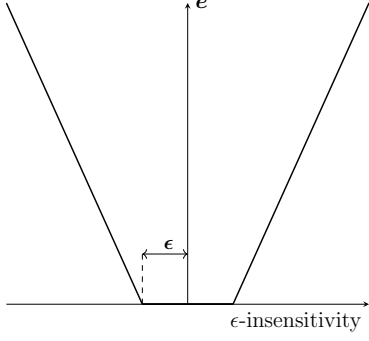


Fig. 2.3.: Vapnik's ϵ -insensitivity loss function.

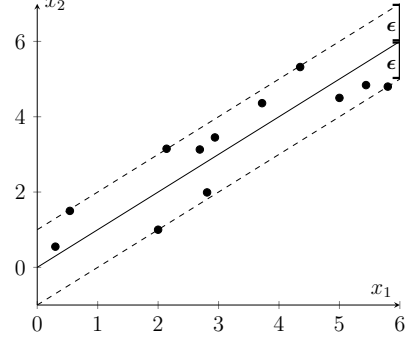


Fig. 2.4.: Support vector regression example solution.

Figure 2.3 and given by:

$$|y_i - o_{(w,b)}(\mathbf{x}_i)|_\epsilon = \begin{cases} 0 & \text{if } |y_i - o_{(w,b)}(\mathbf{x}_i)| \leq \epsilon \\ |y_i - o_{(w,b)}(\mathbf{x}_i)| - \epsilon & \text{otherwise.} \end{cases} \quad (2.8)$$

The loss is equal to 0 if the difference between the predicted and true output values is less than ϵ . Vapnik's ϵ -insensitivity function, shown in Equation 2.8, defines an ϵ -tube, illustrated in Figure 2.4. If the predicted value is within the tube, no loss is incurred [54]. Estimating a linear regression hyperplane is achieved by minimizing:

$$\min_{(w,b) \in \mathcal{H}_o \times \mathbb{R}} R = \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_{i=1}^n (|y_i - o_{(w,b)}(\mathbf{x}_i)|_\epsilon). \quad (2.9)$$

Equation 2.9 is equivalent to the following, where non-negative slack variables are introduced:

$$\begin{aligned} \min_{(w,b,\xi,\xi^*)} \quad & \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_{i=1}^n (\xi_i + \xi_i^*) \\ \text{s.t.} \quad & y_i - \mathbf{w} \cdot \mathbf{x}_i - b \leq \xi_i + \epsilon, \quad \forall i = \{1, \dots, n\} \\ & \mathbf{w} \cdot \mathbf{x}_i + b - y_i \leq \xi_i^* + \epsilon, \quad \forall i = \{1, \dots, n\} \\ & \xi_i, \xi_i^* \geq 0, \quad \forall i = \{1, \dots, n\}. \end{aligned} \quad (2.10)$$

Note that the constant C influences the trade-off between approximation error and the model generalizability, similar to the classification setting. The optimization function given in 2.10

can be solved more easily in its dual formulation and is key to extending the SVR to learn from non-linear functions [75]. It is as follows:

$$\begin{aligned}
& \max_{(\boldsymbol{\alpha}, \boldsymbol{\alpha}^*)} \quad -\frac{1}{2} \sum_{i,j=1}^n (\alpha_i - \alpha_i^*)(\alpha_j - \alpha_j^*) \mathcal{K}(\mathbf{x}_i, \mathbf{x}_j) - \epsilon \sum_{i=1}^n (\alpha_i + \alpha_i^*) + \sum_{i=1}^n y_i (\alpha_i - \alpha_i^*) \\
& \text{s.t.} \quad \sum_{i=1}^n (\alpha_i - \alpha_i^*) = 0, \alpha_i, \alpha_i^* \in [0, C], \forall i = \{1, \dots, n\},
\end{aligned} \tag{2.11}$$

where $\boldsymbol{\alpha}$ and $\boldsymbol{\alpha}^*$ correspond to the SVR dual variables.

2.3 Support Vector Machine Solvers

Although support vector machines represent a major development in machine learning algorithms, in the case of large-scale problems (hundreds of thousands to several millions of samples), the design of SVM training algorithms still has room for improvement.

Interior Point (IP) methods pose the SVM learning problem as a quadratic optimization problem subject to linear constraints, which are then replaced with a barrier function [20]. The resulting unconstrained problem can then be optimized using Newton or Quasi-Newton methods. However, IP methods typically require a run time of $[O(n^3)]$. Moreover, the memory requirements of IP methods are $[O(n^2)]$, rendering the use of IP methods impractical when the training set consists of a large number of samples.

The first attempts of overcoming the quadratic memory requirement of IP methods and speeding up their training time were aimed at decomposing the underlying SVMs quadratic programming problem. First, Boser et al. [15] implemented Vapnik's *chunking* method. *Sequential Minimal Optimization* (SMO) by Platt [71], its improvement by Keerthi et al. [58], and SVM-Light [52] are alternative approaches to *decomposing* the QP problem. SMO, implemented in the popular, widely used software package LIBSVM [24], is an iterative procedure that divides the SVM dual problem into a series of sub-problems, which are solved analytically by finding the optimal $\boldsymbol{\alpha}$ values that satisfy the Karush-Kuhn-Tucker conditions [20]. Although SMO is guaranteed to converge, heuristics are used to choose $\boldsymbol{\alpha}$ values in order to accelerate the convergence rate. This is a critical step because the convergence speed of the

SMO algorithm is highly dependent on the dataset size and SVM hyperparameters [75].

Most existing approaches, including the methods described above, focus on solving the dual of Equation 2.3, for two main reasons: firstly, the dual formulation provides a convenient way of dealing with the constraints. Secondly, the dual formulation can be written in terms of dot products, allowing the use of kernel functions [18]. However, these two conveniences are not restrictions for solving primal SVM problem. For example, Chappelle [25] showed that even though optimizing the primal and dual are equivalent in terms of solutions and time, optimizing the primal when dealing with approximate solutions is far more superior. The primal problem can be cast as an unconstrained problem by using linear or non-linear kernels and the Representer theorem [74]; mainly, reparametrizing the weight vector. Chappelle [25] investigated solving the primal objective with smooth loss functions, rather than using the hinge loss function, and suggested using methods such as conjugate gradient descent and Newton’s method.

Some advancements in handling large scale problems are based on a geometric interpretation of SVM problem. Some of these geometric SVMs include approaches that use convex hulls [10] and minimum enclosing balls such as *Core Vector Machines* (CVM) [84]. Tsang et al. [83] then improved the scalability of CVMs by introducing *Ball Vector Machines* (BVM) which do not require a QP solver. Other geometric approaches include the novel algorithms introduced by Strack [81], known as the *Sphere Support Vector Machine* (SphereSVM) and *Minimal Norm Support Vector Machine* (MNSVM), which utilize the connection between minimal enclosing balls and convex hull problems, while demonstrating a high capability for learning from large datasets.

The *Non-Negative Iterative Single Data Algorithm* (NNISDA) [102] is an efficient approach for solving the SVM problem, shown to be faster than SMO and equal in terms of accuracy [57]. NNISDA is an iterative algorithm that finds a solution to the L2-SVM using coordinate descent, inspired by *Iterative Single Data Algorithm* (ISDA) [50] which was originally introduced by Kecman et al. [55].

Recently, several authors have proposed the use of a standard stochastic (or online) gradient descent (SGD) approach for SVMs to optimize large-scale learning problems [46, 59, 75, 76, 78]. Kivinen et al. [60] and Bousquet and Bottou [19] showed that stochastic algorithms can be both the fastest, and have the best generalization performances. It has also been shown that the SGD runtime for solving the SVM unconstrained primal problem is inversely proportional to the size of the training set [70, 77]. Shalev-Shwartz and Ben-David [76] have demonstrated that the basic SGD algorithm is very effective when data are sparse, taking less than linear $[O(d)]$ time and space per iteration to optimize a system with d parameters. It can greatly surpass the performance of more sophisticated batch methods on large data sets. The previously mentioned approaches are extended variants of a classic kernel perceptron algorithm [29].

Notable representatives of this method of learning include the *Naïve Online R Minimization Algorithm* (NORMA) by Kivinen et al. [60] and the *Primal Estimated Sub-Gradient Solver for SVM* (PEGASOS) by Shalev-Shwartz et al. [78]. NORMA is an online kernel based algorithm designed to utilize SGD for solving the SVM problem, exploiting the kernel trick in an online setting. It can be regarded as a generalization of the kernel perceptron algorithm with regularization [60]. PEGASOS solves the primal SVM problem using stochastic sub-gradient descent, implementing both linear and non-linear kernels, and showed that the algorithm does not directly depend on the size of the data, making it suitable for large-scale learning problems.

A more recent approach, named *OnLine Learning Algorithm* (OLLA) [53] is a unification, simplification, and expansion of the somewhat similar approaches presented in [46, 59, 75, 76, 78] and [56, 67, 65]. This algorithm is unique because it is not only designed to optimize the SVM cost function, but also the cost functions of several other popular nonlinear (kernel) classifiers using SGD in the primal domain. Collobert and Bengio [29] provided justification for not using regularization, and thus OLLA was designed to handle cost functions with and without the regularization term. Comparisons of performances of OLLA with the

popular SMO algorithm highlighted the merits of OLLA in terms of speed, as well as accuracy, when the number of samples was increased, making it suitable for large-scale learning. Comparisons using various different classifiers against SMO were also shown [53], but for the scope of this thesis the L1-SVM was mentioned.

Although the SGD approaches mentioned above have many merits when it comes to solving large-scale machine learning problems, stochastic procedures also have their disadvantages. One of them stems from the lack of meaningful stopping criteria. The only specified stopping criteria is a user defined input for the number of iterations, which gives rise to the question of what it should be set to. Another unknown parameter that requires tuning is the gradient step size, which in some cases, directly affects the algorithm convergence rate. Moreover, a third disadvantage of kernelized online algorithms is that the training time for each update increases superlinearly with the number of samples [17].

2.4 Why Support Vector Machines: Form & Norm

The support vector machine problem shares similarities to classical statistical inference such as Neural Networks (NNs), however, there are several very important differences between their approaches and assumptions.

One of the major differences is that these traditional classification and regression statistical techniques are based on the strict assumption that the data distribution is known and is that of a Gaussian distribution. Another assumption is that this data can be modeled by a set of linear parameter functions. Following this, the induction paradigm for parameter estimation is the maximum likelihood estimation method, which can be reduced to the minimization of the sum-of-errors-squared cost function [54].

The previously stated assumptions, on which the classic statistical paradigm relied, turned out to be inappropriate for many contemporary problems for a couple of reasons [87]. Real-world problems are, more often than not, high-dimensional. If the underlying mapping is not smooth, the linear paradigm needs an exponentially increasing number of terms,

thus increasing the dimensionality of the input space, also known as ‘*the curse of dimensionality*’. Another issue is that the data generation process might be very different from the normal distribution. Due to these grave concerns, the maximum likelihood estimator or sum-of-errors-squared cost function, should be replaced by a new induction paradigm to be able to model non-Gaussian distributions, or rather, to have a distribution-free method of classification or regression for high-dimensional, sparse data [54]. This was the foundational reason for developing support vector machines.

The main differences between SVMs and classical statistical techniques, such as NNs, can be identified by analyzing their *form* and *norm*. With respect to both types of models’ *form*, all machine learning models are the same, i.e. sums of weighted basis functions. The difference in the two approaches stems from their *norm* (cost functions) and how these models learn their function parameters (e.g. how many functions should be used, what their parameters are, what the value of their weights should be).

For example, NNs minimize the sum-of-errors-squared in output space, i.e. the L2 norm, and SVMs maximize the margin in input space by minimizing the L2 norm of the weight vector. For SVMs, the model parameters are not predefined and their number depends on the training data used. Rather than choosing the appropriate structure of the model, keeping the estimation error fixed, and minimizing the training error, as done by classical techniques; SVMs keep the training error fixed or set to some appropriate level and minimize the estimation error. This is the paradigm of structural risk minimization (SRM) introduced by Vapnik and Chervonenkis and their colleagues, which led to the new learning algorithm. This approach has been proven, both experimentally and theoretically, to be superior (or comparable) to NNs and other statistical methods for contemporary real-world problems, which are often very sparse datasets (i.e. small number of samples in high dimensional spaces).

CHAPTER 3

MULTI-TARGET SVR USING MAXIMUM CORRELATION CHAINS

This contribution presents three multi-target support vector regression (SVR) models. The first involves building independent, single-target SVR models for each output variable. The second builds an ensemble of random chains using the first method as a base model, named SVR with Random Chains (SVRRC), inspired by the classification MT method, Ensemble of Random Chains Corrected (ERCC) [79]. The third calculates the targets' correlations and forms a maximum correlation chain, which is used to build a single chained model named SVR with Correlation Chaining (SVRCC). The experimental study compares the performance of the three approaches with six other prominent MT regressors. The experimental results are then analyzed using non-parametric statistical tests. The results show that the maximum correlation SVR approach improves the performance of using ensembles of random chains.

This chapter is organized as follows: Section 3.1 describes the notation used throughout this chapter and reviews related works on multi-target regression. Section 3.2 presents the three multi-target support vector regression approaches. Section 3.3 presents the experimental study. Section 3.4 discusses the results and the statistical analysis. Finally, Section 3.5 shows the main conclusions of this work.

3.1 Multi-Target Regression Background

This section first defines the notation that will be used throughout this chapter, and then formally describes the multi-target regression problem along with relevant popular algorithms used within this paradigm.

Table 3.1.: Multiple-Target Learning Notation

Definition	Notation
Number of Samples	\mathcal{N}
Number of Input Attributes	d
Input Space	$\mathbf{X} = \{\mathbf{X}_1, \dots, \mathbf{X}_i, \dots, \mathbf{X}_d\} \in \mathbb{R}^{\mathcal{N} \times d}, 1 \leq i \leq d$
Input Instance	$\mathbf{x}^{(l)} = (x_1^{(l)}, \dots, x_d^{(l)}) \in \mathbf{X}, 1 \leq l \leq \mathcal{N}$
Number of Dataset Targets/Outputs	m
Target Space	$\mathbf{Y} = \{\mathbf{Y}_1, \dots, \mathbf{Y}_j, \dots, \mathbf{Y}_m\} \in \mathbb{R}^{\mathcal{N} \times m}, 1 \leq j \leq m$
Predicted Target Space	$\hat{\mathbf{Y}} = \{\hat{\mathbf{Y}}_1, \dots, \hat{\mathbf{Y}}_j, \dots, \hat{\mathbf{Y}}_m\} \in \mathbb{R}^{\mathcal{N} \times m}, 1 \leq j \leq m$
Target Instance	$\mathbf{y}^{(l)} = (y_1^{(l)}, \dots, y_m^{(l)}) \in \mathbf{Y}, 1 \leq l \leq \mathcal{N}$
Full Multi-Target (MT) Training Dataset	$\mathcal{D} = \{(x_1^{(1)}, y_1^{(1)}), \dots, (x_d^{(\mathcal{N})}, y_m^{(\mathcal{N})})\}$
Single-Target (ST) Dataset with j^{th} Target	$\mathcal{D}_j = \{(x_1^{(1)}, y_j^{(1)}), \dots, (x_d^{(\mathcal{N})}, y_j^{(\mathcal{N})})\} \in \mathcal{D}, 1 \leq j \leq m$
Number of Cross-Validation (CV) Sets	k
ST Test Dataset with j^{th} Target, i^{th} CV Fold	$\mathcal{D}_j^{(i)} = \{(x_1^{(i)}, y_j^{(i)}), \dots, (x_d^{(i)}, y_j^{(i)})\} \in \mathcal{D}_j, i \in \{1, \dots, \mathcal{N}\}$
ST Training Dataset with j^{th} Target, Excluding the i^{th} CV Fold	$\mathcal{D}_j^{(k-i)} = \mathcal{D}_j \setminus \mathcal{D}_j^{(i)}$
ST Regression Model	$h : \mathbf{X} \times \mathbf{Y}$
MT Regression Model	$h_j : \mathbf{X} \times \mathbf{Y}_j, 1 \leq j \leq m$
Unknown Sample	$\mathbf{x}^{(\mathcal{N}')} = \{\mathbf{x}^{(\mathcal{N}+1)}, \dots, \mathbf{x}^{(\mathcal{N}')}\}$
Predicted Values for Unknown Sample	$\mathbf{y}^{(\mathcal{N}')} = \{\mathbf{y}^{(\mathcal{N}+1)}, \dots, \mathbf{y}^{(\mathcal{N}')}\}$

3.1.1 Notation

Let \mathcal{D} be a training dataset of n instances. Let $\mathbf{X} \in \mathcal{D}$ be a matrix consisting of d input variables and n samples, such that $\mathbf{X} \in \mathbb{R}^{n \times d}$. Let $\mathbf{Y} \in \mathcal{D}$ be a matrix consisting of m continuous target variables and n samples, where $\mathbf{Y} \in \mathbb{R}^{n \times m}$. Table 3.1 summarizes the notation used throughout this chapter.

3.1.2 Multi-Target Regression

As mentioned previously, there are two main approaches to solving multiple-output problems: problem transformation and algorithm adaptation.

Problem Transformation

Algorithm Adaptation

Using *problem transformation* algorithms for a domain of t target variables, t predictive models must be constructed, each predicting a single-target variable. Prediction for an unseen

sample would be obtained by running each of the t single-target models and concatenating their results. Conversely, when using *algorithm adaptation* algorithms for the same domain of t target variables, a single model would need to be constructed which would output all t predictions.

In the multi-target learning paradigm, the *problem transformation* approach involves training m independent, single-target models for each target output on datasets $\mathcal{D}_j = \{\mathbf{X}, \mathbf{Y}_j\}$, $\forall j \in \{1, \dots, m\}$, and is considered as a baseline for measuring model performance [79]. Many *problem transformation* methods have been proposed for solve multi-target problems, however, the main issue with this type of approach is that the relationships between the targets is lost once independent models are built for each target. Examples of *problem transformation* approaches include Linear Target Combinations for MT Regression [86], and Multi-Objective Random Forests (MORF) [61].

The RC, MTS, MTSC, ERC, and ERCC methods are introduced by Spyromitros et. al. in [79]. The idea behind these algorithms was to investigate whether advances in multi-label learning can be successfully used in a multi-target regression setting and shed light on modeling target dependencies. These methods involve two stages of learning, the first being building ST models. The second uses the knowledge gained by the first step to predict the target variables while using possible relationships the targets might have with one another.

The two stages of training in MTS involve firstly, training m independent single-target models, like in ST. In the second step, a second set of m meta models are learned for each target variable, \mathbf{Y}_j , $1 \leq j \leq m$. These meta models are learned on a transformed dataset, where the input attributes space is expanded by adding the approximated target variables obtained in the first stage, excluding the j^{th} target being predicted.

The ERC method is somewhat similar to the MTS method. In the training of a Regression Chain (RC) model, a random chain, or sequence, of the set of target variables is selected and for each target in the chain, models are built sequentially by using the output of the previous model as input for the next [80]. If the default, ordered chain is $C = \{\mathbf{Y}_1, \mathbf{Y}_2, \dots, \mathbf{Y}_m\}$,

the first model $h_1 : \mathbf{X} \rightarrow \mathbb{R}$ is trained for \mathbf{Y}_1 , as in ST. For the subsequent models $h_{j,j>1}$, the dataset is transformed by sequentially appending the true values of each of the previous targets in the chain to the input vectors. For a new input vector, the target values are unknown. So once the models are trained, the unseen input vector will be appended with the approximated target values, making the models dependent on the approximated values obtained in each step. One of the issues associated with this method is that, if a single random chain is used, the possible relationships between the targets at the head of the chain and the end of the chain are not exploited due to the algorithm’s sequential nature. Also, prediction error in the earlier stages of the models will be propagated as the rest of the models are trained, which is why the Ensemble of Regressor Chains was proposed in [79]. Instead of a single chain, k chains are created at random, and the final prediction values are obtained by taking the mean values of the k predicted values for each target.

In the methods described above, the estimated target variables (meta-variables) are used as input in the second stage of training. In both methods, the models are trained using these meta-variables that become noisy at prediction time, and thus the relationship between the meta-variables and target variable is muddled. Dividing the training set into sets, one for each stage, would not help this situation because both methods would be trained on training sets of decreasing size. Due to these issues, *Spyromitros et. al.* proposed modifications, in [79], to both methods that resembles k -fold cross-validation (CV) to be able to obtain unbiased estimates of the meta-variables. These methods are called Regression Chains Corrected (RCC) and Multi-Target Stacking Corrected (MTSC).

The ERCC and MTSC procedures involve repeating the RCC and MTS procedures k times, respectively, with k randomly ordered chains for ERCC, and k different modified training sets for MTSC. The algorithms were tested and compared using Bagging of 100 regression trees as their base regression algorithm with ERC and ERCC ensemble size of 10, and 10-fold cross-validation. The corrected methods exhibited better performance than their original variants, as well as ST models. The ERCC algorithm had the best overall per-

formance, as well as being statistically significantly more accurate of all the methods tested. These methods can be found and used through the open-source Java library, Mulan [85]; to replicate the results found in [79].

3.2 Three Novel SVMs for Multi-Target Regression

Three novel models have been implemented for the purposes of multi-target regression. The base model is the SVR model, where m single-target soft margin non-linear support vector regressors (NL-SVR) are built for each target variable \mathbf{Y}_j .

For NL-SVR, the regularized soft margin loss function given in equation (2.10) is minimized. This contribution involves solving the dual of this formulation given by (2.11). Using the dual formulation, the multi-target problem is solved by transforming it into m single-

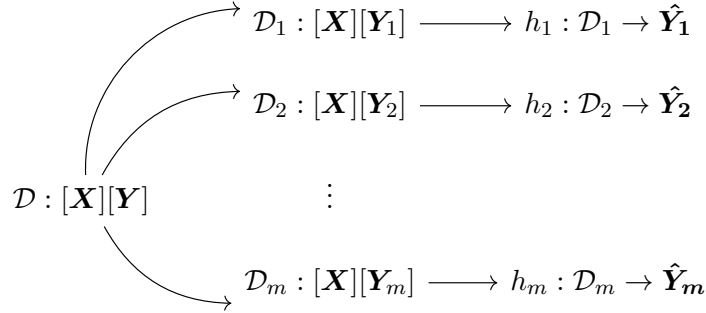


Fig. 3.1.: SVR Flow Diagram. Firstly, MT dataset is divided into m ST datasets, $\mathcal{D}_1, \mathcal{D}_2, \dots, \mathcal{D}_m$. Then m models, h_1, h_2, \dots, h_m , are independently trained for each ST dataset.

Algorithm 3.1 MT Support Vector Regression (SVR)

Input: Training dataset \mathcal{D}

Output: ST models $h_j, j = 1, \dots, m$

```

1: for  $j = 1$  to  $m$  do
2:    $\mathcal{D}_j = \{\mathbf{X}, \mathbf{Y}_j\}$  ▷ Get ST data
3:    $h_j : \mathbf{X} \rightarrow \mathbb{R}$  ▷ Build ST model for the  $j^{th}$  target
4: end for
5: return  $h_j, j = 1, \dots, m$ 

```

Algorithm 3.2 Build Chained Model

Input: Training dataset \mathcal{D} , random chain \mathbf{C} **Output:** A chained model $h_j, j = \{1, \dots, m\}, c \leq 10$

```
1:  $\mathcal{D}_1 = \{\mathbf{X}, \mathbf{Y}_{C_1}\}$  ▷ Initialize first dataset
2: for  $j = 1$  to  $m$  do ▷ For each target in chain  $\mathbf{C}$ 
3:    $h_j : \mathcal{D}_j \rightarrow \mathbb{R}$  ▷ Train model on appended dataset
4:   if  $j < m$  then
5:      $\mathcal{D}_{j+1} = \{\mathcal{D}_j, \mathbf{Y}_{C_j}\}$  ▷ Append new target in chain to dataset
6:   end if
7: end for
8: return  $h_j, j = 1, \dots, m$ 
```

target problems, as shown in Algorithm 3.1 and Figure 3.1. This algorithm will output m single-target models, $h_j, \forall j = 1, \dots, m$, for a given dataset \mathcal{D} . It first splits the dataset into m separate ones, \mathcal{D}_j , each with a single-target variable \mathbf{Y}_j , and then builds a distinct SVR model for each of the datasets.

Building m ST models is a good base-line, but as mentioned previously, it does not capture possible correlations between the target attributes during training. If these correlations are not exploited, this could retract from the model’s potential performance. Therefore, creating an ensemble model using a series of random chains was proposed, using the base-line SVR method, named SVR Random Chains (SVRRC).

For SVRRC, ensembles of at most 10 m -sized random chains, \mathcal{C} , are built from different and distinct permutations of the target variable indices. When chaining target values, there are two main options: using the predicted value as input for the following target, or using the true value of the target variable as input of the subsequent targets. The main problem with the former approach is that errors are propagated throughout the chained model, therefore SVRCC employs chaining of the true values.

For each random chain, a new model is trained by predicting the first target variable in the chain. Next, the first target’s true value, \mathbf{Y}_j , is appended to the training set. This chaining process is repeated for all the target indices in the chains, $\{\mathbf{C}_1, \dots, \mathbf{C}_c\} \in \mathcal{C}, c \leq 10$. This process will be repeated for each random chain generated, returning an ensemble of chained SVRs. Algorithm 3.2 describes the process of building a chained model given chain



Fig. 3.2.: SVRRC Flow Diagram on a dataset with 3 targets. SVRRC first builds the 6 random chains of the target's indices (3 examples are shown). It then constructs a chained model by proceeding recursively over the chain, building a model, and appending the current target to the input space to predict the next target in the chain.

Algorithm 3.3 MT SVR with Random-Chains (SVRRC)

Input: Training dataset \mathcal{D} , c random chains \mathcal{C}

Output: An ensemble of chained models $h_{\mathcal{C}}$

- 1: **for each** $\mathbf{C} \in \mathcal{C}$ **do** ▷ For each random chain
 - 2: $h_{\mathbf{C}} = \text{build chained model}(\mathcal{D}, \mathbf{C})$ ▷ build a chained model for chain \mathbf{C}
 - 3: **end for**
 - 4: **return** $h_{\mathcal{C}}$
-

$\mathbf{C} \in \mathcal{C}$, and Algorithm 3.3 shows the steps taken by SVRRC.

Given this ensemble of chained models, the predicted values for a given unseen instance are calculated by taking the mean of the multiple models generated using different random chains. Since the unseen input has no known target value, the predicted value at each step of the chain $\hat{\mathbf{Y}}_j$ is appended to the input at each step of the chain.

Due to the computational complexity of building $m!$ distinct chains and training $(m!) \times m$ models, the number of ensembles and chains are limited to a maximum of 10. However, if the number of target variables is less than 3, i.e. $m! \leq 10$, all $m!$ random chains are constructed.

A disadvantage of building an ensemble of 10 random chains stems from the fact that: when the number of output variables increases, the number of possible chains increases factorially. Therefore, there is no guarantee that the 10 random chains generated will truly

reflect the relationships among the target variables. Additionally, building an ensemble of regressors is computationally expensive. Finding a heuristic that allows the identification of a single, most appropriate chain, which fully reflects the output variable interrelations would improve the scalability of training the ensemble.

The third proposal was designed to remedy this issue. It builds a single chain based on the maximization of the correlations among the target variables. By calculating the correlation of the target variables and imposing it on the order of the chain, this ensures that each appended target provides some additional knowledge on the training of the next. With SVRRC, there is no reasoning behind the generation of these chains, and since the number of random chains generated is limited to 10, there is no way of ensuring that the 10 chains fully represent the targets' dependencies. Calculating and using the correlations of the targets would break this uncertainty. Algorithm 3.4 presents the SVR Correlation Chain (SVRCC) method. The computational complexity and hardware constraints (memory size) are negligible during the construction of the targets' correlation matrix, since the correlation matrix would be an $(m \times m)$ matrix, and the likelihood that the number of targets is large enough to cause a memory issue is minimal.

To calculate the correlation coefficients of the targets, the targets' co-variance matrix, Σ , is first calculated as shown in Equation 3.1:

$$\Sigma_{ij} = cov(\mathbf{Y}_i, \mathbf{Y}_j) = \mathbf{E}[(\mathbf{Y}_i - \mu_i)(\mathbf{Y}_j - \mu_j)], \quad (3.1)$$

where $\mu_i = \mathbf{E}(\mathbf{Y}_i)$, and $\mathbf{E}(\mathbf{Y}_i)$ is the expected value of \mathbf{Y}_i , $\forall i, j \in \{1, \dots, m\}$. This matrix will show how the targets change together.

The correlation coefficients matrix, \mathbf{P} , is then calculated as shown in Equation 3.2:

$$\mathbf{P} = corrcoeff(\mathbf{Y}) = \frac{\Sigma_{ij}}{\sqrt{\Sigma_{ii}\Sigma_{jj}}}, \forall i, j \in \{1, \dots, m\}. \quad (3.2)$$

It will describe the linear relationship among the target variables. The coefficients are then sorted in decreasing order, creating the maximum correlation chain.



Fig. 3.3.: SVRCC Flow Diagram on a sample dataset with 3 targets. SVRCC first finds the direction of maximum correlation among the targets and uses that order as the only chain. It then constructs the chained model as done in SVRRC.

Algorithm 3.4 MT SVR with Max-Correlation Chain (SVRCC)

- | | |
|---|--|
| 1: $\mathbf{P} = \text{corrcoef}(\mathbf{Y})$ | ▷ Find correlation coefficient matrix for target variables |
| 2: $\mathbf{C} = \sum_{i=1}^n \mathbf{P}_{ij}, \forall j = 1, \dots, m$ | ▷ Sum row elements of the correlation coefficient matrix |
| 3: $\mathbf{C} = \text{sort}(\mathbf{C}, \text{decreasing})$ | ▷ Sort sums in decreasing order |
| 4: $h_{\mathbf{C}} = \text{build chained model}(\mathcal{D}, \mathbf{C})$ | ▷ build a chained model for max correlation chain \mathbf{C} |
| 5: return $h_{\mathbf{C}}$ | |
-

3.3 Experimental Environment

Although many interesting applications of multi-target regression exist, there are not many publicly available datasets to use. The datasets used in the experimental study were collected from the Mulan website [85], as well as the UCI Machine Learning Repository [5]. Information on the 24 datasets used is summarized in Table 3.2, where the number of samples, attributes (dimensionality), and targets are shown.

Experiments were performed over the RC, ST, MTS, MTSC, ERC, ERCC, and MORF algorithms, which have also been used in the experimental study conducted in [79]. These algorithms were chosen because they have shown considerable performance in training multi-target models. They have also made their framework readily available for reproducing their results. All three SVR algorithms are implemented within the general framework of Mulan’s MTRRegressor¹ [85], which was built on top of Weka² [45]. LIBSVM’s Epsilon-SVR [24] implementation was used as the base SVR model. The parameters experimented with for

¹<http://mulan.sourceforge.net>

²<http://www.cs.waikato.ac.nz/ml/weka>

Table 3.2.: Multi-Target (MT) Regression datasets

Dataset	Samples (n)	Attributes (d)	Targets (m)
EDM	145	16	2
Enb	768	8	2
Jura	359	11	7
Osales	639	413	12
Scpf	1137	23	3
Slump	103	7	3
Solar Flare 1	323	10	3
Solar Flare 2	1,066	10	3
Water Quality	1,060	16	14
OES97	323	263	16
OES10	403	298	16
ATP1d	201	411	6
ATP7d	188	411	6
Andro	49	30	6
Wisconsin Cancer	198	34	2
Stock	950	10	3
California Housing	20,640	7	2
Puma8NH	8,192	8	3
Puma32H	8,192	32	6
Friedman	500	25	6
Polymer	41	10	4
M5SPEC	80	700	3
MP5SPEC	80	700	3
MP6SPEC	80	700	3

the SVR regression task are the penalty parameter C , the Gaussian kernel parameter γ , and the error or tube parameter ϵ given by Equations (3.3a) to (3.3c), referred to as (3.3).

$$C \in \{1, 10, 100\} \quad (3.3a)$$

$$\gamma \in \{1^{-9}, 1^{-7}, 1^{-5}, 1^{-3}, 1^{-1}, 1, 5, 10\} \quad (3.3b)$$

$$\epsilon \in \{0.01, 0.1, 0.2\} \quad (3.3c)$$

To ensure a controlled environment when conducting the performance comparisons, the experimental environment for running the competing algorithms was the same as what was done in [79]. This includes the following. The ST base-line model used was Bagging [21] of 100 regression trees [94]. The MTSC and ERCC methods are run using 10-fold cross-

validation, and the ensemble size for the ERC and ERCC methods was set to 10. The ensemble size of 100 trees was used for MORF, and the rest of its parameters were set as recommended by [62].

The performance metrics used to analyze our contributions' performances are shown in Equations 3.4 to 3.7. For unseen or test datasets of size \mathcal{N}_{test} , the performances are evaluated by taking the run time (seconds) each algorithm takes to build a classifier, as well as the following metrics, where the upwards arrow \uparrow indicates maximizing the metric and the downwards arrow \downarrow indicates minimizing the metric.

- The average correlation coefficient (aCC \uparrow):

$$\frac{1}{m} \sum_{j=1}^m \frac{\sum_{l=1}^{\mathcal{N}_{test}} (y_j^{(l)} - \bar{y}_j)(\hat{y}_j^{(l)} - \bar{\hat{y}}_j)}{\sqrt{\sum_{l=1}^{\mathcal{N}_{test}} (y_j^{(l)} - \bar{y}_j)^2 \sum_{l=1}^{\mathcal{N}_{test}} (\hat{y}_j^{(l)} - \bar{\hat{y}}_j)^2}} \quad (3.4)$$

- The mean squared error (MSE \downarrow):

$$\frac{1}{m} \sum_{j=1}^m \frac{1}{\mathcal{N}_{test}} \sum_{l=1}^{\mathcal{N}_{test}} (y_j^{(l)} - \hat{y}_j^{(l)})^2 \quad (3.5)$$

- The average root mean squared error (aRMSE \downarrow):

$$\frac{1}{m} \sum_{j=1}^m \sqrt{\frac{\sum_{l=1}^{\mathcal{N}_{test}} (y_j^{(l)} - \hat{y}_j^{(l)})^2}{\mathcal{N}_{test}}} \quad (3.6)$$

- The average relative root mean squared error (aRRMSE \downarrow):

$$\frac{1}{m} \sum_{j=1}^m \sqrt{\frac{\sum_{l=1}^{\mathcal{N}_{test}} (y_j^{(l)} - \hat{y}_j^{(l)})^2}{\sum_{l=1}^{\mathcal{N}_{test}} (y_j^{(l)} - \bar{y}_j)^2}} \quad (3.7)$$

The predicted output is represented by $\hat{\mathbf{y}}$, the average of the predicted output is $\bar{\hat{\mathbf{y}}}$, and the average of the true output target variable is $\bar{\mathbf{y}}$. The test dataset is the hold-out set during cross validation. This ensures our model is evaluated on data that it has not been trained on, and thus unbiased towards the training datasets. It also contributes to the generalizability and robustness of the model.

3.4 Results & Statistical Analysis

Tables 3.3, 3.5, 3.7, 3.9, and 3.11 show the results of our algorithm implementations compared with those of *RC*, *MORF*, *ST*, *MTS*, *MTSC*, *ERC*, and *ERCC*. Each subsection discusses a single metric along with the statistical analysis of the results. The best metric value obtained on each dataset is typeset in bold. Non-parametric statistical tests are then used to validate the experiments results obtained. To determine whether significant differences exist among the performance and results of the algorithms, the Iman-Davenport non-parametric test is run to rank the algorithms over the datasets used, according to the Friedman test. The average ranks are presented in the last row of the results tables. The Bonferroni-Dunn post-hoc test [36] is then used to find these differences that occur between the algorithms. Below each result table, a diagram highlighting the critical distance (in gray) between each algorithm is shown. The Wilcoxon, Nemenyi, and Holm [93] tests were run for each of the result metrics to compute multiple pairwise comparisons among the algorithms used in the experimental study. Tables 3.4, 3.6, 3.8, 3.10, and 3.12 show the sum of ranks R^+ and R^- of the Wilcoxon rank-sum test, and the p -values for the 3 tests, which show the statistical confidence rather than using a fixed α value.

3.4.1 Average Correlation Coefficient

Table 3.3 shows that our proposed methods perform the best on 15 out of the 24 datasets. Specifically, the maximum correlation chain method, SVRCC, performs the best on 11, which is better than the total number of datasets the competing methods performed better at (9). The Iman-Davenport statistic, distributed according to the F-distribution with 9 and 207 degrees of freedom is 6.72, with a p -value of $1.9E^{-8}$ which is significantly less than 0.01, implying a statistical confidence larger than 99%. Therefore, we can conclude that there exist statistically significant differences between the aCC results of the algorithms.

Figure 3.4 shows the mean rank values of each algorithm along with the critical difference value, 2.4236, for $\alpha = 0.05$. The algorithms that are to the right of the critical

Table 3.3.: Average Correlation Coefficient (aCC) for MT regressors

Datasets	MORF	ST	MTS	MTSC	RC	ERC	ERCC	SVR	SVRRC	SVRCC
Slump	0.6965	0.7062	0.7163	0.6977	0.6956	0.6977	0.7023	0.7245	0.7339	0.7457
Polymer	0.7305	0.7336	0.7371	0.7228	0.7015	0.7029	0.7222	0.7634	0.7857	0.7905
Andro	0.7349	0.6454	0.6793	0.6581	0.6915	0.6806	0.6653	0.6880	0.6951	0.7056
EDM	0.6722	0.6352	0.6412	0.6354	0.6355	0.6379	0.6354	0.6484	0.6565	0.6567
Solar Flare 1	0.1083	0.1258	0.1034	0.1193	0.1492	0.1387	0.1292	0.1066	0.0857	0.1152
Jura	0.7854	0.7907	0.7880	0.7882	0.7877	0.7884	0.7897	0.7789	0.7921	0.7983
Enb	0.9828	0.9832	0.9822	0.9829	0.9813	0.9823	0.9837	0.9858	0.9867	0.9868
Solar Flare 2	0.2357	0.2295	0.2375	0.2343	0.2302	0.2351	0.2432	0.1470	0.1648	0.1656
Wisconsin Cancer	0.3362	0.3587	0.3652	0.3588	0.3628	0.3609	0.3590	0.3187	0.3208	0.3373
California Housing	0.7705	0.7720	0.7149	0.7451	0.7007	0.7844	0.8065	0.7847	0.7949	0.8007
Stock	0.9785	0.9747	0.9755	0.9752	0.9753	0.9757	0.9763	0.9825	0.9829	0.9822
SCPF	0.5827	0.5508	0.5503	0.5477	0.5569	0.5656	0.5515	0.5891	0.5975	0.5946
Puma8NH	0.5424	0.4828	0.4942	0.4205	0.4677	0.4656	0.4650	0.6041	0.5975	0.6038
Friedman	0.1507	0.1609	0.1548	0.1667	0.1558	0.1608	0.1632	0.1710	0.1748	0.1752
Puma32H	0.3085	0.2934	0.2890	0.2504	0.2754	0.2870	0.2797	0.3358	0.3351	0.3385
Water Quality	0.4303	0.4063	0.4019	0.4051	0.3992	0.4052	0.4147	0.3545	0.3828	0.3857
M5SPEC	0.8161	0.8346	0.8134	0.8228	0.8333	0.8340	0.8308	0.9451	0.9452	0.9472
MP5SPEC	0.8315	0.8536	0.8244	0.8535	0.8524	0.8526	0.8542	0.9560	0.9602	0.9633
MP6SPEC	0.8317	0.8531	0.8231	0.8531	0.8507	0.8515	0.8541	0.9444	0.9500	0.9528
ATP7d	0.8260	0.8408	0.8422	0.8474	0.8273	0.8351	0.8464	0.8305	0.8407	0.8400
OES97	0.7829	0.7995	0.7990	0.8001	0.7986	0.7990	0.7999	0.8116	0.8134	0.8137
Osales	0.7186	0.6912	0.7104	0.7076	0.6357	0.7136	0.7193	0.6511	0.6433	0.6677
ATP1d	0.8961	0.9066	0.9051	0.9075	0.9048	0.9081	0.9071	0.9092	0.9130	0.9100
OES10	0.8708	0.8808	0.8805	0.8806	0.8804	0.8804	0.8809	0.8911	0.8924	0.8963
Average	0.6508	0.6462	0.6429	0.6409	0.6396	0.6476	0.6492	0.6634	0.6685	0.6739
Ranks	6.4167	5.8958	6.6042	6.4792	7.5208	5.8958	4.8542	4.7917	3.7083	2.8333

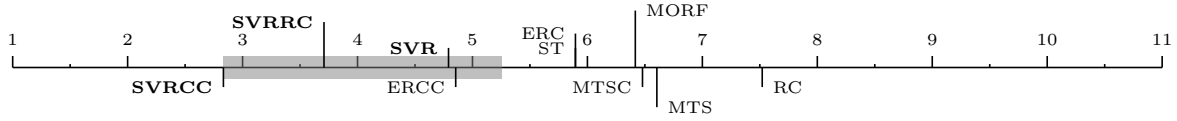


Fig. 3.4.: Bonferroni-Dunn test for aCC

Table 3.4.: Wilcoxon, Nemenyi, and Holm tests for aCC

SVRCC vs.	Wilcoxon R^+	Wilcoxon R^-	Wilcoxon p -value	Nemenyi p -value	Holm p -value
MORF	224.0	76.0	$3.4E^{-2}$	$4.1E^{-5}$	$8.3E^{-3}$
ST	239.0	61.0	$9.6E^{-3}$	$4.6E^{-4}$	$1.3E^{-2}$
MTS	242.0	58.0	$7.2E^{-3}$	$1.6E^{-5}$	$6.3E^{-3}$
MTSC	238.0	62.0	$1.1E^{-2}$	$3.0E^{-5}$	$7.1E^{-3}$
RC	250.0	50.0	$3.1E^{-3}$	0.0000	$5.6E^{-3}$
ERC	229.0	71.0	$2.3E^{-2}$	$4.6E^{-4}$	$1.0E^{-2}$
ERCC	221.0	79.0	$4.3E^{-2}$	$2.1E^{-2}$	$1.7E^{-2}$
SVR	297.0	3.00	$6.0E^{-7}$	$2.5E^{-2}$	$2.5E^{-2}$
SVRRC	266.5	33.5	$4.0E^{-4}$	$3.2E^{-1}$	$5.0E^{-2}$

difference rectangle are the ones with significantly different results. Therefore, the 6 out of 10 algorithms beyond the critical difference perform significantly worse than our control algorithm, SVRCC. Table 3.4 provides complementary analysis of the results. According

to the Wilcoxon test, SVRCC is shown to have significantly better performance over all algorithms with $p\text{-value} < 0.05$. The Nemenyi and Holm tests show that SVRCC performs significantly better than 6 out of the 9 algorithms with $p\text{-value} \leq 5.6E^{-3}$ and $\leq 1.7E^{-2}$, respectively. The exact confidence for algorithm SVRCC against all others is 0.95.

3.4.2 Mean Square Error

Table 3.5 shows that our proposed methods perform the best on 15 out of the 24 datasets. In this case, SVRCC also performs the best on 11 versus the 9 that the competing methods performed better at. The Iman-Davenport statistic, distributed according to the F-distribution with 9 and 207 degrees of freedom is 6.57, with a $p\text{-value}$ of $3.1E^{-8}$, implying statistically significant differences among the MSE results.

Figure 3.5 shows the mean rank values of each algorithm along with the critical difference value, 2.4236, for $\alpha = 0.05$. According to the critical difference bar, there are 6 out of 10 algorithms beyond that perform significantly worse than our control algorithm, SVRCC. According to the Wilcoxon test, shown in Table 3.6, SVRCC is shown to have significantly better performance over all algorithms with $p\text{-value} < 0.05$. The Nemenyi and Holm tests show that SVRCC performs significantly better than 6 out of the 9 algorithms with $p\text{-values} \leq 5.6E^{-3}$ and $\leq 1.7E^{-2}$ respectively, and has an exact confidence of 0.95 against all others.

3.4.3 Average Root Mean Square Error

Table 3.7 shows that our proposed methods perform the best on 18 out of the 24 datasets. In this case, SVRCC performs the best on 15 versus the 6 that the methods compared performed better at. The Iman-Davenport statistic is 7.6, with a $p\text{-value}$ of $1.3E^{-9}$, implying statistically significant differences in the aRMSE results.

Figure 3.6 shows the mean rank values of each algorithm along with the critical difference value, 2.4236, for $\alpha = 0.05$. According to the critical difference bar, there are 7 out of 10 algorithms that perform significantly worse than our control algorithm, SVRCC.

According to the Wilcoxon test, shown in Table 3.8, SVRCC is shown to have significantly better performance over all algorithms with p -value < 0.01 . The Nemenyi test shows that SVRCC performs significantly better than 7 out of the 9 algorithms with p -value

Table 3.5.: Mean Square Error (MSE) for MT regressors

Datasets	MORF	ST	MTS	MTSC	RC	ERC	ERCC	SVR	SVRRC	SVRCC
Slump	1.4388	1.4161	1.3667	1.4414	1.4602	1.4727	1.4183	1.2991	1.1726	1.1614
Polymer	1.6718	1.8120	1.5446	1.6726	1.8259	1.9999	1.6873	1.1874	1.1068	1.0796
Andro	1.4930	2.1467	1.4714	1.7525	2.2603	2.0812	1.8707	1.5406	1.2847	1.2187
EDM	0.8342	0.9373	0.9352	0.9418	0.9389	0.9326	0.9393	0.9092	0.8650	0.8817
Solar Flare 1	3.3458	3.1196	3.1193	3.0524	3.0357	3.0381	3.0594	2.9912	3.0176	3.0129
Jura	1.0973	1.0595	1.0732	1.0695	1.0744	1.0694	1.0632	1.1167	1.0435	1.0315
Enb	0.0381	0.0361	0.0407	0.0377	0.0452	0.0403	0.0343	0.0255	0.0216	0.0214
Solar Flare 2	2.9619	2.8532	2.7732	2.8282	2.8510	2.8273	2.8110	2.9518	2.9204	2.8713
Wisconsin Cancer	1.7666	1.7155	1.7156	1.7256	1.7119	1.7146	1.7195	1.8171	1.7915	1.7692
California Housing	0.8665	0.8221	0.9642	0.8673	1.0125	0.8952	0.7513	0.7477	0.6987	0.6726
Stock	0.0841	0.1039	0.0990	0.1008	0.0998	0.0987	0.0949	0.0578	0.0596	0.0554
SCPF	2.2244	2.3173	2.3661	2.3517	2.3923	2.3025	2.3295	2.2960	2.2510	2.3179
Puma8NH	1.9678	2.1133	2.0989	2.2024	2.1413	2.1473	2.1467	1.8242	1.8728	1.8299
Friedman	5.4573	5.3357	5.3478	5.3260	5.3482	5.3253	5.3210	5.3038	5.2942	5.2812
Puma32H	5.3419	4.9499	4.9627	5.0405	4.9905	4.9662	4.9805	5.2711	5.2749	5.1306
Water Quality	11.3143	11.5621	11.6276	11.5931	11.6495	11.6022	11.5004	12.2974	12.2042	12.0593
M5SPEC	1.0081	0.8754	1.0336	0.9421	0.8847	0.8824	0.8903	0.2578	0.2597	0.2575
MP5SPEC	1.1483	0.9817	1.1953	0.9970	0.9886	0.9880	0.9882	0.2261	0.1979	0.2136
MP6SPEC	1.1626	0.9928	1.1906	0.9992	1.0115	1.0045	0.9905	0.2926	0.2903	0.2954
ATP7d	1.7859	1.7348	1.6435	1.6460	1.8521	1.7888	1.6739	1.7820	1.7433	1.7098
OES97	4.6331	4.8340	4.8379	4.8082	4.8573	4.8591	4.8187	3.1440	3.0633	3.0499
Osales	7.3631	6.6850	5.8848	6.0850	7.8575	6.4746	5.9155	7.0727	7.3153	7.1374
ATP1d	1.0589	0.9056	0.9053	0.8982	0.9125	0.8783	0.9004	0.9091	0.8837	0.8922
OES10	3.6471	3.8931	3.8952	3.8909	3.9031	3.9063	3.8869	2.2623	2.1608	2.1320
Average	2.6546	2.6334	2.5872	2.5946	2.7127	2.6373	2.5747	2.3993	2.3664	2.3368
Ranks	6.5833	5.6667	6.0833	6.2500	7.8333	6.1250	5.1250	4.6667	3.6250	3.0417

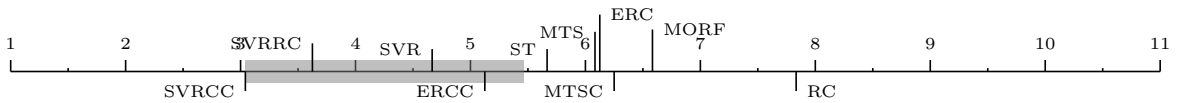


Fig. 3.5.: Bonferroni-Dunn test for MSE

Table 3.6.: Wilcoxon, Nemenyi, and Holm tests for MSE

SVRCC vs.	Wilcoxon R^+	Wilcoxon R^-	Wilcoxon p -value	Nemenyi p -value	Holm p -value
MORF	268.0	32.0	$3.2E^{-4}$	$5.1E^{-5}$	$6.3E^{-3}$
ST	241.0	59.0	$7.9E^{-3}$	$2.7E^{-3}$	$1.3E^{-2}$
MTS	224.0	76.0	$3.4E^{-2}$	$5.0E^{-4}$	$1.0E^{-2}$
MTSC	226.0	74.0	$2.9E^{-2}$	$2.4E^{-4}$	$7.1E^{-3}$
RC	263.0	37.0	$6.5E^{-4}$	0.0000	$5.6E^{-3}$
ERC	234.0	66.0	$1.5E^{-2}$	$4.2E^{-4}$	$8.3E^{-3}$
ERCC	224.0	76.0	$3.4E^{-2}$	$1.7E^{-2}$	$1.7E^{-2}$
SVR	262.0	38.0	$7.4E^{-4}$	$6.3E^{-2}$	$2.5E^{-2}$
SVRRC	245.0	55.0	$5.3E^{-3}$	$5.1E^{-1}$	$5.0E^{-2}$

Table 3.7.: Average Root Mean Square Error (aRMSE) for MT regressors

Datasets	MORF	ST	MTS	MTSC	RC	ERC	ERCC	SVR	SVRRC	SVRCC
Slump	0.6711	0.6652	0.6456	0.6699	0.6787	0.6793	0.6649	0.5561	0.5345	0.5337
Polymer	0.5277	0.5409	0.5042	0.5336	0.5536	0.5803	0.5319	0.4403	0.4062	0.4060
Andro	0.4649	0.5420	0.4414	0.4871	0.5390	0.5317	0.5039	0.4326	0.4061	0.3989
EDM	0.6372	0.6715	0.6705	0.6729	0.6722	0.6704	0.6721	0.6449	0.6411	0.6366
Solar Flare 1	0.9777	0.9274	0.9271	0.9089	0.8921	0.9016	0.9121	0.8856	0.8844	0.8801
Jura	0.5800	0.5686	0.5720	0.5706	0.5726	0.5712	0.5693	0.5794	0.5687	0.5622
Enb	0.1212	0.1166	0.1237	0.1214	0.1272	0.1253	0.1140	0.0981	0.0914	0.0903
Solar Flare 2	0.8725	0.8420	0.8127	0.8305	0.8313	0.8300	0.8304	0.8418	0.8349	0.8345
Wisconsin Cancer	0.9290	0.9163	0.9158	0.9187	0.9153	0.9160	0.9173	0.9422	0.9362	0.9306
California Housing	0.6541	0.6366	0.6889	0.6530	0.7053	0.6632	0.6079	0.6038	0.5859	0.5755
Stock	0.1643	0.1830	0.1774	0.1790	0.1790	0.1777	0.1739	0.1357	0.1329	0.1308
SCPF	0.7113	0.7235	0.7342	0.7255	0.7285	0.7143	0.7227	0.7155	0.7081	0.7048
Puma8NH	0.7855	0.8139	0.8114	0.8307	0.8196	0.8202	0.8203	0.7650	0.7740	0.7671
Friedman	0.9382	0.9203	0.9219	0.9199	0.9219	0.9197	0.9193	0.9203	0.9195	0.9183
Puma32H	0.9395	0.8700	0.8713	0.8778	0.8739	0.8716	0.8727	0.9353	0.9356	0.9331
Water Quality	0.8921	0.9015	0.9041	0.9025	0.9051	0.9030	0.8990	0.9284	0.9293	0.9271
M5SPEC	0.5707	0.5324	0.5761	0.5515	0.5347	0.5339	0.5376	0.2745	0.2744	0.2740
MP5SPEC	0.5315	0.4914	0.5426	0.4947	0.4930	0.4928	0.4928	0.2337	0.2176	0.2177
MP6SPEC	0.5344	0.4939	0.5416	0.4943	0.4982	0.4967	0.4927	0.2627	0.2460	0.2497
ATP7d	0.5216	0.4956	0.4752	0.4765	0.5194	0.5024	0.4824	0.5141	0.5066	0.5018
OES97	0.4652	0.4634	0.4635	0.4622	0.4643	0.4644	0.4627	0.3794	0.3768	0.3749
Osales	0.7190	0.6912	0.6496	0.6615	0.7591	0.6772	0.6515	0.7212	0.7343	0.7121
ATP1d	0.4053	0.3608	0.3587	0.3591	0.3653	0.3562	0.3596	0.3693	0.3638	0.3507
OES10	0.3954	0.3896	0.3897	0.3892	0.3901	0.3903	0.3889	0.3085	0.3039	0.3038
Average	0.6254	0.6149	0.6133	0.6121	0.6225	0.6162	0.6083	0.5620	0.5547	0.5506
Ranks	7.3333	5.7708	5.8125	6.0625	7.6250	6.0208	4.8542	5.0625	3.9167	2.5417



Fig. 3.6.: Bonferroni-Dunn test for aRMSE

Table 3.8.: Wilcoxon, Nemenyi, and Holm tests for aRMSE

SVRCC vs.	Wilcoxon R^+	Wilcoxon R^-	Wilcoxon p -value	Nemenyi p -value	Holm p -value
MORF	286.0	14.0	$1.3E^{-5}$	0.0000	$6.3E^{-3}$
ST	259.0	41.0	$1.1E^{-3}$	$2.2E^{-4}$	$1.3E^{-2}$
MTS	247.0	53.0	$4.3E^{-3}$	$1.8E^{-5}$	$1.0E^{-2}$
MTSC	251.0	49.0	$2.8E^{-3}$	$5.6E^{-5}$	$7.1E^{-3}$
RC	270.0	30.0	$2.4E^{-4}$	0.0000	$5.6E^{-3}$
ERC	255.0	45.0	$1.8E^{-3}$	$6.9E^{-5}$	$8.3E^{-3}$
ERCC	246.0	54.0	$4.8E^{-3}$	$8.2E^{-3}$	$2.5E^{-2}$
SVR	296.0	4.00	$8.3E^{-7}$	$3.9E^{-3}$	$1.7E^{-2}$
SVRRC	284.0	16.0	$2.0E^{-5}$	$1.2E^{-1}$	$5.0E^{-2}$

$\leq 5.6E^{-3}$, while the stricter Holm test shows that it performs significantly better than 8 out of the 9 algorithms with p -value ≤ 0.05 .

3.4.4 Average Relative Root Mean Square Error

Table 3.9 shows that our proposed methods perform the best on 16 out of the 24 datasets. In this case, SVRCC performs the best on 11 versus the 6 that the competing methods

Table 3.9.: Average Relative Root Mean Square Error (aRRMSE) for MT regressors

Datasets	MORF	ST	MTS	MTSC	RC	ERC	ERCC	SVR	SVRRC	SVRCC
Slump	0.6939	0.6886	0.6690	0.6938	0.7019	0.7022	0.6886	0.5765	0.5545	0.5560
Polymer	0.6159	0.5971	0.5778	0.6493	0.6270	0.6544	0.6131	0.5573	0.5253	0.5116
Andro	0.5097	0.5979	0.5155	0.5633	0.5924	0.5885	0.5666	0.4856	0.4651	0.4455
EDM	0.7337	0.7442	0.7413	0.7446	0.7449	0.7452	0.7443	0.7058	0.7070	0.6978
Solar Flare 1	1.3046	1.1357	1.1168	1.0758	0.9951	1.0457	1.0887	0.9917	0.9455	0.9320
Jura	0.5969	0.5874	0.5906	0.5892	0.5910	0.5896	0.5880	0.5952	0.5764	0.5885
Enb	0.1210	0.1165	0.1231	0.1211	0.1268	0.1250	0.1139	0.0977	0.0910	0.0899
Solar Flare 2	1.4167	1.1503	0.9483	1.0840	1.0092	1.0522	1.0928	1.0385	1.0253	1.0298
Wisconsin Cancer	0.9413	0.9314	0.9308	0.9336	0.9305	0.9313	0.9323	0.9555	0.9483	0.9427
California Housing	0.6611	0.6447	0.6974	0.6630	0.7131	0.6690	0.6146	0.6130	0.5945	0.5852
Stock	0.1653	0.1844	0.1787	0.1803	0.1802	0.1789	0.1752	0.1364	0.1337	0.1388
SCPF	0.8273	0.8348	0.8436	0.8308	0.8263	0.8105	0.8290	0.8164	0.8037	0.8013
Puma8NH	0.7858	0.8142	0.8118	0.8311	0.8199	0.8205	0.8207	0.7655	0.7744	0.7676
Friedman	0.9394	0.9214	0.9231	0.9210	0.9231	0.9209	0.9204	0.9218	0.9208	0.9196
Puma32H	0.9406	0.8713	0.8727	0.8791	0.8752	0.8729	0.8740	0.9364	0.9367	0.9319
Water Quality	0.8994	0.9085	0.9109	0.9093	0.9121	0.9097	0.9057	0.9343	0.9310	0.9045
M5SPEC	0.5910	0.5523	0.5974	0.5671	0.5552	0.5542	0.5558	0.2951	0.2935	0.2925
MP5SPEC	0.5522	0.5120	0.5683	0.5133	0.5145	0.5143	0.5119	0.2484	0.2323	0.2358
MP6SPEC	0.5553	0.5152	0.5686	0.5119	0.5198	0.5187	0.5109	0.2850	0.2669	0.2623
ATP7d	0.5563	0.5308	0.5141	0.5142	0.5558	0.5397	0.5182	0.5455	0.5371	0.5342
OES97	0.5490	0.5230	0.5229	0.5217	0.5239	0.5237	0.5222	0.4641	0.4618	0.4635
Osales	0.7596	0.7471	0.7086	0.7268	0.8318	0.7258	0.7101	0.7924	0.7924	0.7811
ATP1d	0.4173	0.3732	0.3733	0.3712	0.3790	0.3696	0.3721	0.3773	0.3707	0.3775
OES10	0.4518	0.4174	0.4176	0.4171	0.4178	0.4180	0.4166	0.3570	0.3555	0.3538
Average	0.6910	0.6625	0.6551	0.6589	0.6611	0.6575	0.6536	0.6039	0.5935	0.5893
Ranks	7.5000	5.7708	5.9375	6.1667	7.4375	6.3750	4.9792	4.7708	3.2708	2.7917

Fig. 3.7.: Bonferroni-Dunn test for aRRMSE

Table 3.10.: Wilcoxon, Nemenyi, and Holm tests for aRRMSE

SVRCC vs.	Wilcoxon R^+	Wilcoxon R^-	Wilcoxon p -value	Nemenyi p -value	Holm p -value
MORF	290.0	10.0	$5.1E^{-6}$	0.0000	$5.6E^{-3}$
ST	261.0	39.0	$8.5E^{-4}$	$6.5E^{-4}$	$1.3E^{-2}$
MTS	239.0	61.0	$9.6E^{-3}$	$3.2E^{-3}$	$1.0E^{-2}$
MTSC	261.0	39.0	$8.5E^{-4}$	$1.1E^{-3}$	$8.3E^{-3}$
RC	275.0	25.0	$1.1E^{-4}$	0.0000	$6.3E^{-3}$
ERC	261.0	39.0	$8.5E^{-4}$	$4.1E^{-5}$	$7.1E^{-3}$
ERCC	254.0	46.0	$2.0E^{-3}$	$1.2E^{-2}$	$1.7E^{-2}$
SVR	291.0	9.00	$3.9E^{-6}$	$2.4E^{-2}$	$2.5E^{-2}$
SVRRC	222.5	77.5	$3.8E^{-2}$	$5.8E^{-1}$	$5.0E^{-2}$

performed better at. The Iman-Davenport statistic is 8.54, with a p -value of $7.6E^{-11}$.

Figure 3.7 shows the mean rank values of each algorithm along with the critical difference value, 2.4236, for $\alpha = 0.05$. According to the critical difference bar, there are 6 out of 10 algorithms beyond that perform significantly worse than our control algorithm, SVRCC.

According to the Wilcoxon test, shown in Table 3.10, SVRCC is shown to have significantly better performance over all algorithms with p -value < 0.05 , and 8 out of the 9 algorithms for p -value < 0.01 . The Nemenyi test shows that SVRCC performs significantly better than 6 out of the 9 algorithms with p -value $\leq 5.6E^{-3}$, and the Holm test shows its performance is significantly better than 8 out of the 9 algorithms with p -value ≤ 0.05 .

3.4.5 Run Time

Table 3.11 shows that our proposed methods perform faster on 16 out of the 24 datasets. In this case, SVR performs the best on 12 versus the 6 of the state-of-the-art methods. The Iman-Davenport statistic 64.41, with a p -value of 0.0 which implies a statistical confidence of 100%. Figure 3.8 shows the mean rank values of each algorithm along with the critical difference value, 2.4236, for $\alpha = 0.05$. According to the critical difference bar, there are 6 out of 10 algorithms beyond that perform significantly worse than our control algorithm, SVR. According to the Wilcoxon test, shown in Table 3.12, SVR is shown to have significantly better performance over all algorithms with p -value < 0.01 . The Nemenyi and Holm tests show that SVRCC performs significantly better than 6 out of the 9 algorithms and 8 out of the 9 algorithms with p -value $\leq 5.6E^{-3}$ and p -value $\leq 1.6E^{-2}$, respectively.

3.4.6 Discussion

Results indicate that our proposed methods perform competitively against the current contemporary methods, specifically SVRCC which exploits relationships among the targets. Firstly, they show that using SVR as a base-line method for multi-target chaining causes a performance improvement in model prediction, compared to other ST base-line models,

Table 3.11.: Run Time (seconds) for MT regressors

Datasets	MORF	ST	MTS	MTSC	RC	ERC	ERCC	SVR	SVRRC	SVRCC
Slump	38.1	2.6	9.9	15.9	1.8	11.1	50.5	0.6	1.9	0.7
Polymer	7.6	2.7	9.1	15.5	1.9	14.9	80.5	0.5	2.6	0.5
Andro	25.7	4.4	15.0	34.2	3.4	33.2	197.9	1.1	6.2	1.1
EDM	24.8	2.8	9.4	18.1	2.1	5.8	19.0	0.9	1.0	0.9
Solar Flare 1	34.1	3.5	13.6	26.7	2.7	17.7	86.9	2.3	9.3	2.6
Jura	64.3	7.9	31.8	74.3	6.4	43.5	254.2	4.7	18.7	5.3
Enb	71.4	6.6	26.1	63.6	5.4	15.6	69.6	11.3	17.7	15.9
Solar Flare 2	55.4	7.4	30.7	68.0	6.3	42.9	241.5	9.4	53.5	15.6
Wisconsin Cancer	51.4	6.1	21.9	53.7	4.9	14.8	61.6	2.0	2.4	2.0
California Housing	93.0	9.7	34.8	75.9	8.2	21.3	102.0	15.8	25.2	23.6
Stock	93.7	11.7	46.8	96.7	11.0	75.4	427.3	18.5	90.5	26.3
SCPF	66.3	19.3	65.9	176.3	15.0	104.2	734.2	32.8	162.8	48.8
Puma8NH	130.4	29.7	106.7	288.6	27.9	201.6	1227.7	94.1	516.6	177.1
Friedman	79.5	27.0	81.2	258.3	25.0	273.7	2871.6	12.3	322.3	18.8
Puma32H	93.9	68.1	181.0	635.0	87.7	667.9	6087.0	32.2	1018.7	53.1
Water Quality	108.4	93.1	262.1	912.3	127.2	925.4	10993.3	110.2	2567.9	189.5
M5SPEC	89.8	68.9	166.3	604.6	73.7	262.3	3132.1	39.2	546.7	45.1
MP5SPEC	84.5	94.6	221.2	888.3	91.5	557.0	6864.1	49.3	1132.1	58.4
MP6SPEC	90.3	93.4	212.6	871.0	89.1	557.6	6761.3	47.2	1227.1	58.5
ATP7d	70.5	262.6	452.1	2319.8	242.1	1779.2	24373.8	80.0	1897.4	136.5
OES97	83.4	485.3	1146.6	4928.9	499.8	5315.0	58072.1	148.2	3759.1	342.6
Osales	92.0	1094.8	2340.7	8322.2	986.5	11361.2	122265.3	437.0	4830.1	843.6
ATP1d	70.7	272.9	476.5	2568.9	261.9	2138.9	26768.9	95.0	2127.8	174.4
OES10	90.0	738.9	1633.6	6682.9	688.5	7150.8	83533.1	229.1	5419.4	577.1
Average	71.2	142.2	316.5	1250.0	136.2	1316.3	14803.2	61.4	1073.2	117.4
Ranks	5.5	3.71	6.0	8.29	3.0	7.08	9.92	1.88	6.71	2.92

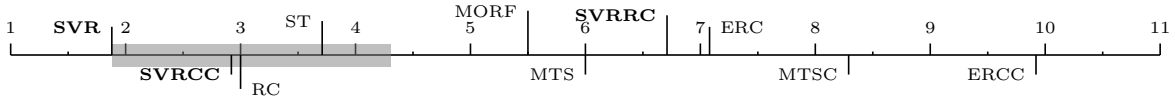


Fig. 3.8.: Bonferroni-Dunn test for Run Time

Table 3.12.: Wilcoxon, Nemenyi, and Holm tests for Run Time

SVRCC vs.	Wilcoxon R^+	Wilcoxon R^-	Wilcoxon p -value	Nemenyi p -value	Holm p -value
SVRCC	295.0	5.00	$1.2E^{-6}$	$2.3E^{-1}$	$5.0E^{-2}$
MORF	225.0	75.0	$3.2E^{-2}$	$3.4E^{-5}$	$1.3E^{-2}$
ST	221.5	78.5	$4.1E^{-2}$	$3.6E^{-2}$	$1.7E^{-2}$
MTS	300.0	0.00	$1.2E^{-7}$	$2.0E^{-6}$	$1.0E^{-2}$
MTSC	300.0	0.00	$1.2E^{-7}$	0.0000	$6.3E^{-3}$
RC	229.0	71.0	$2.3E^{-2}$	$2.0E^{-1}$	$2.5E^{-2}$
ERC	300.0	0.00	$1.2E^{-7}$	0.0000	$7.1E^{-3}$
ERCC	300.0	0.00	$1.2E^{-7}$	0.0000	$5.6E^{-3}$
SVRRC	300.0	0.00	$1.2E^{-7}$	0.0000	$8.3E^{-3}$

as well as most MT methods. This demonstrates the advantages of using the SVR method as a base-line for multi-target learning, thus increasing the performance of the ensemble of

regressor chains, SVRRC, compared to ERCC. More importantly, the results highlight the major advantage of capturing and exploiting the targets’ relationships during model training. Using an ensemble of randomly generated chains does not ensure the targets’ correlations are fully captured; however, using a maximum correlation chain improves the performance in terms of quality metrics as well as run time. The run time of SVR was shown to be the fastest, due to the fact that its complexity is mostly dependent on the number of targets. However, this method does not consider any of the correlations that might exist among the target variables, but SVRCC does take them into account and does not have a significant impact on run time. The most noteworthy finding that highlights advantage of using the base-line SVR and the maximum correlation method, SVRCC, rather than random chaining as done in ERCC, are the run time results and their analysis. ERCC had the worst run time across all datasets, whereas our proposals, SVR and SVRCC, performed the fastest. This emphasizes the advantage of using a single chain rather an ensemble of random chains, especially when the single chain is ordered in the direction of the targets maximum correlation.

3.5 Conclusions

This contribution proposed three novel methods for solving multi-target regression problems. The first method takes a *problem transformation* approach, which generates m ST models, each trained independently. This base-line approach was shown to perform the best in terms of run time, but its drawback is that it does not take the possible correlations between the target variables into account during training. The second implements SVR as an ensemble model of randomly generated chains, inspired by the classification method ERCC. This was done to investigate the effects of exploiting correlations among the target variables during model training. Due to the random nature of this method, capturing target correlations is not guaranteed. The third proposal, SVRCC, generates a single chain that is ordered in the direction of the targets’ maximum correlation, ensuring the correlations among targets are taken into account within the learning process.

The experimental study compared the proposed methods' performances to 7 popular, contemporary methods on 24 MT regression datasets. Firstly, the results show the superior performance of using the SVR method as a base-line model, rather than regression trees as used in MORF. The results for SVRRC show an increase in performance when random chaining is used to develop an ensemble model. This indicates the importance of the relationship among the target variables during training. Finally, the results show the superiority of using the SVRCC method, which was ranked the best in all quality metrics and second best in terms of run time. SVRCC performed better than the single-target SVR model and the randomly chained ensemble model SVRRC, showing that the targets' maximum correlation does positively contribute toward model training. The statistical analysis supports and shows the significance of the results obtained by our experiments. They demonstrated that statistically significant differences exist between the proposed algorithms against the methods compared. SVRCCs competitive performance, as well as speed, shows that it is a powerful learning algorithm for multi-target problems. The research outcomes of this chapter have been published in [68].

CHAPTER 4

MULTI-INSTANCE SVM USING BAG-REPRESENTATIVES

This contribution of this chapter proposes a novel support vector machine (SVM) formulation under the multiple-instance learning (MIL) paradigm. It also presents a novel algorithm and bag representative selector that trains the SVM using bag-level information, named Multi-Instance Representative SVM (MIRSVM). The contribution is able to identify instances that highly impact classification, i.e. the *bag-representatives*, for both positive and negative bags, while finding the optimal class separation hyperplane. Unlike other multi-instance SVM methods, this approach eliminates possible class imbalance issues by allowing both positive and negative bags to have at most one representative, which constitute as the most contributing instances to the model. The experimental study evaluates and compares the performance of this proposal against 11 popular and widely used multi-instance methods over 15 datasets, and the results are validated through non-parametric statistical analysis. The results indicate that bag-based learners outperform the instance-based and wrapper methods, and emphasize this proposal’s overall superior performance against other multi-instance SVM models.

4.1 Multi-Instance Classification Background

This section defines the notation that will be used throughout this chapter and reviews related works pertaining to multiple-instance learning, specifically the concepts of instance-based and bag-based learners are discussed and compared, along with algorithms within those paradigms.

Table 4.1.: Summary of Multiple-Instance Learning Notation

Definition	Notation
Number of Bags	n
Number of Instances	m
Number of Input Attributes	d
Set of Bags	$\mathcal{B} = \{\mathcal{B}_1, \dots, \mathcal{B}_n\}$
Bag Index Set	$I \in \mathbb{Z}_+^n$
Input Space	$\mathbf{X} \in \mathbb{R}^{m \times d}$
Bag Labels	$\mathbf{Y} \in \{-1, 1\}^n$
Input Instance i from Bag I	$\mathbf{x}_i \in \mathbf{x}_I = (x_{i1}, \dots, x_{id}), \forall i \in \{1, \dots, n\}, i \in I$
Unknown Individual Instance Label i	$y_i \in \{-1, 1\}$
Bag I	$\mathcal{B}_I = \{\mathbf{x}_i \mid \forall i \in I\}$
Full Multi-Instance Training Dataset	$\mathcal{D} = \{(\mathcal{B}_1, Y_1), \dots, (\mathcal{B}_n, Y_n)\}$

4.1.1 Notation

Let \mathcal{D} be a training dataset of n bags. Let $\mathbf{Y} \in \mathcal{D}$ be a vector of n labels corresponding to each bag, having a domain of $\mathbf{Y} \in \{-1, 1\}^n$. Let $\mathbf{X} \in \mathcal{D}$ be a matrix consisting of d input variables and m instances, $\mathbf{x}_i \in \mathbf{X}, \forall i \in \{1, \dots, m\}$, having a domain of $\mathbf{X} \in \mathbb{R}^{m \times d}$. Let \mathcal{B} be the set of bags which contain $|\mathcal{B}_I|$ number of instances, sometimes of different size and usually non-overlapping, such that $\mathcal{B}_I = \{\mathbf{x}_1, \dots, \mathbf{x}_{|\mathcal{B}_I|}\}$ for index set $I \in \{1, \dots, n\}$.

4.1.2 Multi-Instance Classification

The difference between MIL and traditional learning lies in the nature of the data. In the traditional binary classification setting, the goal is to learn a model that maps input samples to labels, $f : \mathbb{R}^{m \times d} \rightarrow Y^m \in \{-1, +1\}$. In the multi-instance setting, samples are called *bags* and each bag contains one or more input instances and is assigned a single bag-level label. Input instances, $\{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_m\}$, are grouped into bags with unique identifiers, $\mathcal{B} = \{\mathcal{B}_1, \mathcal{B}_2, \dots, \mathcal{B}_n \mid \mathcal{B}_I = \{\mathbf{x}_i \mid \forall i \in I\}, \forall I \in \{1, \dots, n\}\}$ and assigned a label, Y_I . The individual instance labels within a bag are unknown. Traditionally, MIL has focused on binary classification problems, however, there are cases where the number of classes can be larger, i.e. $\mathbf{Y} \in \mathbb{Z}^n$. The scope of this chapter will consist of binary multiple instance

classification problems.

Using the training dataset $\mathcal{D} = \{(\mathcal{B}_1, Y_1), \dots, (\mathcal{B}_n, Y_n)\}$, the goal is to train a classifier that predicts the label of an unseen bag, $f(\mathcal{B}_{n+1}) \rightarrow Y_{n+1}$ [3]. In order to build a classifier without any knowledge of the individual training instance labels, Dietterich et. al. [32] proposed the *standard MI* (SMI) hypothesis, shown in Equation (4.1), which states that a bag is labeled positive if and only if at least one of the instances in the bag is positive, and is labeled negative otherwise.

$$Y_I = \begin{cases} +1 & \text{if } \exists y_i = +1, \forall i \in I \\ -1 & \text{otherwise.} \end{cases} \quad (4.1)$$

This implies that individual instance labels y_i exist, but are not known (for positive bags) during training. Equation (4.1) can also be rewritten as Equation (4.2) for simplicity:

$$Y_I = \operatorname{argmax}_{\forall i \in I} y_i. \quad (4.2)$$

In addition to the SMI assumption, alternative MI assumptions have been proposed to date [40]. A recent review [3] describing the taxonomy of multi-instance classification presents various methods and algorithms used in literature, which are categorized based on their approach to handling the MI input space. Instance-based classifiers that fall under the *instance-space paradigm*, aim to separate instances in positive bags from those in negative ones. Bag-level classifiers (*bag-space paradigm*) treat each bag as a whole entity, implicitly extracting information from each bag in order to accurately predict their labels. Methods that fall under the *embedded-space paradigm* map the input bags to a single feature vector that explicitly encapsulates all the relevant information contained within each bag.

Instance-based methods that follow the SMI assumption attempt to identify desirable instance properties that make a bag positive. A simple and natural way of addressing this type of learning problem is to assume that each instance in a bag has the same label as the bag itself. After that, a single-instance classifier can be trained on the transformed dataset,

and finally, the SMI assumption can be applied over the predicted instance labels of unseen bags [48]. Methods that employ this type of approach are called *Wrapper* methods; they act as an interface between the instance and bag levels. One algorithm that employs this approach is called Simple-MI [33]. Simple-MI represents each bag with the mean vector of the instances within it. Another example is MIWrapper [41], which introduces weights to treat instances from different bags differently. The major disadvantage of wrapper techniques is that they assume the distribution the instances in positive bags is positive, when it may not be, thus imposing noise over the positive class.

One traditional instance-based method that takes a different approach is the Axis-Parallel Rectangle (APR) [32], which trains a model that assigns a positive label to an instance if it belongs to an axis-parallel rectangle in feature space, and assigns a negative label otherwise. The APR method is optimized by maximizing the number of positive bags in the training set containing at least one instance in the APR, while concurrently maximizing the number of negative bags that do not contain any instance in the APR.

Another similar and popular approach that falls under maximum likelihood-based methods, is the Diverse Density (DD) [64] framework. The DD metric is maximized for instances in feature space that are near at least one instance in a positive bag and far from all instances in negative bags. In the Expectation-Maximization Diverse Density (EM-DD) algorithm, Zhang and Goldman [99] propose a similar framework that iteratively maximizes the DD measure. Auer and Ortner [6] present a boosting approach that uses balls centered around positive bags to solve the MI problem called Multi-Instance Optimal Ball (MIOptimalBall). This approach is similar to that of APR and DD, except that Auer and Ortner [6] propose computing optimal balls per positive bags. A major challenge affecting these methods is that the distributions of the positive and negative bags affect their performance. Methods based on the DD metric [22, 26, 27] assume the positive instances form a cluster, which may not be the case. Alternatively, Fu et al.[43] models the distribution of negative bags with Gaussian kernels, which can prove difficult when the quantity of data is limited.

As mentioned previously in brief, the classical method of boosting [42, 73] has been adapted to a multi-instance instance-based algorithm. For example, Viola et al. [97] proposed an adaptation of boosting to the multi-instance paradigm based on the standard MI assumption, named MILBoost. Later, inspired by the MILBoost [97] and Online-ADABOOST [69] algorithms, Babenko et al. [7] developed a novel online MI boosting method which lead to a robust and stable model in the area of object tracking. All the works mentioned, including many other such as citeqi2011online,xie2012online, have positively contributed and inspired many approaches in the area of visual detection and tracking.

An extension of traditional single-instance k -nearest neighbors method (k -NN) was proposed by Wang and Zucker [90] to be applied to the bag-level, named CitationKNN. This method uses a distance function between bags in order to determine bag similarities. Not only are the set of closest bags to a single bag considered, but also how many bags is the single bag closest to. A voting scheme is then used to determine the bag class labels.

Blockeel et al. [13] introduced the Multi-Instance Tree Inducer (MITI), based on the standard MI assumption, which uses decision trees as a heuristic to solve the MI problem. This approach aims to identify whether an instance within a bag is truly positive and eliminate false positives within the same bag. The disadvantage of this approach stems from removing instances considered as false positives from partially grown trees without updating the existing tree structure. Bjerring and Frank [11] then enhanced this approach by creating the method Multi-Instance Rule Induction (MIRI). The algorithm aims to eliminate any possibility of a suboptimal split because the tree is discarded and regrown.

The MI adaptation of the SVM presents two contexts for solving the problem: the instance-level and the bag-level. The first tries to identify instances, either all within a bag or just key instances, that help find the optimal separating hyperplane between positive and negative bags. The latter uses kernels defined over whole bags to optimize the margin [34].

Andrews et. al. [4] proposed a mixed-integer quadratic program that solves the MI problem at an instance-level, using a support vector machine, named MISVM, that can be

solved heuristically. Rather than maximizing the margin of separability between instances of different classes, this instance-based method maximizes the margin between bags. It tries to identify the key instance from each positive bag that makes the bag positive by assuming it has the highest margin value. Instances from positive bags are selected as bag-representatives, and the algorithm iteratively creates a classifier that separates those representatives from all instances from the negative bags. Using bag-representatives from one class and all instances from the other is an example of an approach that combines rules from the SMI assumption and the collective assumption. A disadvantage of this approach stems from the assumption that all instances within positive bags are also positive, which is an implicit step in the initialization of MISVM. Andrews et. al. [4] also proposed a second mixed-integer instance-level approach, named mi-SVM, which does not discard the negative instances of the positive bags. It rather tries to identify the instances within positive bags that are negative and utilize them in the construction of the negative margin. The main disadvantage of these approaches is that they create an imbalanced class problem that favors the negative class, resulting in a biased classifier.

Asymmetric SVM (ASVM), presented by Yang et al. [47], was designed to use an asymmetric loss function under the SMI assumption. This approach was based on the idea that the cost of misclassification is different for positive and negative bags. For example, a false negative instance in a positive bag would not necessarily introduce an error on the bag label, assuming there are many positive instances within the bag. However, a false positive instance in a negative bag would definitely lead to an error. Using rules such as these, ASVM aims to minimize false positives, while ensuring all negative instances are on the negative side of the hyperplane.

The approach presented by Cheung et al. [28] presented a loss function that takes the cost associated with bag labels and the cost (under the SMI assumption) between prediction of beach bag and its instances into account. They also presented an SVM regularization scheme as well which, rather than using a heuristic method, used concave-convex optimization,

ensuring local-optimum convergence.

An example of using the approach of a bag-level kernel would coincidentally be one of the first bag-level approaches to the multi-instance SVM problem, proposed by Gärtner et. al. [44]. A bag-level kernel determines the similarity between two bags in a higher dimensional space. Blaschko et. al. [12] proposed conformal kernels which manipulate each attribute’s dimension based on its importance, without affecting the angles between vectors in the transformed space. These type of bag level kernels transform the bags into a single-instance representation which enables standard SVMs to be directly applied to multi-instance data.

Unlike the bag-based methods mentioned previously, which have a the SMI assumption embedded in their design, mapping-based algorithms do not assume a specific relationship exists between the labels of each bag and its instances. Rather, the relationship is learned from the data. An example of such methods is the Two-Level Classifier (TLC) [92]. Another example, which includes the use of kernels in the multi-instance bag-space setting, would be approach taken by Zhou et al. [101]. The basic idea behind their method is to treat instances in an non-i.i.d. manner, thus exploiting relationships among instances using a graph kernel.

Wang and Zucker [91] proposed CitationKNN which extends the traditional k -nearest neighbors (k -NN) method to the bag-level. To classify a new bag \mathcal{B} , CitationKNN uses a distance function that operates at the bag level to determine which bags are the closest to \mathcal{B} . It not only considers the bags that are closest, but also the bags for which \mathcal{B} is in the set of closest bags to others. Any bag-based distance function can be used by CitationKNN.

For most of the methods described above, implicit or explicit assumptions have been made about the distribution of the data. Selecting a method that is robust for a problem such as MIL can be difficult when little is known about the nature of the data, especially considering the unknown distribution of the instances within bags [3]. The proposed method, MIRSVM, is a general method that uses support vector machines to design a MIL model without making prior assumptions about the data. Classifiers of this type are known to provide better generalization capabilities and performance, as well as sparser models.

4.2 MIRSVM: A Novel SVM for Multi-Instance Classification

MIRSVM is based on the idea of selecting representative instances from both positive and negative bags which are used to find an unbiased, optimal separating hyperplane. A representative is iteratively chosen from each bag, and a new hyperplane is formed according to the representatives until they converge. Based on the SMI hypothesis, only one instance in a bag is required to be positive for the bag to adopt a positive label. Due to the unknown distribution of instances within positive bags, MIRSVM is designed to give preference to negative bags during training, because their distribution is known, i.e. all instances are guaranteed to be negative. This is evident during the representative selection process, by taking the index of the maximum output value within each bag based on the current hyperplane using the following rule, $s_I = \operatorname{argmax}_{i \in I} (\langle \mathbf{w}, \mathbf{x}_i \rangle + b)$, $\forall I \in \{1, \dots, n\}$. In other words, the most positive instance is chosen from each positive bag and the least negative instance is chosen from each negative bag (instances with the largest output value based on the current hyperplane), pushing the decision boundary towards the positive bags.

Equation (4.3) presents the primal MIRSVM optimization problem:

$$\min_{\mathbf{w}, b, \xi} \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_I \xi_I, \quad (4.3)$$

$$\text{s.t. } Y_I (\langle \mathbf{w}, \mathbf{x}_{s_I} \rangle + b) \geq 1 - \xi_I, \forall I \in \{1, \dots, n\} \quad (4.3a)$$

$$\xi_I \geq 0, \forall I \in \{1, \dots, n\}, \quad (4.3b)$$

where \mathbf{S}_I is the set of the bag representatives' indices and \mathbf{x}_{s_I} is the instance representative of bag \mathcal{B}_I . Note the variables in MIRSVMs formulation are the similar to those of the traditional SVM, except they are now representing each bag as an instance. Solving the optimization problem given in Equation (4.3) using a quadratic programming solver is a computationally expensive task due to the number of constraints, which scales by the number of bags n , as well as the calculation of the inner product between two d -dimensional vectors in constraint (4.3a). The proposed solution for these problems was deriving and solving the

dual of the optimization problem given by Equation (4.3).

The dual can be formed by taking the Lagrangian of (4.3), given by Equation (4.4), where α and β are the non-negative Lagrange multipliers.

$$\mathcal{L}(\mathbf{w}, b, \xi, \alpha, \beta) = \frac{1}{2} \sum_{j=1}^d w_j^2 + C \sum_I \xi_I - \sum_I \beta_I \xi_I - \sum_I \alpha_I \left(Y_I \left(\sum_{j=1}^d w_j x_{s_I j} + b \right) - 1 + \xi_I \right) \quad (4.4)$$

At optimality [20], $\nabla_{\mathbf{w}, b, \xi} \mathcal{L}(\mathbf{w}, b, \xi, \alpha, \beta) = 0$ and the following conditions are met:

$$\frac{\partial \mathcal{L}}{\partial w_j} : w_j = \sum_I \alpha_I Y_I x_{s_I j}, \forall j \in \{1, \dots, d\} \quad (4.5)$$

$$\frac{\partial \mathcal{L}}{\partial b} : \sum_I \alpha_I Y_I = 0, \quad (4.6)$$

$$\frac{\partial \mathcal{L}}{\partial \xi_I} : \alpha_I + \beta_I = C, \forall I \in \{1, \dots, n\} \quad (4.7)$$

By substituting Equations (4.5), (4.6), and (4.7) into the Lagrangian in (4.4), the dual MIRSVM formulation becomes the following:

$$\max_{\alpha, \beta} \sum_I \alpha_I - \frac{1}{2} \sum_I \sum_{K \in I} \sum_{j=1}^d \alpha_I \alpha_K Y_I Y_K x_{s_I j} x_{s_K j} \quad (4.8)$$

$$\text{s.t.} \sum_I \alpha_I Y_I = 0 \quad (4.8a)$$

$$\alpha_I + \beta_I = C, \forall I \in \{1, \dots, n\} \quad (4.8b)$$

$$\alpha_I \geq 0, \forall I \in \{1, \dots, n\} \quad (4.8c)$$

$$\beta_I \geq 0, \forall I \in \{1, \dots, n\}, \quad (4.8d)$$

where s_I is computed for each bag, as shown in Equation (4.9):

$$s_I = \underset{i \in I}{\operatorname{argmax}} \left(\sum_{K \in I} \sum_{j=1}^d \alpha_K Y_K x_{s_K j} x_{ij} + b \right), \forall I \in \{1, \dots, n\}. \quad (4.9)$$

The implicit constraints (4.8b) through (4.8d) imply three possible cases for α_I values:

1. If $\alpha_I = 0$, then $\beta_I = 0$ and $\xi_I = 0$, implying the instance is correctly classified and outside the margin.

2. If $0 < \alpha_I < C$, then $\xi_I = 0$, and the instance is a support vector and site on the margin boundary, i.e. is an *bounded support vector*.
3. If $\alpha_I = C$, then there is no restriction for $\xi_I > 0$. This also indicates that the instance is a support vector, but one that is *unbounded*. If $0 < \xi_I < 1$, then the instance is correctly classified, and is misclassified if $\xi \geq 1$.

The dual is then kernelized by replacing the inner product of samples in feature space with their corresponding kernel value, $\mathcal{K}(\mathbf{x}_{s_I}, \mathbf{x}_{s_K})$. The dual function is now written as:

$$\max_{\boldsymbol{\alpha}} \sum_I \alpha_I - \frac{1}{2} \sum_I \sum_{K \in I} \alpha_I \alpha_K Y_I Y_K \mathcal{K}(\mathbf{x}_{s_I}, \mathbf{x}_{s_K}) \quad (4.10)$$

$$\text{s.t.} \sum_I \alpha_I Y_I = 0 \quad (4.10a)$$

$$0 \leq \alpha_I \leq C, \forall I \in \{1, \dots, n\}. \quad (4.10b)$$

One of the biggest advantages of the dual SVM formulation is the sparseness of the resulting model. This is because support vectors, instances that have their corresponding $\alpha_I \neq 0$, are only considered when forming the decision boundary. MIRSVM uses a Gaussian RBF kernel, given by Equation (4.11), where σ is the Gaussian shape parameter.

$$\mathcal{K}(\mathbf{x}_i, \mathbf{x}_j) = e^{-\frac{\|\mathbf{x}_i - \mathbf{x}_j\|^2}{2\sigma^2}} \quad (4.11)$$

To evaluate the output vector, \mathbf{o}_I , of bag I using the kernel, the following equation is used, where \mathcal{B}_I are the instances of bag I , \mathbf{X}_S are the optimal bag representatives, and \mathbf{Y}_S are the representative bag labels.

$$\mathbf{o}_I = \mathcal{K}(\mathcal{B}_I, \mathbf{X}_S) * (\boldsymbol{\alpha} \cdot \mathbf{Y}_S) + b \quad (4.12)$$

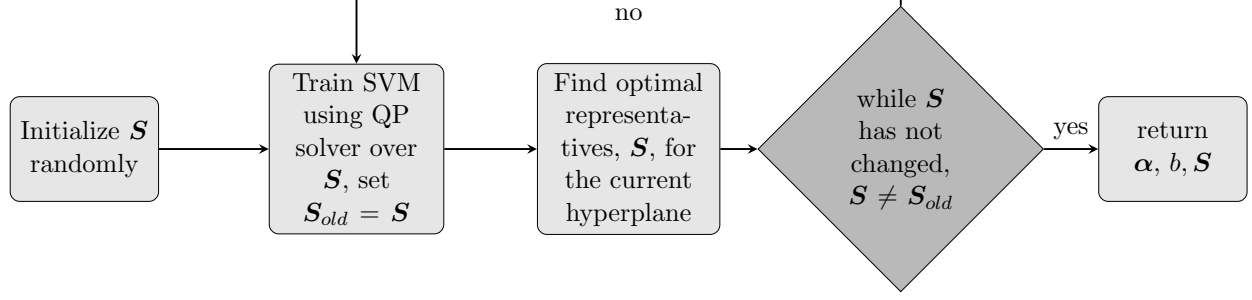


Fig. 4.1.: A summary of the steps performed by MIRSVM. The representatives are first randomly initialized and continuously updated according to the current hyperplane. Upon completion, the model is returned along with the optimal bag-representatives.

Algorithm 4.1 Multi-Instance Representative SVM (MIRSVM)

Input: Training dataset \mathcal{D} , SVM Parameters C and σ

Output: SVM model parameters α and b , Bag Representative IDs \mathbf{S}

```

for  $I \in \{1, \dots, n\}$  do
     $\mathbf{S}_I \leftarrow \text{rand}(|\mathcal{B}_I|, 1, 1)$  ▷ Assign each bag a random instance
end for
while  $\mathbf{S} \neq \mathbf{S}_{old}$  do
     $\mathbf{S}_{old} \leftarrow \mathbf{S}$ 
     $\mathbf{X}_S \leftarrow \mathbf{X}(\mathbf{S}), \mathbf{Y}_S \leftarrow \mathbf{Y}(\mathbf{S})$  ▷ Initialize the representative dataset
     $\mathbf{G} \leftarrow (\mathbf{Y}_S \times \mathbf{Y}_S) \cdot \mathcal{K}(\mathbf{X}_S, \mathbf{X}_S, \sigma)$  ▷ Build Gram matrix
     $\alpha \leftarrow \text{quadprog}(\mathbf{G}, -\mathbf{1}^n, \mathbf{Y}_S, \mathbf{0}^n, \mathbf{0}^n, C^n)$  ▷ Solve QP Problem
     $\mathbf{sv} \leftarrow \text{find}(0 < \alpha \leq C)$  ▷ Get the support vector indices
     $n_{sv} \leftarrow \text{count}(0 < \alpha \leq C)$  ▷ Get the number of support vectors
     $b \leftarrow \frac{1}{n_{sv}} \sum_{i=1}^{n_{sv}} (\mathbf{Y}_{sv} - \mathbf{G}_{sv} * (\alpha_{sv} \cdot \mathbf{Y}_{sv}))$  ▷ Calculate the bias term
    for  $I \in \{1, \dots, n\}$  do
         $\mathbf{G}_I \leftarrow (\mathbf{Y}_I \times \mathbf{Y}_S) \cdot \mathcal{K}(\mathcal{B}_I, \mathbf{X}_S, \sigma)$ 
         $\mathbf{S}_I \leftarrow \text{argmax}_{i \in I} (\mathbf{G}_I * \alpha + b)$  ▷ Select optimal bag-representatives
    end for
end while
  
```

The bias term b is calculated as shown in Equation (4.13), where \mathbf{sv} is the vector of support vector indices and n_{sv} is the number of support vectors [50].

$$b = \frac{1}{n_{sv}} \sum_{sv} Y_{sv} - \mathcal{K}(\mathbf{X}_{sv}, \mathbf{X}_{sv}) * (\alpha_{sv} \cdot Y_{sv}) \quad (4.13)$$

Algorithm 4.1 shows the procedure for training the multi-instance representative SVM classifier and obtaining the optimal representatives from each bag. During training, the representatives, \mathbf{S} , are first initialized by randomly selecting an instance from each bag. A hyperplane is then obtained using the representative instances, and new optimal rep-



Fig. 4.2.: Bag representative convergence plots on 9 datasets. The blue line shows the number of bag representatives that are equal from one iteration to the next. The red dashed line represents the total number of bags.

representatives are found with respect to the current hyperplane, by using the rule given in Equation (4.9). At each step, the previous values in \mathcal{S} are stored in \mathcal{S}_{old} . The training procedure ends when the bag representatives stop changing from one iteration to the next ($\mathcal{S} = \mathcal{S}_{old}$). Examples of the convergence of bag-representatives are shown in Figure 4.2. During the testing procedure, each bag produces an output vector based on the hyperplane found in the training procedure. The bag label is then assigned by taking the sign of the output vector's maximum value, following the SMI assumption.

This formulation is designed to utilize and select representatives from positive and negative bags, unlike MISVM, which only optimizes over representatives from positive bags, while flattening the negative bag instances. MISVM allows multiple representatives to be chosen from negative bags and limits positive bag-representatives to be one, while MIRSVM

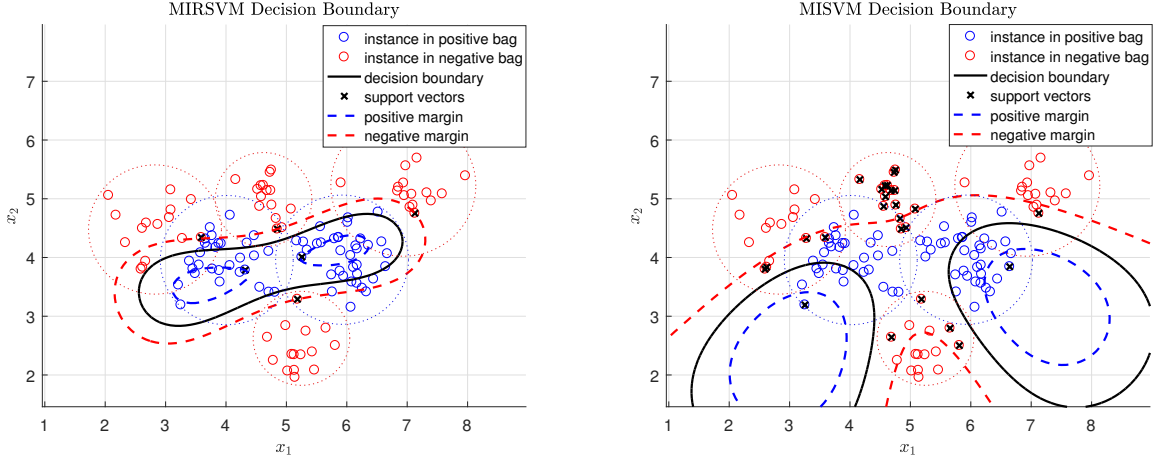


Fig. 4.3.: Difference between MIRSVM and MISVM on a random 2-dimensional toy dataset. Note the differing number of support vectors produced by the two methods. MIRSVM has 6, one for each bag, and MISVM has 29. Also note the smoother representation of the data distribution given by MIRSVM’s decision boundary, unlike MISVM whose decision boundary was greatly influenced by the larger number of support vectors belonging to the negative class with respect to the only 2 positive support vectors.

allows for balanced bag-representative selection, where each bag is allowed one. MISVM also uses a wrapper method to initialize the positive bag-representatives by taking the mean vector of the instances within each positive bag. This is an implicit assumption that the instances within the positive bags are all positive, whereas MIRSVM’s initialization procedure selects an instance from all bags at random, ensuring no noise is added by any wrapper techniques during initialization and no assumptions are made about the instances. Due to the constraints on the representatives, MIRSVM produces sparser models while MISVM has the freedom to select as many negative support vectors as it needs and restricts the support vectors chosen from positive bags to be one. Figure 4.3 shows the decision boundaries produced by MIRSVM and MISVM to highlight the differences in their solutions. As Figure 4.3 shows, MISVM produces a larger number of support vectors from the negative bags, which greatly influences the final decision boundary in favor of the negative class.

Table 4.2.: Multi-Instance (MI) Classification datasets

Dataset	Attributes	Positive Bags	Negative Bags	Total	Instances	Avg. Bag Size
Suramin	20	7	6	13	2898	222.92
EastWest	24	10	10	20	213	10.65
WestEast	24	10	10	20	213	10.65
Musk1	166	47	45	92	476	5.17
Musk2	166	39	62	101	6728	66.61
Webmining	5863	21	92	113	3423	30.29
Mutagenesis-atoms	10	125	63	188	1618	8.61
Mutagenesis-bonds	16	125	63	188	4081	21.71
Mutagenesis-chains	24	125	63	188	5424	28.85
TRX	8	25	168	193	26611	137.88
Elephant	230	100	100	200	1391	6.96
Fox	230	100	100	200	1320	6.60
Tiger	230	100	100	200	1188	5.94
Component	200	423	2707	3130	36894	11.79
Function	200	443	4799	5242	55536	10.59

4.3 Experimental Environment

This section presents the experimental setup and comparison of our contribution, as well as 11 other widely used methods on 15 different benchmark datasets. The main aim of the experiments is to compare our contribution to other multi-instance support vector machines, contemporary multi-instance learners, and ensemble methods.

Table 4.2 presents a summary of the 15 datasets used throughout the experiments, where the number of attributes, bags, and total number of instances are shown. The datasets were obtained from the Weka¹ [45] and KEEL² [2] dataset repositories.

The experimental environment was designed to test the difference in performance of the proposed method against 11 competing algorithms, contrasting instance-level, bag-level, and ensemble methods. Instance-level methods include MIOptimalBall, MIBoost, MISVM, MIDD, and MIWrapper. Bag-level methods include MISMO, SimpleMI, miGraph, and TLC. The ensemble-based bag-space methods, Bagging and Stacking, were also used. The base

¹<http://www.cs.waikato.ac.nz/ml/weka>

²<http://sci2s.ugr.es/keel/datasets.php>

algorithms selected for the ensembles Bagging and Stacking were TLC, and TLC and SimpleMI, respectively. These algorithms were chosen because they have shown considerable performance in learning multi-instance models, while also having their frameworks readily available for reproducing their results through MILK, the Multi-Instance Learning Kit³ [95], used in conjunction with the Weka framework. Experiments were run on an Intel i7-6700k CPU with 32GB RAM. MIRSVM was implemented in MATLAB while the referenced algorithms are available in the Java implementation of Weka with the exception of miGraph which was made available by Zhou et al.⁴ and tested in MATLAB.

We have compared the models trained on the different hyper-parameters using the cross-validation (CV) procedure which ensures that the models performances are accurately assessed and the model built is not biased towards the full dataset. The tuning of the model includes finding the best penalty parameter, C , as well as the best shape parameter for the Gaussian radial basis function kernel, σ . The best hyper-parameters were chosen from the following 6×6 possible combination runs, shown in Equations (4.14a) and (4.14b), referred to as (4.14).

$$C \in \{0.1, 1, 10, 100, 1000, 10000\} \quad (4.14a)$$

$$\sigma \in \{0.1, 0.5, 1, 2, 5, 10\} \quad (4.14b)$$

These parameters were also used for the compared SVM methods. This was done in order to keep the experimental environment controlled and ensure fair evaluation of the multi-instance SVM algorithms. The parameters for the referenced algorithms used throughout the experiments were those specified by their authors.

³<http://www.cs.waikato.ac.nz/ml/milk>

⁴http://lamda.nju.edu.cn/code_miGraph.ashx

4.4 Results & Statistical Analysis

The classification performance was measured using five metrics: Accuracy (4.15a), Precision (4.15b), Recall (4.15c), Cohen’s kappa rate (4.15d), and Area under ROC curve (AUC) (4.15e). The Precision and Recall measures were reported because Accuracy alone can be misleading when classes are imbalanced, as is the case with the *component* and *function* datasets, which respectively have six and ten times as many negative bags than positive. Cohen’s Kappa Rate and the AUC measures are used as complementary measures in order to evaluate the algorithms comprehensively. Cohen’s kappa rate, shown in Equation (4.15d), evaluates classifier merit according to the class distribution and ranges between -1 (full disagreement), 0 (random classification), and 1 (full agreement). The AUC metric highlights the trade-off between the true positive rate, or recall, and the false positive rate, as shown in Equation (4.15e). The values of the true positive (TP), true negative (TN), false positive (FP), and false negative samples (FN) were first collected for each of the classifiers, then the metrics were computed using the equations shown in (4.15) on the n' bags of the test data, where $n' = TP + FP + TN + FN$. The run times (training and testing times) of each algorithm are also reported to analyze the scalability and speed of each of the algorithms across differently sized datasets.

The results for the following are shown in Tables 4.3, 4.5, 4.7, 4.9, 4.11, and 4.13.

$$\text{Accuracy} \quad \frac{TP + TN}{n'} \quad (4.15a)$$

$$\text{Precision} \quad \frac{TP}{TP + FP} \quad (4.15b)$$

$$\text{Recall} \quad \frac{TP}{TP + FN} \quad (4.15c)$$

$$\text{Cohen's Kappa Rate} \quad \frac{n' - \frac{(TP + FN) * (TP + FP)}{n'}}{1 - \frac{(TP + FN) * (TP + FP)}{n'}} \quad (4.15d)$$

$$\text{Area Under ROC Curve} \quad \frac{1 + \frac{TP}{TP + FN} - \frac{FP}{FP + TN}}{2} \quad (4.15e)$$

In order to analyze the performances of the multiple models, non-parametric statistical tests are used to validate the experimental results obtained. The Iman-Davenport non-parametric test is run to investigate whether significant differences exist among the performance of the algorithms by ranking them over the datasets used, using the Friedman test. The algorithm ranks for each metric in Equations (4.15) are presented in the last row of the results tables, and the lowest (best) rank value is typeset in bold. Table 4.14 contains the ranks and meta-rank of all methods, which helps determine and visualize the best performing algorithms across all datasets and metrics.

After the Iman-Davenport test indicates significant differences, the Bonferroni-Dunn post-hoc test [36] is then used to find where they occur between algorithms by assuming the classifiers' performances are different by at least some critical value. Below each result table, a figure highlighting the critical distance (in gray), from the best ranking algorithm to the rest, is shown. The algorithms to the right of the critical distance bar perform statistically significantly worse than the control algorithm, MIRSVM. Figures 4.4, 4.5, 4.6, 4.7, 4.8, 4.9 show the results of the Bonferroni-Dunn post-hoc procedure over the metrics in (4.15), as well as the meta-rank results in Table 4.14. The Holm (multiple) and Wilcoxon

(pairwise) rank-sum post-hoc tests [49] were then run for each of the metrics to compute multiple and pairwise comparisons between the proposed algorithm and the other methods compared, investigating whether statistical differences exist among the algorithms' results. Tables 4.4, 4.6, 4.8, 4.10, and 4.12 show the p -values for the Holm test for $\alpha = 0.05$, and the rank-sums and adjusted p -values for the Wilcoxon test.

4.4.1 Accuracy

The results for accuracy indicate that the bag-based and ensemble learners perform better than the instance-based and wrapper methods. MIRSVM achieves the best accuracy over 5 of the 15 datasets with a competitive average against miGraph, Bagging, Stacking, and TLC. Note that MIRSVM performs better than MISVM for all datasets, indicating that using representatives from each bag and limiting the number of support vectors per negative bag improves the classification performance. The instance-level classifiers and wrapper methods,

Table 4.3.: Accuracy for MI classifiers

Datasets	MIRSVM	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
suramin	0.8000	0.8462	0.5000	0.7250	0.4250	0.5000	0.7250	0.5000	0.5000	0.6000	0.6650	0.4615
eastWest	0.8000	0.7000	0.5000	0.7250	0.6125	0.5000	0.7125	0.5625	0.5000	0.6000	0.6000	0.4500
westEast	0.7500	0.7500	0.5000	0.3750	0.4500	0.5000	0.7375	0.4125	0.5000	0.5625	0.9649	0.6375
musk1	0.9022	0.8152	0.5109	0.7717	0.8804	0.5109	0.7826	0.7609	0.5109	0.8587	0.8142	0.8587
musk2	0.8218	0.7426	0.6139	0.7723	0.7228	0.6139	0.7030	0.7129	0.6139	0.6238	0.8756	0.6733
webmining	0.8500	0.8142	0.8142	0.7699	0.8142	0.8142	0.8407	0.6903	0.8142	0.8142	0.9358	0.8053
trx	0.8860	0.8964	0.8705	0.9016	0.8808	0.8705	0.8705	0.8705	0.8705	0.8756	0.6450	0.8860
mutagenesis-atoms	0.7714	0.7606	0.6649	0.6436	0.7074	0.6649	0.6915	0.6649	0.6649	0.7766	0.7766	0.7606
mutagenesis-bonds	0.8252	0.7872	0.6649	0.6915	0.7713	0.6649	0.7979	0.6649	0.6649	0.8351	0.8351	0.8564
mutagenesis-chains	0.8411	0.7926	0.6649	0.6702	0.7766	0.6649	0.8351	0.6649	0.6649	0.8404	0.8404	0.8351
tiger	0.7750	0.7950	0.5000	0.5000	0.7100	0.5000	0.7200	0.7550	0.5000	0.6650	0.8000	0.7250
elephant	0.8300	0.8300	0.5000	0.5000	0.7900	0.5000	0.8100	0.8000	0.5000	0.8000	0.5625	0.8250
fox	0.6550	0.6300	0.5000	0.5000	0.5800	0.5000	0.5250	0.4750	0.5000	0.6450	0.8587	0.6500
component	0.9366	0.9153	0.8649	0.8696	0.8780	0.8649	0.8968	0.8703	0.8649	0.9358	0.6000	0.9355
function	0.9523	0.9405	0.9155	0.9138	0.9193	0.9155	0.9376	0.9195	0.9155	0.9649	0.6238	0.9647
Average	0.8264	0.8010	0.6390	0.6886	0.7279	0.6390	0.7724	0.6883	0.6390	0.7598	0.7598	0.7550
Rank	2.2000	3.8667	9.6000	7.8667	6.5667	9.6000	5.3333	8.5667	9.6000	4.7000	4.8667	5.2333



Fig. 4.4.: Bonferroni-Dunn test for Accuracy

Table 4.4.: Holm and Wilcoxon tests for Accuracy

MIRSVM vs.	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
Holm p -value	0.0500	0.0045	0.0071	0.0083	0.0050	0.0100	0.0063	0.0056	0.0250	0.0167	0.0125
Wilcoxon p -value	0.0279	0.0001	0.0001	0.0001	0.0001	0.0001	0.0001	0.0001	0.0067	0.3028	0.0103
Wilcoxon R^+	98.500	120.00	119.00	120.00	120.00	120.00	120.00	120.00	106.00	79.000	104.00
Wilcoxon R^-	21.500	0.0000	1.0000	0.0000	0.0000	0.0000	0.0000	0.0000	14.000	41.000	16.000

such as MIBoost, MIWrapper, and SimpleMI perform the worst. This behavior emphasizes the importance of not making prior assumptions about the positive bags’ distributions.

Figure 4.4 and Table 3.4 show the results for the statistical analysis on the accuracy results. The algorithms with ranking higher than 5.63 (MIRSVM rank + Bonferroni-Dunn critical value), to the right of the grey bar in Figure 4.4, perform statistically worse than MIRSVM. Table 3.4 shows the p -values of the Holm and Wilcoxon tests and their results complement one another. Holm’s procedure rejects those hypotheses having a p -value ≤ 0.01 , thus indicating that MIRSVM performs significantly better than all methods except miGraph, Bagging, Stacking, and TLC. The Wilcoxon p -values show significant differences exist among all algorithms except miGraph, Bagging, and Stacking. They also show that MIRSVM has significantly better accuracy than MIBoost, MIOptimalBall, MIDD, MIWrapper, MISMO, MISVM, and SimpleMI, each having respectively small p -values, highlighting MIRSVM’s superior classification accuracy.

4.4.2 Precision & Recall

Precision and recall are conflicting metrics that must be evaluated together in order to observe their behavior, since they are both used to measure relevance. The results for MIWrapper and SimpleMI indicate that they are unstable classifiers, exhibiting extreme variance in behavior, making them unsuitable for real-world applications. It is also interesting to analyze the performance on the mutagenesis datasets which have a larger number of positive bags than negative, where MISVM, MIBoost, MIWrapper, and SimpleMI predict all bags as negative. Additionally, while MISMO obtains unbiased results on these datasets, MIRSVM significantly outperforms it over both precision and recall, achieving a better trade-off.

Table 4.5.: Precision for MI classifiers

Datasets	MIRSVM	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
suramin	0.7778	0.7778	1.0000	1.0000	0.2857	1.0000	1.0000	0.5000	1.0000	0.6429	0.6514	0.4000
eastWest	0.7143	0.7000	0.5000	0.8750	0.5882	0.5000	0.7429	1.0000	0.5000	0.6053	0.6053	0.4444
westEast	0.7272	0.7273	0.5000	0.2727	0.4600	0.5000	0.6939	0.3600	0.5000	0.5581	0.9729	0.6038
musk1	0.8519	0.7778	1.0000	0.9286	0.9048	1.0000	0.8049	0.8108	1.0000	0.8478	0.8817	0.8478
musk2	0.7059	0.7826	0.6139	0.7826	0.7576	0.6139	0.7424	0.7538	0.6139	0.7400	0.9138	0.7164
webmining	0.7500	1.0000	0.8142	0.8173	0.8142	0.8142	0.8936	1.0000	0.8142	0.8817	0.9462	0.8500
trx	1.0000	0.8571	0.8705	0.9306	0.9191	0.8705	0.8705	0.8705	0.8705	0.9138	0.6747	0.9011
mutagenesis-atoms	0.7872	0.7985	1.0000	0.4630	0.6111	1.0000	0.5439	1.0000	1.0000	0.7059	0.7059	0.6667
mutagenesis-bonds	0.8468	0.8195	1.0000	0.5385	0.7500	1.0000	0.6812	1.0000	1.0000	0.7857	0.7857	0.8333
mutagenesis-chains	0.8571	0.8116	1.0000	0.5091	0.7059	1.0000	0.7759	1.0000	1.0000	0.7705	0.7705	0.7581
tiger	0.7365	0.7323	0.5000	0.5000	0.6944	0.5000	0.7444	0.7802	0.5000	0.6514	0.8000	0.7320
elephant	0.8576	0.8750	0.5000	0.5000	0.7959	0.5000	0.8444	0.7679	0.5000	0.8000	0.5581	0.8283
fox	0.6040	0.6275	0.5000	0.5000	0.5833	0.5000	0.5287	0.4854	0.5000	0.6747	0.8478	0.6705
component	0.9866	0.7782	0.8649	0.8778	0.8902	0.8649	0.8958	0.8696	0.8649	0.9462	0.6429	0.9449
function	0.8459	0.6775	0.9155	0.9202	0.9317	0.9155	0.9376	0.9197	0.9155	0.9729	0.7400	0.9726
Average	0.8033	0.7828	0.7719	0.6944	0.7128	0.7719	0.7800	0.8079	0.7719	0.7665	0.7665	0.7447
Rank	5.3333	6.1333	7.1000	7.3333	7.3667	7.1000	5.8667	5.8667	7.1000	5.9000	6.3333	6.5667



Fig. 4.5.: Bonferroni-Dunn test for Precision

Table 4.6.: Holm and Wilcoxon tests for Precision

MIRSVM vs.	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
Holm p -value	0.0125	0.0056	0.0050	0.0045	0.0063	0.0250	0.0500	0.0071	0.0167	0.0100	0.0083
Wilcoxon p -value	0.4212	0.5614	0.0946	0.0256	0.5614	0.4212	0.8039	0.5614	0.1354	0.4543	0.1354
Wilcoxon R^+	75.000	71.000	90.000	99.000	71.000	75.000	55.000	71.000	87.000	74.000	87.000
Wilcoxon R^-	45.000	49.000	30.000	21.000	49.000	45.000	65.000	49.000	33.000	46.000	33.000

Table 4.7.: Recall for MI classifiers

Datasets	MIRSVM	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
suramin	1.0000	1.0000	0.0000	0.4500	0.1000	0.0000	0.4500	0.5000	0.0000	0.4500	0.7100	0.3333
eastWest	1.0000	0.7000	0.7000	0.5250	0.7500	0.7000	0.6500	0.1250	1.0000	0.5750	0.5750	0.4000
westEast	0.8000	0.8000	0.9000	0.1500	0.5750	0.9000	0.8500	0.2250	1.0000	0.6000	0.9892	0.8000
musk1	0.9787	0.8936	0.0000	0.5778	0.8444	0.0000	0.7333	0.6667	0.0000	0.8667	0.8913	0.8667
musk2	0.9231	0.4615	1.0000	0.8710	0.8065	1.0000	0.7903	0.7903	1.0000	0.5968	0.9464	0.7742
webmining	0.2857	0.0000	1.0000	0.9239	1.0000	1.0000	0.9130	0.6196	1.0000	0.8913	0.9815	0.9239
trx	0.4833	0.2400	1.0000	0.9583	0.9464	1.0000	1.0000	1.0000	1.0000	0.9464	0.5600	0.9762
mutagenesis-atoms	0.8880	0.8560	0.0000	0.3968	0.3492	0.0000	0.4921	0.0000	0.0000	0.5714	0.5714	0.5714
mutagenesis-bonds	0.8960	0.8720	0.0000	0.5556	0.4762	0.0000	0.7460	0.0000	0.0000	0.6984	0.6984	0.7143
mutagenesis-chains	0.9120	0.8960	0.0000	0.4444	0.5714	0.0000	0.7143	0.0000	0.0000	0.7460	0.7460	0.7460
tiger	0.8700	0.9300	0.5000	1.0000	0.7500	0.5000	0.6700	0.7100	1.0000	0.7100	0.8000	0.7100
elephant	0.9100	0.7700	0.6000	1.0000	0.7800	0.6000	0.7600	0.8600	1.0000	0.8000	0.6000	0.8200
fox	0.9000	0.6400	0.7000	1.0000	0.5600	0.7000	0.4600	0.8300	1.0000	0.5600	0.8667	0.5900
component	0.5839	0.5225	1.0000	0.9867	0.9797	1.0000	0.9967	1.0000	1.0000	0.9815	0.4500	0.9826
function	0.5327	0.5643	1.0000	0.9919	0.9840	1.0000	0.9983	0.9994	1.0000	0.9892	0.5968	0.9894
Average	0.7976	0.6764	0.5600	0.7221	0.6982	0.5600	0.7483	0.5551	0.6667	0.7322	0.7322	0.7465
Rank	4.8667	6.5667	6.8667	6.3333	7.3667	6.8667	6.7000	7.4333	4.8333	7.3667	6.0667	6.7333



Fig. 4.6.: Bonferroni-Dunn test for Recall

Table 4.8.: Holm and Wilcoxon tests for Recall

MIRSVM vs.	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
Holm p -value	0.0125	0.0063	0.0167	0.0050	0.0071	0.0100	0.0045	0.0500	0.0056	0.0250	0.0083
Wilcoxon p -value	0.0060	0.2077	0.5614	0.4543	0.2077	0.6387	0.1070	0.6603	0.5995	0.1354	0.5721
Wilcoxon R^+	106.50	83.000	71.000	74.000	83.000	69.000	89.000	60.000	70.000	87.000	62.000
Wilcoxon R^-	13.500	37.000	49.000	46.000	37.000	51.000	31.000	45.000	50.000	33.000	43.000

Figure 4.5 and 4.6 show that there are no significant differences between the precision and recall results obtained by all algorithms. Note, MIRSVM outperforms both ensemble methods according to recall, despite them exhibiting good accuracy and precision, indicating they are strongly conservative towards predicting positive bags. Holm’s test indicates significant differences exist between MIRSVM and all algorithms except miGraph, MISMO, MISVM, and TLC for precision, and all the above along with SimpleMI, MIOptimalBall, and Bagging for recall. The Wilcoxon test does not reflect significant differences for precision, does for recall. The tests are severely biased due to the classifier’s extreme unbalanced behavior, whereas MIRSVM demonstrates proper balance of the precision-recall trade-off.

4.4.3 Cohen’s Kappa Rate

Table 4.9 shows the Cohen’s Kappa rate results obtained by the algorithms. These results support the accuracy achieved by the algorithms, in the sense that the instance-based

Table 4.9.: Cohen’s Kappa Rate for MI classifiers

Datasets	MIRSVM	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
suramin	0.6829	0.6829	0.0000	0.4500	-0.1500	0.0000	0.4500	0.0000	0.0000	0.2000	0.3300	-0.0964
eastWest	0.6000	0.4000	0.0000	0.4500	0.2250	0.0000	0.4250	0.1250	0.0000	0.2000	0.2000	-0.1000
westEast	0.5000	0.5000	0.0000	-0.2500	-0.1000	0.0000	0.4750	-0.1750	0.0000	0.1250	0.7529	0.2750
musk1	0.8036	0.6290	0.0000	0.5396	0.7604	0.0000	0.5642	0.5197	0.0000	0.7174	0.3744	0.7174
musk2	0.6540	0.4123	0.0000	0.5031	0.4039	0.0000	0.3613	0.3856	0.0000	0.2492	0.3858	0.2940
webmining	0.3468	0.0000	0.0000	0.0246	0.0000	0.0000	0.4535	0.3771	0.0000	0.3744	0.6945	0.2458
trx	0.2100	0.3375	0.0000	0.5228	0.4224	0.0000	0.0000	0.0000	0.0000	0.3858	0.2900	0.3364
mutagenesis-atoms	0.5395	0.4431	0.0000	0.1709	0.2654	0.0000	0.2909	0.0000	0.0000	0.4738	0.4738	0.4431
mutagenesis-bonds	0.5699	0.5070	0.0000	0.3131	0.4356	0.0000	0.5569	0.0000	0.0000	0.6195	0.6195	0.6659
mutagenesis-chains	0.6303	0.5094	0.0000	0.2359	0.4738	0.0000	0.6225	0.0000	0.0000	0.6391	0.6391	0.6285
tiger	0.5500	0.5900	0.0000	0.0000	0.4200	0.0000	0.4400	0.5100	0.0000	0.3300	0.6000	0.4500
elephant	0.7000	0.6600	0.0000	0.0000	0.5800	0.0000	0.6200	0.6000	0.0000	0.6000	0.1250	0.6500
fox	0.3100	0.2600	0.0000	0.0000	0.1600	0.0000	0.0500	-0.0500	0.0000	0.2900	0.7174	0.3000
component	0.6644	0.5795	0.0000	0.1613	0.2836	0.0000	0.3656	0.0675	0.0000	0.6945	0.2000	0.6906
function	0.6292	0.5838	0.0000	0.0966	0.2801	0.0000	0.4083	0.0933	0.0000	0.7529	0.2492	0.7507
Average	0.5594	0.4730	0.0000	0.2145	0.2973	0.0000	0.4056	0.1635	0.0000	0.4434	0.4434	0.4167
Rank	2.6333	4.2000	10.1667	7.0000	6.5333	10.1667	5.2333	8.3667	10.1667	4.2667	4.2667	5.0000



Fig. 4.7.: Bonferroni-Dunn test for Cohen’s Kappa rate

Table 4.10.: Holm and Wilcoxon tests for Cohen’s Kappa rate

MIRSVM vs.	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
Holm p -value	0.0500	0.0045	0.0071	0.0083	0.0050	0.0100	0.0063	0.0056	0.0167	0.0250	0.0125
Wilcoxon p -value	0.0121	0.0001	0.0012	0.0012	0.0001	0.0006	0.0001	0.0001	0.1205	0.2077	0.0946
Wilcoxon R^+	91.500	120.00	113.00	113.00	120.00	115.00	119.00	120.00	88.000	83.000	90.000
Wilcoxon R^-	13.500	0.0000	7.0000	7.0000	0.0000	5.0000	1.0000	0.0000	32.000	37.000	30.000

and wrapper methods perform worse than bag-based and ensemble learners. MIRSVM's kappa values all fall within the range (0.5-1], indicating that its merit as a classifier agrees with the class distribution and is not random. Note that MIOptimalBall, MIDD, MISVM, MISMO, and Stacking contain some negative kappa values, indicating performance worse than the default-hypothesis. MIBoost, SimpleMI, and MIWrapper are shown to randomly classify all 15 datasets. Figure 4.7 and Table 4.10 show the results of the statistical analysis on the Cohen's Kappa Rate results. The Holm and Wilcoxon procedures reflect results similar to the Bonferroni-Dunn test, where MIRSVM performs significantly better than MIOptimalBall, MIDD, MISVM, MIWrapper, MIBoost, and SimpleMI, having p -values < 0.01. This supports MIRSVM's performance as a competitive classifier.

Table 4.11.: AUC for MI classifiers

Datasets	MIRSVM	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
suramin	0.8333	0.8333	0.5000	0.7250	0.4250	0.5000	0.7250	0.5000	0.5000	0.6000	0.6650	0.4524
eastWest	0.8000	0.7000	0.5000	0.7250	0.6125	0.5000	0.7125	0.5625	0.5000	0.6000	0.6000	0.4500
westEast	0.7500	0.7500	0.5000	0.3750	0.4500	0.5000	0.7375	0.4125	0.5000	0.5625	0.8456	0.6375
musk1	0.9005	0.8135	0.5000	0.7676	0.8797	0.5000	0.7816	0.7589	0.5000	0.8589	0.6837	0.8589
musk2	0.8406	0.6904	0.5000	0.7432	0.6981	0.5000	0.6772	0.6900	0.5000	0.6317	0.6732	0.6435
webmining	0.6320	0.5000	0.5000	0.5096	0.5000	0.5000	0.7184	0.8098	0.5000	0.6837	0.8123	0.6048
trx	0.6500	0.6170	0.5000	0.7392	0.6932	0.5000	0.5000	0.5000	0.5000	0.6732	0.6450	0.6281
mutagenesis-atoms	0.7106	0.7137	0.5000	0.5824	0.6186	0.5000	0.6420	0.5000	0.5000	0.7257	0.7257	0.7137
mutagenesis-bonds	0.7856	0.7455	0.5000	0.6578	0.6981	0.5000	0.7850	0.5000	0.5000	0.8012	0.8012	0.8211
mutagenesis-chains	0.8252	0.7417	0.5000	0.6142	0.7257	0.5000	0.8051	0.5000	0.5000	0.8170	0.8170	0.8130
tiger	0.7750	0.7950	0.5000	0.5000	0.7100	0.5000	0.7200	0.7550	0.5000	0.6650	0.8000	0.7250
elephant	0.8200	0.8300	0.5000	0.5000	0.7900	0.5000	0.8100	0.8000	0.5000	0.8000	0.5625	0.8250
fox	0.6550	0.6300	0.5000	0.5000	0.5800	0.5000	0.5250	0.4750	0.5000	0.6450	0.8589	0.6500
component	0.7855	0.7496	0.5000	0.5536	0.6033	0.5000	0.6272	0.5201	0.5000	0.8123	0.6000	0.8081
function	0.7563	0.7698	0.5000	0.5298	0.6015	0.5000	0.6391	0.5268	0.5000	0.8456	0.6317	0.8434
Average	0.7680	0.7253	0.5000	0.6015	0.6390	0.5000	0.6937	0.5874	0.5000	0.7148	0.7148	0.6983
Rank	2.7667	4.2667	10.1667	7.0000	6.5333	10.1667	5.2333	8.2333	10.1667	4.2667	4.2667	4.9333

Fig. 4.8.: Bonferroni-Dunn test for AUC

Table 4.12.: Holm and Wilcoxon tests for AUC

MIRSVM vs.	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
Holm p -value	0.0167	0.0045	0.0071	0.0083	0.0050	0.0100	0.0063	0.0056	0.0250	0.0500	0.0125
Wilcoxon p -value	0.0166	0.0001	0.0002	0.0003	0.0001	0.0012	0.0009	0.0001	0.2523	0.3028	0.0781
Wilcoxon R^+	90.000	120.00	118.00	117.00	120.00	113.00	114.00	120.00	81.000	79.000	91.500
Wilcoxon R^-	15.000	0.0000	2.0000	3.0000	0.0000	7.0000	6.0000	0.0000	39.000	41.000	28.500

4.4.4 Area Under ROC Curve

Table 4.11 shows AUC results obtained by the algorithms, which complement the accuracy and kappa rate, emphasizing the better performance of bag-based methods. MIRSVM achieves the best AUC score on 5 of the 15 datasets, while MIBoost, SimpleMI, and MIWrapper obtain the worst results. Their AUC score indicates random predictor behavior, having values = 0.5. Bag-level methods all obtain scores between 0.7 and 0.77 indicating a high true positive rate and a low false positive rate, which is reflected by the precision and recall results. Figure 4.8 and Table 4.12 show that MIRSVM performs significantly better than 6 out of the 11 competing algorithms. Holm’s procedure indicates that significant differences exist between MIRSVM and all algorithms except miGraph, TLC, Bagging, and Stacking. MISVM’s true positive rate could be affected because of the possible imbalance of support vectors from the positive and negative classes (favoring the negative). Note that the Wilcoxon p -values for MIWrapper, MIBoost, and SimpleMI are 0.0001.

4.4.5 Overall Comparison

Table 4.13 shows the run times, in seconds, for each algorithm. MIRSVM has the fastest run time and is ranked second. MIRSVM shows very good scalability considering the number of features, such as in the webmining dataset which comprises of 5863 attributes. Additionally, taking into account the number of instances as seen in the two largest datasets, component and function, MIRSVM displays superior scalability. It is important to note that quadratic programming solvers are not the most efficient tools for solving optimization problems in terms of run time, and yet MIRSVM still is shown to perform competitively against the current widely used algorithms. The scalability of MIRSVM is founded on the speedy rate of bag-representative convergence, as shown previously in Figure 4.2.

SimpleMI achieves the highest rank and competitive run times because, rather than use the instances in each bag to train a model, it takes the mean value of the instances in a bag and uses that for training. Even though SimpleMI has fast run-times, its performance

Table 4.13.: Run Time (seconds) for MI classifiers

Datasets	MIRSVM	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
suramin	0.1	19.7	8.8	30.5	7922.0	9.5	52.3	333.9	7.2	35.5	183.0	90.6
eastWest	0.1	3.0	5.5	9.4	217.1	6.3	14.8	21.4	5.8	15.4	15.4	15.2
westEast	0.1	2.8	6.5	7.8	79.7	6.5	14.7	99.5	6.0	16.6	12128.1	10.8
musk1	0.4	56.8	13.4	32.1	3542.6	20.6	89.7	198.4	11.1	93.0	86272.6	759.5
musk2	2.3	452.3	97.3	782.9	126016.8	208.3	1799.4	26093.5	16.1	1772.2	2229.3	16759.0
webmining	300.6	302.5	45745.4	60474.8	47601.4	68736.7	51923.6	105622.3	2685.9	86272.6	9861.5	592948.9
trx	61.8	2206.4	17.6	682.3	339110.5	19.3	8670.3	134622.1	7.4	2229.3	243.3	11927.9
mutagenesis-atoms	9.8	193.1	8.8	99.2	2623.0	8.0	55.0	53.5	6.4	44.0	44.0	153.9
mutagenesis-bonds	8.3	410.3	10.2	310.2	17538.7	12.3	457.4	2794.8	8.4	131.1	131.1	853.1
mutagenesis-chains	19.3	513.4	12.0	525.0	48982.7	14.8	2451.9	6637.4	7.2	224.4	224.4	1619.0
tiger	29.5	302.8	44.5	157.8	23220.5	56.2	208.0	608.8	16.2	183.0	212.1	1085.0
elephant	47.7	306.7	45.5	243.9	56456.2	69.7	232.1	1114.3	20.8	212.1	16.6	1462.2
fox	81.0	303.1	44.2	206.1	27773.8	66.0	369.6	891.5	23.5	243.3	93.0	1729.1
component	231.7	3091.0	572.5	228209.6	96263.9	1096.9	629366.4	37224.6	144.0	9861.5	35.5	79149.8
function	740.3	8162.7	935.5	768458.0	350124.7	1887.5	1052225.3	565026.4	232.8	12128.1	1772.2	185918.5
Average	102.2	1088.4	3171.2	70682.0	76498.2	4814.6	116528.7	58756.2	213.3	7564.1	7564.1	59632.2
Rank	2.3	6.2	3.1	7.2	11.1	4.3	8.5	10.1	1.9	7.2	6.5	9.7

Table 4.14.: Overall ranks comparison for MI classifiers

Ranks	MIRSVM	miGraph	MIBoost	MIOptimalBall	MIDD	MIWrapper	MISMO	MISVM	SimpleMI	TLC	Bagging	Stacking
Accuracy	2.2000	3.8667	9.6000	7.8667	6.5667	9.6000	5.3333	8.5667	9.6000	4.7000	4.8667	5.2333
Precision	5.3333	6.1333	7.1000	7.3333	7.3667	7.1000	5.8667	5.8667	7.1000	5.9000	6.3333	6.5667
Recall	4.8667	6.5667	6.8667	6.3333	7.3667	6.8667	6.7000	7.4333	4.8333	7.3667	6.0667	6.7333
Kappa	2.6333	4.2000	10.1667	7.0000	6.5333	10.1667	5.2333	8.3667	10.1667	4.2667	4.2667	5.0000
AUC	2.7667	4.2667	10.1667	7.0000	6.5333	10.1667	5.2333	8.2333	10.1667	4.2667	4.2667	4.9333
Time	2.2667	6.2000	3.1000	7.2000	11.0667	4.3000	8.5333	10.1333	1.8667	7.2000	6.4667	9.6667
Average	3.3444	5.2056	7.8333	7.1222	7.5722	8.0333	6.1500	8.1000	7.2889	5.6167	5.3778	6.3556
Rank	1.3333	3.6667	8.9167	7.7500	9.2500	9.0833	5.9167	8.7500	7.3333	5.2500	4.2500	6.5000



Fig. 4.9.: Bonferroni-Dunn test for overall ranks comparison

over the previous metrics has been shown to be random and not as effective as the bag-level methods.

Table 4.14 shows the ranks achieved by each of the metrics along with the average and meta-ranks, to illustrate the overall performance across all metrics. MIRSVM has the best meta-rank (rank of the ranks) and the miGraph method has the second best. The meta-ranks also highlight the better performance of bag-level methods over instance-level and wrapper methods, emphasizing the importance of training at the bag-level. Not only does MIRSVM use bag-level information during classification, but it also optimizes over the instances within the bag, which helps determine which instances contribute the most information about the bags label. SimpleMI, MIWrapper, MIBoost, MISVM, and MDD have the worst performance compared to MIRSVM and miGraph. Specifically, it is evident from the precision and recall

results that MIBoost, MIWrapper, and SimpleMI, for example, classify all bags as negative for datasets that have imbalanced class distributions which favor the negative class. This emphasizes the disadvantage of using wrapper methods and assuming the data distribution of the instances within positive bags. Although these algorithms are popular in literature, the experimental study clearly shows that recent bag-level and ensemble methods easily overcome traditional multi-instance learning algorithms.

In summary, MIRSVM offers improvement in terms of both accuracy and run-time when compared to referenced methods, especially those utilizing SVM-based algorithms.

4.5 Conclusions

This proposal consisted of a novel formulation and algorithm for the multiple-instance support vector machine problem, which optimizes bag classification via bag-representative selection. First, the primal formulation was posed and its dual was then derived and solution computed using a quadratic programming solver. This formulation was designed to utilize bag-level information and find an optimal separating hyperplane between bags, rather than individual instances, using the standard multi-instance assumption. The SMI assumption states that a bag is labeled positive if and only if at least one instance within a bag is positive, and is negative otherwise. The key features of the proposed algorithm MIRSVM are its ability to identify instances within positive and negative bags, i.e. the support vectors or representatives, that highly impact the decision boundary and margin, as well as avoiding uncertainties and issues caused by techniques that flatten, subset, or under-represent positive instances within positively labeled bags. Additionally, it exhibits desirable convergence and scalability, making it suitable for large-scale learning tasks.

The experimental study showed the better performance of MIRSVM compared with existing multi-instance support vector machines, traditional multi-instance learners, as well as ensemble methods. The results, according to a variety of performance metrics, were compared and further validated using statistical analysis with non-parametric tests which

highlight the advantages of using bag-level based and ensemble learners, such as miGraph, Bagging, and Stacking, while showing the instance-level based learners performed poorly in comparison or were deemed as strongly biased and unstable classifiers. Our proposal, MIRSVM, performs statistically better, neither compromising accuracy nor run-time while displaying a robust performance across all of the evaluated datasets. The research outcomes of this chapter have been published in [66].

CHAPTER 5

NOVEL ONLINE SVM USING WORST-VIOLATORS

Due to the ever-growing nature of dataset sizes, the need for scalable and accurate machine learning algorithms has become evident. Stochastic gradient descent methods are popular tools used to optimize large-scale learning problems because of their generalization performance, simplicity, and scalability. This chapter proposes a novel stochastic, also known as online, learning algorithm for solving the L1 support vector machine (SVM) problem, named *OnLine Learning Algorithm using Worst-Violators* (OLLAWV). The scope of this chapter is concerned with developing a unique learning algorithm for large data problems without the use of parallelization techniques and distributed systems. Unlike other stochastic methods, OLLAWV eliminates the need for specifying the maximum number of iterations and the use of a regularization term. OLLAWV uses early stopping for controlling the size of the margin instead of the regularization term. The experimental study, performed under very strict nested cross-validation (a.k.a., double resampling), evaluates and compares the performance of this proposal with state-of-the-art SVM kernel methods that have been shown to outperform traditional and widely used approaches for solving L1-SVMs such as *Sequential Minimal Optimization*. OLLAWV is also compared to 5 other traditional non-SVM algorithms. The results over 23 datasets show OLLAWV's superior performance in terms of accuracy, scalability, and model sparseness, making it suitable for large-scale learning.

5.1 Online Learning Background

Since a comprehensive overview of contemporary online learning algorithms were presented in Chapter 2, this section will present the concept of Stochastic Gradient Descent (SGD) for convex learning problems such as support vector machines.

Table 5.1.: Summary of notation used throughout the paper

Definition	Notation
Number of Samples	n
Number of Input Attributes	d
Input Space	$\mathbf{X} \in \mathbb{R}^{n \times d}$
Labels	$\mathbf{Y} \in \{-1, 1\}^n$
Sample i	$\mathbf{x}_i = (x_{i1}, \dots, x_{id}), \forall i \in \{1, \dots, n\}$
Sample Label i	$y_i \in \{-1, 1\}, \forall i \in \{1, \dots, n\}$
Full Training Dataset	$\mathcal{D} = \{(\mathbf{x}_1, y_1), \dots, (\mathbf{x}_i, y_i), \dots, (\mathbf{x}_n, y_n)\}$

5.1.1 Notation

Let \mathcal{D} be the full training dataset of n d -dimensional samples. Let $\mathbf{Y} \in \mathcal{D}$ be a vector of n labels corresponding to each sample, such that $\mathbf{Y} \in \{-1, 1\}^n$. In the non-binary (more than two classes) classification cases, $\mathbf{Y} \in \mathbb{Z}^n$. Let $\mathbf{X} \in \mathcal{D}$ be a matrix consisting of n samples that are d -dimensional, $\mathbf{X} \in \mathbb{R}^{n \times d}$.

5.1.2 Stochastic Gradient Descent

In the context of convex learning problems, such as RLM, the advantage of SGD is that it is an efficient and simple algorithm that has a good sample complexity. In this section we will first describe gradient descent, subgradients for non-differentiable function, and finally describe the SGD algorithm with some variants.

The approach for minimizing a convex, differentiable function $f(\mathbf{w})$ is an iterative one where the gradient of f is taken at each step and used to update the variable to be minimized. The gradient of function $f : \mathbb{R}^d \rightarrow \mathbb{R}$ at \mathbf{w} , is denoted and defined by $\nabla f(\mathbf{w}) = \left(\frac{\partial f(\mathbf{w})}{\partial w[1]}, \dots, \frac{\partial f(\mathbf{w})}{\partial w[d]} \right)$, the vector of partial derivatives of f with respect to \mathbf{w} . The Gradient Descent (GD) algorithm is as follows: \mathbf{w} is first initialized to some random value (say $\mathbf{w}^{(1)} = 0$), and at each iteration, a step is taken in the negative direction of the gradient at that point, as shown in equation 5.1.

$$\mathbf{w}^{(t+1)} = \mathbf{w}^{(t)} - \eta \nabla f(\mathbf{w}^{(t)}) \quad (5.1)$$

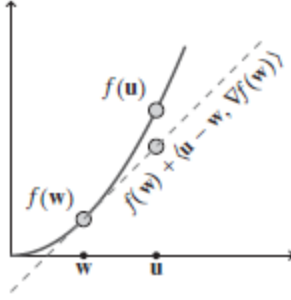


Fig. 5.1.: Right-hand side of Equation 5.2 is the tangent of f at \mathbf{w} , where for convex functions, the tangent lower bounds f .

After T iterations, the algorithm returns the averaged weight vector $\bar{\mathbf{w}} = \frac{1}{T} \sum_{t=1}^T \mathbf{w}^{(t)}$. Different versions of the weight could also be returned, such as the last weight vector or the average of the last 25% of changes, but this will be elaborated on in section ???. The scalar parameter $\eta > 0$ will also be discussed later.

The Gradient Descent algorithm requires that the function being minimized is differentiable. Some loss functions are not differentiable, and in these cases, the subgradient of $f(\mathbf{w})$ at $\mathbf{w}^{(t)}$ can be taken instead of the gradient. A vector \mathbf{v} that satisfies Equation 5.2 is called a *subgradient* of f at \mathbf{w} and the set of subgradients is called the *differential set* denoted as $\partial f(\mathbf{w})$. For every $\mathbf{w} \in S$, where S is an open convex set, there exists a \mathbf{v} such that

$$f(\mathbf{u}) \geq f(\mathbf{w}) + \langle \mathbf{u} - \mathbf{w}, \mathbf{v} \rangle, \forall \mathbf{u} \in S. \quad (5.2)$$

For the stochastic gradient descent algorithm, the update direction does not need to be based exactly on the gradient. It is instead allowed to be a random vector, and at each iteration, its *expected value* is required to be a subgradient of the function at the current vector. An example of the difference between the SGD and the GD procedure for minimization of convex function is shown in Fig. 5.2. As stated previously in ??? and in, if a function f is λ -strongly convex for every vector \mathbf{w}, \mathbf{u} , and $\mathbf{v} \in \partial f(\mathbf{w})$ then we have $\langle \mathbf{w} - \mathbf{u}, \mathbf{v} \rangle \geq f(\mathbf{w}) - f(\mathbf{u}) + \frac{\lambda}{2} \|\mathbf{w} - \mathbf{u}\|^2$ implying that the norms will never be greater than λ . It is then shown in that an unbiased estimate of the function's subgradient can be chosen. The update

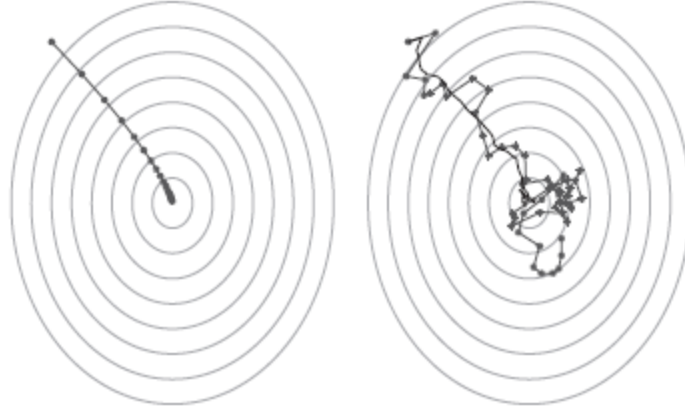


Fig. 5.2.: A plot of the gradient descent algorithm (left) and the stochastic gradient descent algorithm (right). The function being minimized is $1.25(x + 6)^2 + (y - 8)^2$. In the figure representing stochastic gradient descent, the solid line shows the averaged value of \mathbf{w} .

rule for these functions can then be rewritten as 5.3.

$$w^{(t+1)} = -\frac{1}{\lambda t} \sum_{i=1}^t v_i, \quad (5.3)$$

where t is the number of iterations. This shows, by using SGD, the regularized loss function can directly be minimized. It is done by sampling a point i.i.d from a distribution, then using a subgradient of the loss function at this point as an unbiased estimate of the subgradient of the risk function.

5.1.3 Issues with Current Online Learning Algorithms

5.2 OLLAWV: OnLine Learning Algorithm using Worst-Violators

OLLAWV is an iterative, online learning algorithm for solving the L1-SVM problem using a novel model update procedure while implementing a self-stopping condition. The inspiration behind OLLAWV came from [53] which presented a generic online learning algorithm tailored not only for SVMs, but also for various other popular classifiers that use different risk functions, with or without a regularization term. The difference, novelty, and advantage of OLLAWV resides in its iterative method, where the weight α_i of the most violating sample i.e., of the *worst-violator*, is only updated in each iteration. A worst violating sample is defined as the sample that has the largest error with respect to the current decision function. Rather than randomly selecting samples to update per iteration, OLLAWV selects (without replacement) the most incorrectly classified sample and updates the model accordingly. By iteratively updating the model using only the worst-violator, the model is essentially finding its support vectors, as well as implicitly defining a stopping criterion. If there are no more violating samples, the algorithm terminates, eliminating the need to define the number of iterations for an algorithm to perform before returning the model, as is the case with most state-of-the-art online algorithms.

At every iteration, the algorithm selects a worst violating sample that has not been previously chosen, stores its index in vector \mathbf{S} , and then updates the model. Equation 5.4 shows the method for selecting the worst-violator, where $y_o \in \mathbb{R}$ is the error value, $wv \in \{1, \dots, n\}$ is the error value's index, $\mathbf{o} \in \mathbb{R}^n$ is the decision function output, and \neg is the 'not' symbol. For the L1-SVM, an error value will always be negative which is why the minimum function is used (i.e. the most negative output value or incorrectly classified sample). The worst violating sample becomes the model's support vector because its weight is updated and non-zero. Therefore, the number of iterations of OLLAWV is equal to the number of support vectors that the resulting model has. This is an interesting property of OLLAWV; if the number of iterations is set beforehand, one is implicitly setting a bound on the number

of support vectors.

$$[yo, wv] = \min \{y_{wv} \cdot o_{wv}\}, \forall wv \in \{\neg \mathbf{S}\} \quad (5.4)$$

Algorithm 5.1 lists OLLAWV's pseudocode and Figure 5.3 illustrates the steps taken by OLLAWV. First, the model parameters $(\boldsymbol{\alpha}, b, \mathbf{S})$ and the algorithm variables $(\mathbf{o}, \text{iteration counter } (t), \text{initial worst-violator index } wv \text{ and its error } yo)$ are first initialized. The worst-violator with respect to the current hyperplane is then found and the model parameters are updated. Once no more violating samples are found or the maximum number of iterations is reached, the model is returned.

OLLAWV performs stochastic gradient descent on the primal L1-SVM objective function given below:

$$\min_{\mathbf{w} \in \mathcal{H}_o \times \mathbb{R}} R = \frac{1}{2} \|\mathbf{w}\|^2 + C \sum_{i=1}^n \max \{0, 1 - y_i o_{(w)}(\mathbf{x}_i)\}, \quad (5.5)$$

where $o_{(w)}(\mathbf{x}_i) = \langle \mathbf{w}, \mathbf{x}_i \rangle$ is a linear predictor, with the bias term excluded for simplicity. Note that the loss function used in Equation 5.5 is non-differentiable, but it has a subgradient due to a knick-point at $yo = 1$. The loss function's gradient after the knick-point equals zero, which leads to a sparse model. Hence, when the value of $yo \geq 1$ the loss is zero, and for $yo < 1$ the loss increases linearly. The subgradient of the above cost function is given by:

$$\frac{\partial R}{\partial \mathbf{w}} = \begin{cases} \mathbf{w} - C \sum_{i=1}^n y_i \mathbf{x}_i & y_i o_i < 1 \\ \mathbf{w} & \text{otherwise.} \end{cases} \quad (5.6)$$

In the stochastic case, the calculation of the gradient needed for the weight update, is *pattern based*, not *epoch based* as in batch gradient descent. It has been shown [54] that the ideal gradient is equal to the sum of the gradients calculated after each sample is presented for fixed weights during the whole epoch. Thus, the stochastic update of \mathbf{w} from the subgradient

shown in Equation 5.6 becomes:

$$\begin{aligned} \mathbf{w} &\leftarrow \mathbf{w} - \eta \frac{\partial R}{\partial \mathbf{w}} \\ \mathbf{w} &\leftarrow \mathbf{w} + \eta \begin{cases} Cy_i \mathbf{x}_i - \mathbf{w} & y_i o_i < 1 \\ -\mathbf{w} & \text{otherwise,} \end{cases} \end{aligned}$$

where $\eta > 0 \in \mathbb{R}$ is the learning rate. According to the Representer Theorem, a vector $\boldsymbol{\alpha} \in \mathbb{R}^n$ exists such that $\mathbf{w} = \sum_{i=1}^n \alpha_i \phi(\mathbf{x}_i)$ is an optimal solution to Equation 5.5, where $\phi(\cdot)$ is a mapping from feature space to Hilbert space [76]. On the basis of the Representer Theorem, Equation 5.5 can be optimized with respect to $\boldsymbol{\alpha}$ instead of \mathbf{w} . By expressing \mathbf{w} this way and mapping input sample \mathbf{x}_i to $\phi(\mathbf{x}_i)$, the kernelized SGD update becomes:

$$\sum_{i=1}^n \alpha_i \phi(\mathbf{x}_i) \leftarrow \sum_{i=1}^n \alpha_i \phi(\mathbf{x}_i) + \eta \begin{cases} Cy_i \phi(\mathbf{x}_i) - \sum_{i=1}^n \alpha_i \phi(\mathbf{x}_i) & y_i o_i < 1 \\ -\sum_{i=1}^n \alpha_i \phi(\mathbf{x}_i) & \text{otherwise.} \end{cases}$$

However, OLLAWV optimizes in a stochastic manner, resulting in the following update:

$$\begin{aligned} \forall i : \alpha_i \phi(\mathbf{x}_i) &\leftarrow \alpha_i \phi(\mathbf{x}_i) + \eta \begin{cases} (Cy_i \phi(\mathbf{x}_i) - \alpha_i \phi(\mathbf{x}_i)) & y_i o_i < 1 \\ (-\alpha_i \phi(\mathbf{x}_i)) & \text{otherwise} \end{cases} \\ \forall i : \alpha_i &\leftarrow \alpha_i + \eta \begin{cases} (Cy_i - \alpha_i) & y_i o_i < 1 \\ (-\alpha_i) & \text{otherwise.} \end{cases} \end{aligned}$$

The case when the worst violating sample is correctly classified $y_o \geq 1$ is OLLAWV's termination condition, i.e. is used as the stopping criterion in the algorithm. Hence the update for $\boldsymbol{\alpha}$ is reduced to the following:

$$\forall i : \alpha_i \leftarrow \alpha_i + \eta (Cy_i - \alpha_i) \quad (5.7)$$

If the bias term b is included in Equation 5.5, it's stochastic update is as follows:

$$\forall i : b \leftarrow b + \eta \frac{Cy_i}{n} \quad (5.8)$$

In this experimental study, $\eta = 2/\sqrt{t}$ is used, where t is the current iteration; however, other learning rates such as $\eta = 1/t$ can also be used. Let $\Lambda = \eta C y_i$ and $P = \eta \alpha_i$ be the update parameters for OLLAWV, and the α update can be expressed as: $\alpha_i \leftarrow \alpha_i + (\Lambda - P)$. Note that Λ is the update resulting from the loss function and P is derived from the regularizer term in Equation 5.5. In the case of OLLAWV, $P = 0$ because samples are never updated more than once and their initial α value is always 0. It is important to note that in OLLAWV's case, Λ never equals 0 because the samples being updated are worst-violators, meaning they are misclassified or incur some loss. The values of the decision function (output vector $\mathbf{o} \in \mathbb{R}^n$ in Algorithm 5.1), from which a worst-violator is found, changes per iteration based on the influence of the support-vectors that have been previously updated. From Equation 2.7, the output vector update becomes the following:

$$\mathbf{o} \leftarrow \mathbf{o} + \Lambda * \mathcal{K}(\mathbf{x}_{-\mathcal{S}}, \mathbf{x}_{wv}, \gamma) + B \quad (5.9)$$

where $\mathcal{K}(\cdot)$ is the Gaussian radial basis function (RBF) kernel, $\gamma \in \mathbb{R}$ is its parameter, and $B = (\Lambda * \beta)/n$ denotes the bias update. Only non-support vector output values are calculated per iteration, as denoted by $\mathbf{x}_{-\mathcal{S}}$ in the kernel function, because samples are never selected to be updated more than once. Because the output values scale with the C value, the stopping criteria for OLLAWV is also set to scale with C , rather than the classic formulation $y_i o_i \geq 1$. If the value of C is very large, $y_i o_i$ will never be greater than 1 and the algorithm will never terminate. Therefore, the stopping criteria is set to be $y_i o_i \geq M$, where $M \in \mathbb{R}$ is a scaled value of C . For the B calculation in Equation 5.9, $\beta \in \{0, 1\}$ indicates whether the bias term is to be used. If b is not a part of the model, it should be omitted from Equations 2.7 and 5.9 by setting $\beta = 0$, otherwise $\beta = 1$.

OLLAWV is a stochastic gradient method (SGM) that has a convex cost function. Its learning rate coefficient can decrease linearly or semi-linearly during the learning stage. Hence, OLLAWV shares the complexity characteristics of SGM methods. Primarily, it can

Algorithm 5.1 OnLine Learning Algorithm using Worst-Violators (OLLAWV)

Input: $\mathcal{D}, C, \gamma, \beta, M$
Output: α, b, \mathcal{S}

```

1:  $\alpha \leftarrow \mathbf{0}, b \leftarrow 0, \mathcal{S} \leftarrow \mathbf{0}$  ▷ Initialize OLLAWV model parameters
2:  $\mathbf{o} \leftarrow \mathbf{0}, t \leftarrow 0$  ▷ Initialize the output vector and iteration counter
3:  $wv \leftarrow 0, yo \leftarrow y_{wv} * \mathbf{o}_{wv}$  ▷ Initialize hinge loss error and worst-violator index
4: while  $yo < M$  do
5:    $t \leftarrow t + 1$ 
6:    $\eta \leftarrow 2/\sqrt{t}$  ▷ Learning rate
7:
8:    $\Lambda \leftarrow \eta * C * y_{wv}$  ▷ Calculate hinge loss update
9:    $B \leftarrow (\Lambda * \beta) / n$  ▷ Calculate bias update
10:   $\mathbf{o} \leftarrow \mathbf{o} + \Lambda * \mathcal{K}(\mathbf{x}_{-\mathcal{S}}, \mathbf{x}_{wv}, \gamma) + B$  ▷ Update output vector
11:   $\alpha_{wv} \leftarrow \alpha_{wv} + \Lambda$  ▷ Update worst-violator's alpha value
12:   $b \leftarrow b + B$  ▷ Update bias term
13:
14:   $\mathcal{S}_t \leftarrow wv$  ▷ Save index of worst-violator
15:   $[yo, wv] \leftarrow \min_{wv \in \{-\mathcal{S}\}} \{y_{wv} \cdot \mathbf{o}_{wv}\}$  ▷ Find the worst-violator
16: end while

```



Fig. 5.3.: A summary of the steps performed by OLLAWV. The model parameters (α, b, \mathcal{S}) and the algorithm variables $(\mathbf{o}, t, wv, \text{ and } yo)$ are first initialized. The worst-violator with respect to the current hyperplane is then found and the model parameters are then updated. Once no more violating samples are found, the model is returned.



Fig. 5.4.: A case of classifying 2-dimensional normally distributed data with different covariance matrices, (left) for 200 and (right) 2000 data points. The theoretical separation boundary (denoted as the Bayes Separation Boundary) is quadratic and is shown as the dashed black curve. The other two separation boundaries shown are the ones obtained by OLLAWV and SMO (implemented within LIBSVM), respectively. In this particular case (left), the difference between the OLLAWV boundary and the SMO boundary is hardly visible. The case presented on the right shows that, with an increase of training samples, the OLLAWV and SMO boundaries converge to the theoretical Bayesian solution.

achieve linear convergence, making it a particularly convenient and practical method for solving very large machine learning problems. OLLAWV also works over a cost function without local minima, always leading towards the global minimum, even though it stops the learning process as soon as all samples are outside the prescribed margin. Figure 5.4 shows the decision boundaries achieved by OLLAWV versus SMO (implemented within LIBSVM) and Bayes for toy datasets.

5.3 Experimental Environment, Results, and Analysis

This section presents two experimental setups of our contribution against other state-of-the-art algorithms on 23 different benchmark datasets. The first study, presented in Section 5.3.1, compares OLLAWV to two other SVM kernel methods, and the second compares

Table 5.2.: Classification Datasets

Dataset	# Samples	# Attributes	# Classes
<i>small datasets</i>			
iris	150	4	3
teach	151	5	3
wine	178	13	3
cancer	198	32	2
sonar	208	60	2
glass	214	9	6
vote	232	16	2
heart	270	13	2
dermatology	366	33	6
prokaryotic	997	20	3
eukaryotic	2,427	20	4
<i>medium datasets</i>			
optdigits	5,620	64	10
satimage	6,435	36	6
usps	9,298	256	10
pendigits	10,992	16	10
reuters	11,069	8,315	2
letter	20,000	16	26
<i>large datasets</i>			
adult	48,842	123	2
w3a	49,749	300	2
shuttle	58,000	7	7
web (w8a)	64,700	300	2
ijcnn1	141,691	22	2
intrusion	5,209,460	127	2

OLLAWV to 5 non-SVM methods, shown in Section 5.3.3. In each section, the experimental setups are first described and the state-of-the-art methods are listed. The results and statistical analysis are then presented and analyzed. The main aim of the experiments is to compare our contribution to other support vector machine solvers that have been shown to surpass popular and widely used SVM kernel methods in terms of memory consumption, runtime, and accuracy. The supplemental experimental study in 5.3.3 was conducted to emphasize the better performance of OLLAWV against non-SVM algorithms.

Table 5.2 presents a summary of the 23 datasets used throughout the experiments, where the number of attributes (dimensionality), classes, and samples are shown. The datasets used and the results obtained are divided into three groups: *small*, *medium* and *large*. The

datasets were obtained from the UCI Machine Learning repository¹, and the LIBSVM², and the LibCVM³ sites [5, 24, 84].

5.3.1 SVM Experimental Setup

The experimental setup was designed to evaluate differences in performance of the proposed OLLAWV method against the state-of-the-art algorithms: *Minimal Norm SVM* (MNSVM) [81] and *Non-Negative Iterative Single Data Algorithm* (NNISDA) [102]. These algorithms were chosen because they have shown considerable performance in runtime, memory consumption, and accuracy against the popular and widely used LIBSVM and LibCVM packages. In [81], it was shown that MNSVM outperforms both the L1 and L2 implementations of LIBSVM, and BVM embedded in LibCVM. NNISDA was then compared to MNSVM in [102], and showed an added improvement in runtime performance. MNSVM was implemented in an open source C++ framework called “GSVM – Command Line Tool for Geometric SVM Training⁴”. Both, NNISDA and OLLAWV were implemented as additional modules within Strack-Kecman’s code, keeping the experimental environment controlled for all three algorithms. The experiments for all methods were run on the same machine containing two Intel Xeon X5680 CPUs (6-core, 3.33 GHz) and 96 GB of RAM.

Experiments were performed using double, or nested, 5-fold cross-validation in order to objectively evaluate the models’ performances and tune hyper-parameters. In the outer loop, the data are separated into 5 equally sized folds and each part is held out in turn as the test set, and the remaining four parts are used as the training set. In the inner loop, 5-fold cross-validation is also used over the training set assigned by the outer loop, where the best hyper-parameters are chosen. The best model obtained by the inner loop is then applied on the outer loop’s test set. This procedure ensures the model’s performance is not

¹<http://archive.ics.uci.edu/ml/index.php>

²<https://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/>

³<http://c2inet.sce.ntu.edu.sg/ivor/cvm.html>

⁴<https://github.com/strackr/gsvm>

optimistically biased as when using a single loop of k -fold cross-validation. It ensures the class labels of the test data will not be seen when tuning the hyper-parameters, which is consistent with real-world applications. Obviously, such a rigorous procedure is computationally expensive, but the goal is to fairly compare different classification models on the same data sets, with the same cross-validation procedure, and hyper-parameters. First, the datasets were normalized by linear transformation of the feature values to the range $[0, 1]$. Then, the training process, also involving model selection using pattern search, was performed. The best hyper-parameters were chosen from the following 8×8 possible combinations, shown in Equations (5.10a) and (5.10b), and were also used for the competing SVM methods.

$$C \in \{4^n\}, \quad n = \{-2, \dots, 5\} \quad (5.10a)$$

$$\gamma \in \{4^n\}, \quad n = \{-5, \dots, 2\} \quad (5.10b)$$

The γ parameter refers to that of the Gaussian RBF kernel, given by:

$$\mathcal{K}(\mathbf{x}_i, \mathbf{x}_j) = e^{-\gamma \|\mathbf{x}_i - \mathbf{x}_j\|^2}. \quad (5.11)$$

To deal with multi-class classification problems, the one-vs-one, or pairwise, approach was used. The pairwise training procedure trains $c(c-1)/2$ binary classifiers, a classifier for each possible pair of classes, where c is the number of classes. During the prediction phase, a voting scheme is used where all $c(c-1)/2$ models predict an unseen data sample and the class that received the highest number of votes is considered to be the samples true class.

5.3.2 SVM Comparison Results and Statistical Analysis

The classification performance was measured using the following metrics: accuracy, runtime, and the percentage of support vectors (size of the model). Table 5.3 displays the results for OLLAWV and the two state-of-the-art methods. The percentage of support vectors was reported for analyzing the complexities of the resulting models over the variously sized datasets. In order to analyze the performances of the multiple models, non-parametric

Table 5.3.: Comparison of OLLAWV vs. NNISDA and MNSVM

Dataset	Accuracy (%)			Runtime (s)			Support Vectors (%)		
	OLLAWV	NNISDA	MNSVM	OLLAWV	NNISDA	MNSVM	OLLAWV	NNISDA	MNSVM
<i>small datasets</i>									
iris	97.33	94.00	96.67	0.05	0.27	3.57	13.50	40.20	29.80
teach	52.32	52.31	52.95	0.12	0.44	8.85	69.19	99.80	87.40
wine	98.87	96.60	96.60	0.28	0.43	4.84	15.02	44.40	48.60
cancer	80.36	81.86	81.38	0.49	0.85	4.46	42.79	83.80	89.60
sonar	92.32	89.48	87.57	0.59	0.98	3.03	31.26	73.00	66.00
glass	72.41	67.81	69.30	0.46	1.01	11.94	62.84	90.80	87.60
vote	96.54	96.11	93.99	0.26	0.46	1.49	13.36	33.20	34.00
heart	82.22	83.33	83.33	0.50	0.91	6.45	37.69	73.00	82.00
dermatology	97.82	98.36	98.36	1.62	2.47	11.68	36.94	59.00	59.80
prokaryotic	88.96	88.86	88.97	6.09	10.64	50.86	29.01	51.20	49.00
eukaryotic	77.38	79.56	81.21	61.95	49.16	342.76	54.11	76.40	72.60
<i>medium datasets</i>									
optdigits	99.11	99.29	99.31	411	528	787	28.64	31.60	30.60
satimage	91.66	92.39	92.35	1,334	687	1,094	20.72	45.00	44.80
usps	97.49	98.05	98.24	10,214	5,245	7,777	11.22	29.40	28.00
pendigits	99.56	99.62	99.61	723	909	1,500	10.27	17.60	16.60
reuters	98.03	98.08	97.99	954	1,368	1,657	8.770	18.20	18.60
letter	96.99	99.11	99.13	5,259	12,009	26,551	43.56	57.60	56.60
<i>large datasets</i>									
adult	84.75	85.07	85.13	21,025	72,552	123,067	34.66	56.00	56.60
w3a	98.86	98.82	98.82	6,532	15,951	24,562	3.270	14.60	12.40
shuttle	99.77	99.83	99.87	2,833	7,420	45,062	2.010	6.00	16.40
web	98.94	99.00	99.00	12,067	30,583	38,040	4.320	13.20	10.80
ijcnn1	98.31	99.34	99.41	162,587	296,917	370,144	16.36	11.00	7.600
intrusion	99.77	99.67	99.66	2,402,804	4,646,810	3,772,113	0.780	2.000	1.700
Average	91.29	91.15	91.25	114,209	221,350	191,861	25.66	44.65	43.79
Ranks	1.739	2.022	2.239	1.217	1.913	2.869	1.087	2.609	2.304

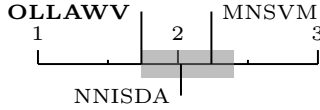


Fig. 5.5.: Bonferroni-Dunn test for Accuracy

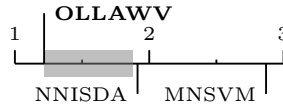


Fig. 5.6.: Bonferroni-Dunn test for Runtime

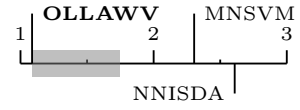


Fig. 5.7.: Bonferroni-Dunn test for % Support Vectors

statistical tests are used to validate the experimental results obtained [31]. The Iman-Davenport non-parametric test is run to investigate whether significant differences exist among the performance of the algorithms by ranking them over the datasets used, using the Friedman test. The algorithm ranks for each metric are presented in the last row of Table 5.3, and the lowest (best) rank value is typeset in bold. After the Iman-Davenport test indicates significant differences (with p -value = 0.2397 for accuracy, and p -value = 0 for runtime and percent support vectors), the Bonferroni-Dunn post-hoc test is then used to find where they occur between algorithms by assuming the classifiers' performances are different by at least some critical value (critical distance is 0.66 for $\alpha = 0.05$). Below Table 5.3, Figures 5.5, 5.6,



Fig. 5.8.: Runtime in seconds versus the number of samples, divided into two groups: small & medium (left) versus large (right). Note OLLAWV’s gradual increase in runtime as the number of samples increases compared to NNISDA and MNSVM’s steeper change. In almost all cases, OLLAWV displays superior runtime over state-of-the-art. Runtime depends upon many characteristics: dimensionality, class-overlapping, complexity of the separation boundary, number of classes as well as upon the number of support vectors, which partly explains the tiny bump in the left figure.

and 5.7 highlight the critical distance (in gray) from the best ranking algorithm to the rest. The algorithms to the right of the critical distance bar perform statistically significantly worse than the control algorithm, OLLAWV.

The results in Table 5.3 indicate that OLLAWV outperforms NNISDA and MNSVM in terms of accuracy, runtime, and model complexity. Although the differences in accuracy between the methods is not very large, on average, OLLAWV is about 2 times faster than NNISDA and MNSVM. As mentioned previously, OLLAWV aims to speed up the learning process without sacrificing the model’s accuracy. This stems from OLLAWV’s ability to produce sparse models, as is shown by the averaged percentage of support vectors. The speedup that OLLAWV presents is proportional to the model complexity and the experimental results show that OLLAWV produces, on average, models that are 1.7 times smaller

than the two state-of-the-art methods used. This highlights the applicability and advantage that OLLAWV has for learning from large datasets.

Figures 4.4, 5.6, and 5.7 show the results of the statistical analysis for accuracy, runtime, and percentage of support vectors. Figures 5.6 and 5.7 show that OLLAWV is statistically significantly better than MNSVM and NNISDA for runtime and percentage of support vectors (model size). At the same time, Figure 4.4 emphasizes what was mentioned earlier: OLLAWV is shown to speed up the learning process without sacrificing model accuracy against the state-of-the-art methods used.

Figure 5.8 plots the correlation of OLLAWV, NNISDA, and MNSVM’s runtime versus number of samples for the small, medium, and large datasets. The figure clearly emphasizes the benefit of using OLLAWV for large-scale learning due to its gradual increase in runtime as the number of samples increases in comparison to NNISDA and MNSVM. Figure 5.9 shows the correlation between OLLAWV, NNISDA, and MNSVM’s percentage of support vectors and the number of samples for all datasets. It highlights OLLAWV’s model sparseness in comparison to the competing methods, while mirroring the runtime results.

5.3.3 Non-SVM Experimental Setup

The supplemental experimental setup was designed to compare the performance of the proposed OLLAWV against the following 5 popular and widely-used non-SVM algorithms: *k-Nearest Neighbors (k-NN)*, *J48*, *JRip*, *Naïve Bayes*, and *Logistic*. These methods have been implemented within the Weka framework [37]. The experiments were performed under the same nested 5-fold cross-validation framework as the SVM algorithm experimental study which was described in Section 5.3.1. The following hyper-parameters shown in Table 5.4 were used for the non-SVM algorithms.

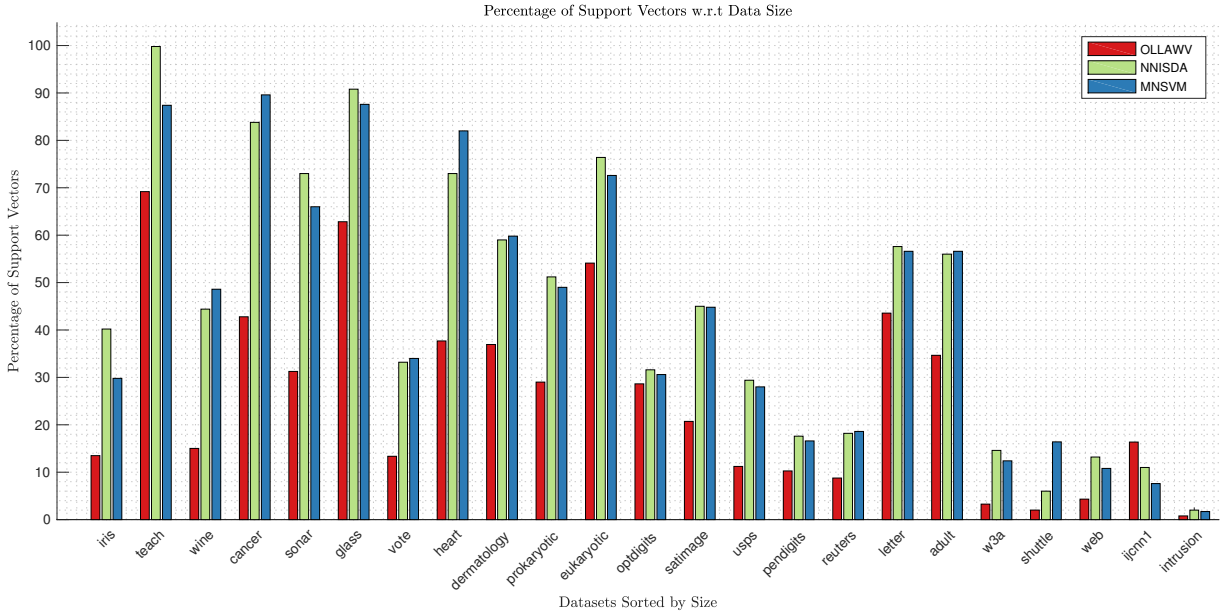


Fig. 5.9.: Size of the model given as percentage of support vectors with respect to the number of samples versus the number of samples. Note that OLLAWVs percentage of support vectors is always smaller (except in one case) than NNISDA’s and MNSVM’s ones.

Table 5.4.: Non-SVM Algorithm Hyper-parameters

Algorithm	Parameters
<i>k</i> -NN	Number of neighbors: $k \in \{1, 3, 5, 7\}$
J48	Pruning: $\{\text{True}, \text{False}\}$, Pruning Confidence: $\{0.1, 0.25, 0.5\}$
JRip	Pruning: $\{\text{True}, \text{False}\}$
Naive Bayes	Use kernel estimation: $\{\text{True}, \text{False}\}$
Logistic	Log-likelihood: $\{1e^{-7}, 1e^{-8}, 1e^{-9}\}$

5.3.4 Non-SVM Results and Statistical Analysis

Table 5.5 displays the accuracy results for OLLAWV and five state-of-the-art methods. The table also shows the standard deviation for accuracy per outer fold, the average values accross all datasets, and the algorithm ranks. As the results indicate, OLLAWV outperforms all other state-of-the-art methods. Figure 5.11 displays the average accuracy results for OLLAWV and the non-SVM methods accross all datasets and highlights OLLAWV’s better

Table 5.5.: Accuracy (%) Comparison for Non-SVM Methods vs. OLLAWV

Dataset	OLLAWV	k-NN	J48	JRip	Naïve Bayes	Logistic
<i>small datasets</i>						
iris	97.33 ± 1.49	96.00 ± 3.65	94.00 ± 2.79	90.67 ± 4.35	96.00 ± 2.79	97.33 ± 2.79
teach	52.32 ± 3.46	59.64 ± 2.89	49.72 ± 7.58	56.75 ± 9.60	53.75 ± 6.46	51.77 ± 6.68
wine	98.87 ± 1.54	97.73 ± 3.72	90.43 ± 5.83	93.24 ± 3.27	96.60 ± 3.14	96.05 ± 2.58
cancer	80.36 ± 5.80	77.32 ± 6.93	73.81 ± 8.57	73.78 ± 5.81	67.73 ± 5.07	77.32 ± 7.78
sonar	92.32 ± 3.11	88.99 ± 4.59	76.16 ± 10.6	75.18 ± 6.77	73.69 ± 7.65	75.18 ± 7.31
glass	72.41 ± 2.28	67.73 ± 5.91	65.06 ± 5.51	65.59 ± 9.66	49.46 ± 5.19	62.04 ± 5.75
vote	96.54 ± 1.87	92.26 ± 3.19	95.70 ± 2.12	96.54 ± 2.45	92.24 ± 3.24	93.54 ± 2.59
heart2	82.22 ± 2.93	79.63 ± 5.71	78.52 ± 2.81	80.74 ± 4.06	84.44 ± 4.46	83.33 ± 3.93
dermatology	97.82 ± 0.05	96.18 ± 1.78	94.52 ± 2.21	91.27 ± 5.08	97.28 ± 1.64	96.98 ± 2.28
prokaryotic	88.96 ± 2.14	87.96 ± 3.01	78.54 ± 1.62	79.13 ± 2.78	62.38 ± 3.54	87.57 ± 2.56
eukaryotic	77.38 ± 1.96	81.42 ± 2.06	65.27 ± 2.92	66.42 ± 3.47	39.27 ± 3.43	69.55 ± 1.34
<i>medium datasets</i>						
optdigits	99.11 ± 0.38	98.74 ± 0.39	90.87 ± 1.09	91.28 ± 0.40	92.42 ± 0.75	95.05 ± 0.91
satimage	91.66 ± 0.80	90.38 ± 0.72	85.64 ± 1.21	85.33 ± 0.77	85.41 ± 0.92	88.14 ± 1.11
usps	97.49 ± 0.22	97.04 ± 0.47	88.73 ± 0.46	89.20 ± 1.00	79.45 ± 0.59	91.88 ± 0.65
pendigits	99.56 ± 0.12	99.33 ± 0.17	96.24 ± 0.31	96.34 ± 0.41	88.34 ± 0.65	95.59 ± 0.18
reuters	98.03 ± 0.22	97.15 ± 0.43	96.90 ± 0.32	97.18 ± 0.44	93.52 ± 0.02	69.54 ± 0.28
letter	96.99 ± 0.21	95.71 ± 0.19	87.34 ± 0.68	87.02 ± 0.66	74.12 ± 0.97	77.45 ± 0.16
<i>large datasets</i>						
adult	84.75 ± 0.26	83.85 ± 0.28	84.38 ± 0.28	83.73 ± 0.17	80.57 ± 0.09	82.46 ± 0.14
w3a	98.86 ± 0.04	98.60 ± 0.06	98.71 ± 0.05	98.41 ± 0.10	96.71 ± 0.20	98.61 ± 0.12
shuttle	99.77 ± 0.03	99.93 ± 0.03	99.97 ± 0.02	99.96 ± 0.02	98.57 ± 0.24	96.83 ± 0.12
web	98.94 ± 0.05	98.89 ± 0.06	98.79 ± 0.09	98.50 ± 0.13	96.71 ± 0.21	98.70 ± 0.08
ijcnn1	98.31 ± 0.07	98.48 ± 0.04	98.40 ± 0.09	98.11 ± 0.10	90.69 ± 0.26	92.29 ± 0.16
intrusion	99.77 ± 0.02	88.20 ± 1.06	58.01 ± 26.6	87.66 ± 3.79	49.75 ± 30.7	65.15 ± 15.7
Average	91.29 ± 1.26	90.05 ± 2.06	84.60 ± 3.64	86.18 ± 2.84	79.96 ± 3.58	84.45 ± 2.83
Ranks	1.500	2.500	4.041	3.9583	5.0625	3.9375



Fig. 5.10.: Bonferroni-Dunn test for Accuracy

performance. The Friedman test indicates that OLLAWV performs significantly better than the competing methods for $\alpha = 0.05$ and is ranked first. Figure 5.10 shows the critical distance bar (which is 1.391), and indicates that all other algorithms perform statistically significantly worse than OLLAWV, except for *k*-NN.

5.4 Conclusions

This paper proposed a novel online learning procedure and algorithm for solving the L1-SVM problem, which is a unique method in terms of both iterating over samples and

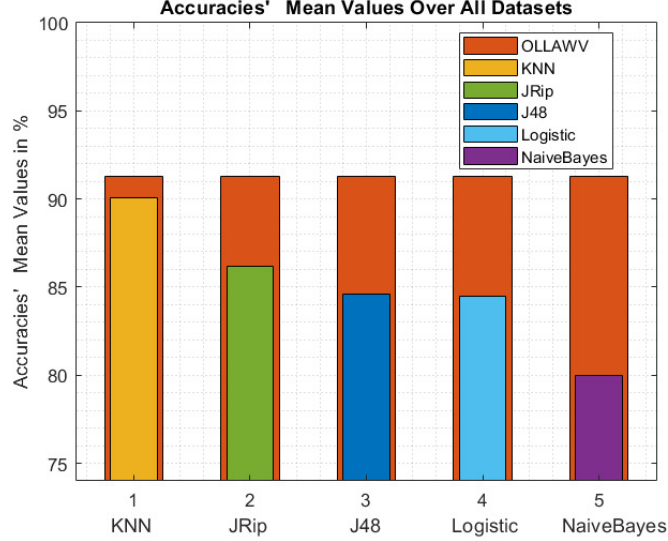


Fig. 5.11.: Mean accuracy over all datasets for OLLAWV and the 5 non-SVM state-of-the-art methods.

updating the model. A new stopping criterion for the stochastic gradient procedure is also proposed. The model is updated by changing the weight α_i of a single worst-violator per iteration and stops when all violating samples i.e., support vectors, are found. Finding the *worst-violators* is done without replacement. Such an approach results in a significant shortening of training time, as well as in a huge decrease in the resulting model size. The key features of the proposed algorithm, OLLAWV, stem from it's implicit ability of finding support vectors and it's self-stopping condition. This design was devised to address the limitations presented by current SVM solvers.

The first experimental study demonstrates the better performance of OLLAWV compared with state-of-the-art SVM solvers (MNSVM and NNISDA) which have been shown to outperform the popular SMO implementation in the LIBSVM package. The results for accuracy, runtime, and percentage of support-vectors, obtained by the strict nested cross-validation procedure, were compared and further validated using statistical analysis with non-parametric tests. They highlighted the advantages and major speedup achieved by OLLAWV against the competing MNSVM and NNISDA. The second, supplemental experi-

mental study evaluated the performance of OLLAWV against 5 popular non-SVM methods, showing the better performance of OLLAWV against all five non-SVM algorithms (k -Nearest Neighbors (k -NN), J48, JRip, Naïve Bayes, and Logistic). The proposal, OLLAWV, performs statistically better in terms of runtime and model size across all 23 evaluated benchmark datasets, without compromising accuracy.

CHAPTER 6

OLLAWV FOR BATCHED DATA STREAMS

- 6.1 Background
- 6.2 DSOLLAWV
- 6.3 Experimental Environment
- 6.4 Results & Analysis
- 6.5 Conclusions

CHAPTER 7

CONCLUSIONS

This thesis introduced four novel SVM algorithms for learning from the non-standard and diverse learning paradigms: multi-target regression and multi-instance classification.

Three unique approaches for multi-target regression were proposed: the baseline problem transformation support vector regressor SVR, an ensemble of randomly generated chains using this base model SVRRC, and a maximally correlated chained model SVRCC. The results highlighted the better performance of SVR as a base model, however, because it is a problem transformation method, the possible correlations amongst the targets are lost on the final model. SVRRC was designed to test whether taking these correlations into account would benefit the final learning model, and the results showed a performance increase. However, due to the random nature of SVRRC, capturing target correlations is not guaranteed. SVRCC was designed to remedy this issue using a maximum correlation chain and proved to capture target correlations accurately, providing the best results among the contributions, as well as against the methods compared.

A novel multiple-instance bag-level formulation and algorithm, dubbed MIRSVM, with a bag-representative selector, are proposed. The algorithm trains the SVM on bag-level information, iteratively selecting the best representative instance for each positive and negative bag, while finding the optimal separating hyperplane. This approach, unlike other existing ones, eliminates possible class imbalance issues by allowing both positive and negative bags to be represented. The experimental and statistical study showed that bag-level learners outperform instance-level learners and wrapper methods. MIRSVM outperformed current contemporary SVM multi-instance algorithms, as well as other algorithms of different classes.

CHAPTER 8

FUTURE WORK

1. Extend OLLAWV to the regression case
2. Apply OLLAWVR to Multi-Target regression problems
3. Use OLLAWV within MIRSVM
4. Experiment with a parallelized/distributed implementation of OLLAWV
5. Experiment with a non-batch version of OLLAWV for Data Streams

The contributions introduced in this work highlight the performance quality of SVMs for solving difficult learning problems. The SVM solvers used were LIBSVM's SMO implementation and MATLAB's standard QP solver. Even though the proposals outperformed existing popular algorithms in each learning paradigm, these two popular and standard solvers have some disadvantages. Although SMO provides an exact solution to the SVM QP problem, its performance is highly dependent on the SVM hyperparameters and MATLAB's QP solver is notoriously dependent on the size and complexity of the data. To address these problems, the future work of this dissertation first aims to deal with these limitations without sacrificing accuracy for scalability with respect to large-scale problems, while maintaining applicability to small and medium sized datasets.

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