Category

Multiple instance learning of Calmodulin binding sites

Fayyaz ul Amir Afsar Minhas^{*} and Asa Ben-Hur^{*}

Department of Computer Science, Colorado State University, Fort Collins, CO 80523-1873, USA

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2 ABSTRACT

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Motivation: Calmodulin (CaM) is a ubiquitously conserved protein 46 4 that acts as a calcium sensor, and interacts with a large number of 47 5 proteins. Detection of CaM binding proteins and their interaction 48 6 sites experimentally requires a significant effort, so accurate meth49 7 ods for their prediction are important. 50

8 **Results:** We present a novel algorithm (MI-1 SVM) for binding site51 9 prediction and evaluate its performance on a set of CaM-bindin 52 10 proteins extracted from the Calmodulin Target Database. Our ap53

proach directly models the problem of binding site prediction as a54 11 12 large-margin classification problem, and is able to take into accoun 55

13 uncertainty in binding site location. We show that the proposed algo56

14 rithm performs better than the standard SVM formulation, and illus57

15 trate its ability to recover known CaM binding motifs. A highly accu58

16 rate cascaded classification approach using the proposed binding 59

17 site prediction method to predict CaM binding proteins in Arabidop60

18 sis thaliana is also presented.

19 Availability: Matlab code for training MI-1 SVM and the cascaded 62 63

20 classification approach is available on request.

21 Contact: fayyazafsar@gmail.com; asa@cs.colostate.edu

22 1 INTRODUCTION

23 Calmodulin (CaM) is an intracellular calcium sensor protein tha68 24 interacts with a large number of proteins to regulate their biologi69 25 cal functions and exhibits sequence conservation across all eukary-70 26 otes (Bouche, Yellin, Snedden, & Fromm, 2005). Ca²⁺ plays a very71 27 important role in many cellular functions ranging from fertilization72 28 and cellular division to neuronal spiking (Reddy, Ben-Hur, & Day73 29 2011). Due to the importance of calcium signaling in cells, identi74 30 fying proteins that bind CaM and determining the location of the75 31 CaM binding site in them can help in gaining a better understand 76 32 ing of cellular function in general, and the role of calcium in dif77 33 ferent cellular processes in particular. This paper presents a highly78 34 accurate computational approach that can identify the location of a7935 CaM binding site in a protein solely on the basis of its amino acid80 36 sequence, helping avoid the significant effort of performing such81 37 experiments in the lab (Reddy, Ben-Hur, & Day, 2011). Our ap82 38 proach uses sequence information alone, which ensures its wide 83 39 applicability in comparison to methods that rely on structural mod84 40 eling (Zhou & Oin, 2007).

- 41 CaM binding sites are known to be contiguous in sequence, often occurring through an amphiphilic alpha helix (O'Neil &8542
- 43 DeGrado, 1990). This makes CaM binding site prediction amena-
- ble to a sliding-window classification approach, as applied in re86 2.1 44

cent work (Radivojac, Vucetic, O'Connor, Uversky, Obradovic, & Dunker, 2006), (Hamilton, Reddy, & Ben-Hur, 2011). The method by Radivojac et al. uses a hierarchical neural network classifier trained on the basis of amino acid properties averaged over a fixedsize window. Hamilton et al. showed that a simple sliding window Support Vector Machine (SVM) trained on average amino acid composition achieves similar performance.

In this paper we present a novel formulation of the binding site prediction problem that is based on the framework of multiple instance learning (MIL) (Dietterich, Lathrop, & Lozano-Perez, 1997). In MIL positive examples come in bags. For a positive bag, it is assumed that at least one of the examples is indeed positive whereas negative bags contain only negative examples. We use this for binding site prediction by forming a positive bag out of fixed-size sequence windows that overlap the annotated binding site. This allows us to model the uncertainty in actual binding site location-experimental methods may not precisely locate a binding site, and may include a region that is larger than the true binding site due to limitations of budget and experimental procedures. Furthermore, modeling binding sites this way facilitates the use of sequence representations that are position dependent, yielding a more detailed model of the binding site. This allows learning of motifs that are characteristic of the binding site.

MIL has been applied in a variety of other problem domains such as object tracking (Babenko, Yang, & Belongie, 2011), protein identification (Tao, Scott, Vinodchandran, & Osugi, 2004), and prediction of protein-ligand binding affinities (Teramoto & Kashima, 2010).

Our results show that the proposed MI-1 SVM has higher accuracy than the classical multiple instance SVM (Andrews, Tsochantaridis, & Hofmann, 2003), and is also faster to train. MI-1 also performs better than a standard SVM, thereby improving on existing work of (Radivojac, Vucetic, O'Connor, Uversky, Obradovic, & Dunker, 2006) and (Hamilton, Reddy, & Ben-Hur, 2011). We also compare the merits of several ways of representing binding sites, and demonstrate the ability of our method to learn motifs that are associated with CaM binding. Finally, we show how the resulting binding site predictor can be used as the basis for a classifier that predicts CaM binding proteins, with improved accuracy over earlier work.

2 **METHODS**

Data Sets and Pre-processing

The data set for CaM binding site prediction and its pre-processing 2 follows (Radivojac, Vucetic, O'Connor, Uversky, Obradovic, & 3 Dunker, 2006). A set of 210 proteins was obtained from the Cal-4 modulin Target database (Yap, Kim, Truong, Sherman, Yuan, & 5 Ikura, 2000). Each of these proteins bind CaM, and one or more 6 binding sites within each protein are annotated. A non-redundant 7 subset of 153 proteins containing 185 binding sites was then cho-8

sen such that no two proteins have more than 40% sequence identi-9 ty and no two binding sites are more than 50% identical.

10 Sequence windows of length 21, the average length of CaM 50 11 binding sites, were extracted from the protein sequences to create 51 12 positive and negative examples. Negative examples were created by sliding a length 21 window in 10 amino acid increments such 52 13 that no part of the window overlaps an annotated binding site53 14 15 Positive examples, on the other hand, were created by sliding 5416 length 21 window over an annotated binding site in increments of 5 17 1 amino acid. Thus, the number of positive examples from an an56 18 notated binding site equals the number of amino acids in the bind57 19 ing site. 58

20 For CaM binding prediction, we used a data set of 236 protein **59** 21 experimentally determined to bind CaM using a protein array60 22 screen that tested around a thousand proteins in Arabidopsis thali61 ana (Popescu, 2007). The remaining 27,140 proteins in the Ara62 23 bidopsis thaliana proteome were used as negative examples (non63 24 25 binders). 64

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26 2.2 Vanilla SVM

As a baseline method we have used a standard binary SVM (Cortes⁶⁷ 27 & Vapnik, 1995). Our labeled dataset consists of N labeled exam $\frac{68}{22}$ 28 ples (x_i, y_i) , where x_i is the sequence of a window, and $y_i \in \{+1, -1\}$ is its associated label indicating whether the central 7129 30 31 residue of x_i lies in a binding site or not. 72

32 The large-margin learning problem can be formulated as:

$$\min_{w,\rho,\xi\geq 0} \frac{1}{2} w^T w + C \sum_{i=1}^{N} \xi_i$$
subject to: $v_i (w^T \phi(x_i) + \rho) \ge 1 - \xi_i, \forall i.$

$$(1)$$

33 Here $\phi(x_i)$ is the feature representation of the window x_i and the 34 cost parameter C controls the trade-off between constraint viola-35 tion and margin maximization. The discriminant function $f(x_i) =$ 36 $w^T \phi(x_i) + \rho$ can then be used to predict whether a given window is part of a binding site or not. The location of a binding site is73 37 predicted by the window that offers the highest value of the dis_{74} 38 39 criminant function for that protein (Hamilton, Reddy, & Ben-Hur75 40 2011). PyML (Ben-Hur, PyML - machine learning in Python76 41 2011) was used for the implementation. 77 78 79

42 2.3 Multiple Instance Learning SVM (mi-SVM)

43 In MIL (both mi-SVM and MI-1-SVM), the positive examples from each binding site are grouped into a single bag. We denote 44 the set of positive examples for a given binding site b as P(b) and $added{delta}$ 45 46 the set of negative examples from the protein to which the binding

- site b belongs as N(b). The mi-SVM approach is formulated a \$547
- 48 follows (Andrews, Tsochantaridis, & Hofmann, 2003):

$$\min_{y \in \{-1,+1\}^N} \left(\min_{w,\rho,\xi \ge 0} \frac{1}{2} w^T w + \frac{C}{N} \sum_{i=1}^N \xi_i \right)$$
subject to: $y_i (w^T \phi(x_i) + \rho) \ge 1 - \xi_i, \forall i$

$$\sum_{i \in P(b)} \frac{y_i + 1}{2} \ge 1, \quad \forall b$$

$$y_i = -1, \quad \forall i \in N(b), \forall b.$$

$$(2)$$

In this formulation $y_i \in \{+1, -1\}$ acts as a label for the window x_i , and the objective is to find the optimal labeling of the examples that comprise the positive bags such that at least one example in each positive bag is labeled as positive $(\sum_{i \in P(b)} (y_i + 1)/2 \ge 1)$. The other constraints ensure correct labeling of the given training examples and that all negative examples are labeled as negative examples. In case of the binding site prediction problem, this means that a trained mi-SVM will choose at least one positive window from the set of positive windows in a binding site. The mi-SVM formulation is a combinatorial optimization problem. We use the heuristic algorithm proposed by (Andrews, Tsochantaridis, & Hofmann, 2003) to solve this problem. The algorithm initially assigns the label of a bag to all examples in it, i.e., all examples in positive bags are assigned a label of +1 whereas all negative examples are assigned -1. It uses these assigned labels to solve a regular SVM learning problem (as in Equation (1)). Labels for all examples in positive bags are then imputed based upon the sign of their discriminant function value. If no example in a positive bag is assigned a positive label (i.e. the constraint $\sum_{i \in P(b)} (y_i + 1)/2 \ge 1$ is violated), the algorithm picks the example in the bag having the largest discriminant function value and sets its label to +1. The algorithm then alternates between label imputation and SVM training until the labels stop changing. This simple algorithm has shown good performance in comparison to more complicated ones



(Andrews, Tsochantaridis, & Hofmann, 2003).

Fig. 1. CaM binding site prediction with MIL. The annotated binding site is shown as a box, and is represented by a "bag" composed of the windows indicated in red above the binding site. The rest of the windows that do not overlap the binding site are negative examples (shown in blue below the protein). The bottom panel illustrates the desired characteristics of the classifier's discriminant function. The dots indicate the score of different examples (positive indicated by solid red circles and negative shown as hollowed blue circles). The score from the trained discriminant function for one window in a binding site should be higher than the scores generated for non-binding site windows within that protein.

2.4 Novel MI SVM Formulation (MI-1 SVM)

86 Accurate prediction of the location of a binding site in a protein requires a less stringent condition than the one used in mi-SVM: at least one window in the true binding site needs to score higher than the negative windows from the same protein (see Fig. 1). This allows us to significantly reduce the complexity of the learning38
 problem in comparison to mi-SVM. The mi-SVM and vanilla39

3 SVM formulations try to classify windows as binding or non40

4 binding without modeling the concept that these windows in fac41

5 lie within a protein. Our proposed MI-1 SVM formulation, on the 42

6 other hand, operates at the protein level. The large-margin formula 43

7 tion of this learning problem, can be expressed as follows:

 $\min_{w,\xi \ge 0} \frac{1}{2} w^T w + \frac{c}{M} \sum_{b}^{b} \xi_{b}$ such that $\forall b$ $\max_{i \in P(b)} w^T \phi(x_i) \ge w^T \phi(x_i) + 1 - \xi_{b}, \forall j \in N(b),$ (3)

8 where, M is the total number of binding sites in the training data.

For a given binding site, this formulation tries to maximize the 449 10 difference between the discriminant function values of the maximum scoring window within the binding site and the non-binding 4511 windows in the rest of the protein containing that binding site.46 12 Since MI-1 SVM simply compares the discriminant function⁴⁷ 13 scores in the binding and non-binding site windows in its $con-\frac{48}{3}$ 14 scores in the binding and non-binding site windows in its con-19 straints, it does not require a bias term. Moreover, the number of $\frac{49}{50}$ slack variables (ξ_b) in MI-1 SVM is equal to the number of bind $\frac{50}{52}$ syvM and the number of training examples, as in the vanilla SVM and the mi-SVM. As a consequence, the number of variables involved in the optimization in ML1 SVM is much smaller than 15 16 17 18 involved in the optimization in MI-1 SVM is much smaller that 5419 that in mi-SVM and this leads to faster training. Using the same ξ_{b55} 20 21 for a single binding site effectively takes the maximum of the

22 scores over all non-binding site windows of the protein to which b

23 belongs. Another important feature of MI-1 SVM is that, like the 56

24 ranking SVM discussed in (Joachims, 2006), MI-1 SVM also ex57

25 plicitly maximizes the area under the Receiver Operating Charac $_{58}$

26 teristics (ROC) curve.

27	Table 1. Heuristic algorithm used for training MI-1	<u> 59</u>
	Initialization	60
	With each binding site b , we associate a representative example x^b w	ith61
	feature representation $\phi(x^b)$ which is initialized to be the mean of	h = 62
	examples in $P(b)$:	63
	$f(w) = \frac{1}{2} \sum_{i=1}^{n} f(w) \times h$	64
	$\varphi(x^{-}) = \frac{\varphi(x_{i})}{ P(b) } \sum_{i=1}^{n} \varphi(x_{i}), \forall b$	- 65
	$i \in P(b)$	66
	Until Convergence, repeat:	
	Solve the following quadratic programming (QP) problem:	6/
	$\min_{w \in \mathcal{A}} \frac{1}{w^T w} + \frac{c}{c} \sum \xi$	- 68
	$MM_{w,\xi \geq 0} 2^{w + w + M} \Delta_{\mu}^{\zeta b}$	69
	such that, $\forall b$	70
	$w^T \phi(x^b) \ge w^T \phi(x_j) + 1 - \xi_b, \forall j \in N(b)$	71
	Update (for all binding sites):	72
	$\phi(x^b) = \phi(x_i)$ such that $i = \operatorname{argmax}_{j \in P(b)} w^T \phi(x_j)$	/3

Similar to mi-SVM, which performs optimization over the labels75 28 of examples in positive bags, MI-1 SVM is also a combinatoriab 29 30 optimization problem because of the maximum operation in its77 31 constraints. We have used the heuristic algorithm given in Table-178 32 to obtain a solution to this problem. The algorithm can be stopped $\frac{1}{79}$ when the representative examples of all binding sites stop chang $\frac{1}{80}$ 33 ing, or on the basis of a user-defined maximum number of itera $\overline{81}$ 34 tions. In all our experiments, the algorithm converged in 10 itera $\frac{1}{82}$ 35 36 tions or less. A trained MI-1 SVM can be used to produce discriga 37 minant function scores for any given residue in a protein.

The quadratic programming problem in the MI-1 algorithm can be solved in the primal or in the dual. The primal formulation of the problem (3) is more efficient than the dual when the dimensionality of the feature vector is smaller than the number of training examples. The dual formulation of the quadratic programming problem (based upon the Lagrange of the primal) is given by:

$$\min_{\alpha} \left(\frac{1}{2} \sum_{b, j \in N(b)} \sum_{c, j \in N(c)} \alpha_{i}^{c} \left(\Delta_{i}^{c^{T}} \Delta_{j}^{b} \right) \alpha_{j}^{b} - \sum_{b, j \in N(b)} \alpha_{j}^{b} \right)$$
such that $\forall b$:

$$\alpha_{j}^{b} \geq 0, j \in N(b) ,$$

$$(4)$$

 $\sum_{j \in N(b)} \alpha_j^{\rm b} \leq \frac{C}{M}.$ Here $\alpha_j^{\rm b}$ is the Lagrange variable corresponding to the primal constraint $w^T \phi(x^b) \geq w^T \phi(x_j) + 1 - \xi_b$ and $\Delta_j^{\rm b} = \phi(x^b) - \phi(x_j)$. The dual formulation reveals some interesting aspects of the MI-1 SVM. It shows that Lagrange variables (α) only exist for negative examples, and that the sum of all α for negative examples from a single protein is constrained to be less than or equal to C/M. This differs from a conventional SVM formulation which requires that each of the α , on its own, should be less than or equal to C/M and the sum of products of α from all training examples with their corresponding labels should be zero. Thus, the MI-1 SVM formulation and this can potentially lead to a better solution.

2.5 CaM Binding Prediction

In this paper, we compare the following two strategies for CaM binding prediction.

2.5.1 Discriminant Function Scoring

The maximum discriminant function score across all windows in a protein can be used as the CaM binding propensity of that protein. This approach was used in (Hamilton, Reddy, & Ben-Hur, 2011) to predict CaM binding of proteins in the *Arabidopsis thaliana* proteome. In their method, the scores were generated using a standard SVM classifier trained for binding site prediction. In this paper, we use the scores from MI-1 SVM instead.

2.5.2 Cascaded Classification

We implemented a two stage cascaded classification approach for CaM binding prediction. In the first stage the window in a given protein with the highest MI-1 SVM discriminant function score is chosen as the most likely binding site window for that protein. This is done for all proteins in the training set. In the second stage, a standard SVM is trained to discriminate between the most likely binding site windows in positive examples (known CaM binding proteins) and negative examples (non CaM-binding proteins). Once the second stage SVM has been trained, the binding propensity of a test protein can be estimated by first finding its most likely binding site window using MI-1 SVM, and then evaluating the discriminant function value of the second stage SVM for the chosen window. A Gaussian kernel was used in the second stage SVM as it performed significantly better than a linear kernel. However, the use of non-linear kernels in MI-1 SVM did not seem to improve performance.

1 2.6 Feature Representations

2 The performance of the learning methods described above for 53

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3 binding site and CaM binding prediction was analyzed using 354

4 number of feature representations which are presented next.

5 2.6.1 *p*-spectrum The *p*-spectrum $\phi(x)$ of a string over an 56

- 6 alphabet Σ is a vector each of whose components $\phi_v(x)$ is the 57
- 7 number of occurrences of each length-*p* substring *v* in the string x58
- 8 The *p*-spectrum kernel between two strings is given by the corre59
- 9 sponding Euclidean dot product (Leslie, Eskin, & Noble, 2002). 60

10 2.6.2 Position dependent *p*-spectrum The position dependent p_{62} 11 spectrum $\phi(x)$ of a string *x* is a vector of indicator variables 12 $\phi_{(v,k)}(x)$ each showing whether the length-*p* substring *v* occurs at 13 position *k* in the string *x*. The resulting position dependent p_{65} 14 spectrum kernel is given by: $K^{PD}(x, x') = \phi(x)^T \phi(x')$. The posified 15 tion dependent kernel takes the relative position of an amino acident 16 in a window into account whereas the *p*-spectrum kernel does not. 68

17 2.6.3 Position dependent gappy triplet This feature representa69 18 tion quantifies the occurrences of motifs of the form $ax^m bx^n c$ **70** 19 where a,b,c are amino acids and x^m indicates m don't-care posi-71 20 tions. For a given string x, the feature vector $\phi^{m,n}(x)$ of the posi**72** 21 tion dependent gappy triplet comprises of variables $\phi_{(a,b,c,k)}^{m,n}(x)_{73}$ which indicate whether the motif $ax^m bx^n c$ starts at position k in 74 22 the string or not. The kernel $K^{m,n}(x, x') = \phi^{m,n}(x)^T \phi^{m,n}(x')$ be-75 23 tween two strings tells us the number of locations in the two strings $\frac{1}{2}$ 24 25 that have the same motif starting at them. We have used multiple77 position dependent gappy triplet kernels as $K^{PDGT}(x, x') = \frac{1}{78}$ 26 $\sum_{m=0}^{4} \sum_{n=0}^{4} K^{m,n}(x, x')$. This kernel allows us to extract meaning 79 27 ful information about motifs for CaM binding sites and is only $\frac{1}{80}$ 28 29 used for binding site prediction for this purpose. 81

30 We perform normalization of any kernel representation using the 82 31 cosine kernel $K_{cos}(x, x') = \frac{K(x, x')}{\sqrt{K(x, x)K(x', x')}}$. 83

32 2.7 Evaluation Methodology

33 We use Leave-One-Protein-Out (LOPO) cross validation in orde86 34 to analyze the performance for binding site prediction. In LOPO87 all examples (positive or negative) from a single protein are held 88 35 out while the classifier is trained on the remaining proteins. The $\frac{89}{2}$ out while the classifier is trained on the remaining proteins. In 90 classifier is then evaluated over the examples from the held 009136 37 protein. We evaluate the following performance metrics and $us \frac{91}{92}$ 38 their average across all proteins to make comparisons between \tilde{p}_3 39 40 methods and kernels: 94

- 41 a. Protein level area under the ROC curve (AUC): The area 95
 42 (expressed as percentage) under the Receiver Operating Char 97
 43 acteristic (ROC) curve (the plot of true positive rate versus 98
- false positive rate) obtained for windows in a given protein. 39
- 45 b. Protein level Area under ROC 10% Curve (AUC_{0.1}): The art100 (expressed as percentage) under the ROC curve based on upor to the first 10% false positives in a protein.
- 48 c. False-Hit Ratio (FH-measure): The percentage of non-bindin 104
 49 site windows (out of the total number of non-binding site
- 50 windows) that have a score higher than the maximum scoring
- 51 window in the known binding site. This measure tells us hopo6

many non-binding site windows are expected with a score higher than the true binding site window.

d. True Hit Probability (TH-measure): For a given protein, a true hit is defined to occur when the residue at the center of the highest scoring window for that protein lies within a binding site. The average number of true hits across all proteins (called the TH-measure) represents the probability of the maximum scoring window predicted by a classifier to lie within a true binding site.

The AUC is a measure of how good a particular method is in ranking binding site windows above non binding sites. AUC_{0.1} gives us a sense of how good are the top scoring windows produced by a classifier. The FH measure represents the chances of a non-binding site window to be ranked higher than a true binding site window. The TH-measure tells us about the chances of the highest scoring window predicted by a classifier to belong to a true binding site. Both the TH and the FH measures provide meaningful information about the accuracy of the method to a biologist who intends to use the proposed prediction scheme to verify potential binding site locations experimentally.

We use AUC as the performance metric for CaM binding prediction. AUC can be directly computed from the estimated CaM binding propensities when using the discriminant function scoring approach. With the cascaded classification approach, AUC is obtained from 5-fold stratified cross validation with nested grid search for model selection. In cross-validation, it was ascertained that two proteins with more than 40% sequence similarity are in the same fold (evaluated using BLASTCLUST from the NCBI BLAST package (Altschul, Gish, Miller, Myers, & Lipman, 1990). Moreover, the data for CaM binding prediction in *A. thaliana* did not include any proteins which were part of the MI-1 training set.

2.8 Model Selection

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In order to perform model selection (the choice of the cost parameter *C*) for the vanilla and MI-1 SVM formulations for binding site prediction, we used nested 5-fold cross validation within each iteration of LOPO cross validation. The TH-measure obtained from the 5-fold cross validation is then used to choose the value of *C* for that iteration of LOPO. The values of C that were used in the nested cross validation are $\{0.01, 0.1, 1.0, 100\}$.

As mi-SVM takes a long time to train, nested cross validation could not be performed. Instead we evaluated the LOPO cross validation performance (TH-measure) of mi-SVM with different values of *C* in {0.01, 0.1, 1.0, 10, 100} and the best results with the optimal value of C=10 are reported. This method for selection of *C* for mi-SVM can potentially lead to over optimistic performance estimates. This is not an issue, since our claim is that the proposed approach performs better.

In the case of CaM binding prediction in *Arabidopsis thaliana* using cascaded classification, we performed a nested (5-fold) grid search within each cross validation fold for selecting the parameter values of the second-stage SVM. Values of *C* in the SVM and γ of the Gaussian kernel ($K(x_1, x_2) = exp(-\gamma ||\phi(x_1) - \phi(x_2)||^2)$)) were chosen from {0.1, 1, 10, 100} and {0.005, 0.02, 0.5, 2.0} respectively.

3 RESULTS & DISCUSSION

Table 2 presents the LOPO cross validation results for the three57 1 2 SVM formulations for the 1-spectrum, position dependent 158 3 spectrum and the combination of the two feature representations 4 for predicting CaM binding sites. We observe that both MIL for-5 mulations (mi-SVM and MI-1 SVM) perform better than the vanil-6 la SVM. This shows the value of expressing binding site prediction 7 as a multiple instance learning problem. This is particularly evident 8 with the use of position dependent feature representations, as they 9 are more sensitive to changes in relative position of an amino acid 10 in a window within the binding site than position independent fea-11 ture representations. It can also be noted that the accuracy of MI-1 12 SVM is noticeably better than mi-SVM. We believe that this im-13 provement stems from the fact that the proposed scheme imple-14 ments a more realistic model of the binding site prediction prob-15 lem. The improvement resulting from switching to a position de-16 pendent feature representation is also larger for MI-1 SVM than 17 that observed in the case of mi-SVM. The higher $AUC_{0,1}$ scores 18 indicate the improved sensitivity and specificity of MI-1 SVM 19 which is also reflected in the ~8% improvement in the TH-20 measures and the decrease in the FH-measure.

The vanilla SVM approach is the same as the method in 59
(Hamilton, Reddy, & Ben-Hur, 2011), which they showed work 60
comparably as the neural network approach of (Radivojac,
Vucetic, O'Connor, Uversky, Obradovic, & Dunker, 2006). There 62
fore we conclude that the proposed scheme performs better than 64
previously reported approaches. 65

27 We also compare the performance of these approaches with 36628 naive local alignment based method for finding CaM binding sites67 In this method, local alignment between a held out protein and the 29 binding sites of the remaining proteins is performed and if the bes $\frac{69}{2}$ 30 70 scoring alignment overlaps (by at least ten residues) with the 31 32 known binding site in the held out protein, it is considered to be a 33 true hit. This approach gives a TH% of 39.5%. This shows that the 34 machine learning approaches presented in this paper use more than 35 sequence similarity to make better predictions.

36 We have also performed an analysis of the stability of the results 37 for the MI-1 and the vanilla SVMs by averaging performance sta-38 tistics of 12 runs of 5-fold cross validation. This analysis was not 39 performed for mi-SVM or for the gappy triplet kernel with MI-1 40 SVM owing to their large time requirements. The 5-fold cross 41 validation results for both the methods are very similar to the 42 LOPO cross validation results. The maximum standard deviation 43 in a particular performance metric across different feature repre-44 sentations obtained from the 5-fold cross validation for vanilla and MI-1 SVMs is given in Table 2. This statistic gives an idea of the $\frac{73}{73}$ 45 46 variability of the results with respect to changes in the data. 74

47 Figure 2 shows the output of the MI-1 SVM for a single protein75 48 for the position dependent and position independent versions of the 76 49 1-spectrum feature representation. It is quite clear that the output 7 for the position independent features is much smoother than that 7850 51 from the position dependent 1-spectrum features. This is because 52 the position independent 1-spectrum feature vector representation 53 changes only slightly as the window is translated by one position, 54 whereas the position dependent feature vector can change dramati-55 cally. Due to the increased resolution power, the position depend-56 ent features lead to a classifier that is able to correctly predict both

binding sites in the example shown in Figure 2, which is not achieved using the position dependent features.



Fig. 2. MI-1 discriminant values along the length of a held-out protein. with the position independent (top) and the position dependent (bottom) 1-spectrum features.

Table 2. Results across methods and kernels. The features are 1-spectrum (1-Spec), position dependent 1-spectrum (PD-1) and the combination (Comb) of the 1-Spec and PD-1 representations. The Max Std. rows show the maximum standard deviation of a particular performance metric using the above feature representations. Results with the position-dependent Gappy triplet kernel (Gappy) with MI-1 SVM are also reported (for a single run due to its longer computational time). Bold numbers indicate the best value (across all methods) for a particular metric using a particular feature representation. (AUC: Area under the ROC curve, AUC_{0.1} AUC for first 10% false positives, TH: True hit, FH: False hit).

		,	,		
Method	Features	AUC	$AUC_{0.1}$	TH %	FH %
	1-Spec	95.5	53.9	66	2.6
Vanilla	PD-1	95.6	54.5	64	2.5
SVM	Comb.	95.9	55.1	65	2.1
	Max. Std.	0.16	0.59	2.2	0.15
	1-Spec	95.5	54.4	64	2.6
mi-SVM	PD-1	96.0	55.8	69	2.1
	Comb.	96.2	55.6	68	1.9
	1-Spec	96.0	54.3	62	2.1
MT 1	PD-1	96.8	58.5	72	1.3
SVM	Comb.	96.9	59.0	75	1.2
5 V IVI	Max Std.	0.14	0.80	3.4	0.11
	Gappy	96.5	58.5	68	1.6

Table 3. Results of CaM binding prediction for Discriminant Function Scoring and Cascaded Classification with an SVM with a Gaussian kernel. The features are 1-spectrum (1-Spec), position dependent 1-spectrum (PD-1) and the combination (Comb) of the 1-Spec and PD-1 feature representations. Using Cascaded Classification with a liner kernel in the second stage SVM instead of the Gaussian kernel, the best AUC was 0.72 with 1-spectrum features. (AUC: Area under the ROC curve).

Method	Features	AUC
Discriminant Function Scoring	1-Spec	71.9
	PD-1	70.1
	Comb.	71.9
Cascaded Classification	1-Spec	75.3
	PD-1	71.1
	Comb.	72.3



Fig. 3. (a) Weights of different amino acids in the (position independent) 1-spectrum feature representation (b) Heat map of the weights of different amino acids versus their position from the MI-1 SVM position dependent 1-spectrum feature representation and (c) Top 100 (in terms of their weights) motifs from the position dependent gappy triplet kernel. The last (numeric) column shows actual weight values.

4 We have also analyzed the weight vectors from different feature31 5 representations in order to extract amino-acid patterns informativ@2 6 of CaM binding sites. The plot of weights from the 1-spectrun33 7 features and the position dependent 1-spectrum features are shownB4 8 in Figure 3a and 3b respectively. The weights for the 1-spectrum 35 9 features closely follow the amino acid propensities in CaM bind36 10 ings sites (Hamilton, Reddy, & Ben-Hur, 2011), with R (Arginine)37 11 K (Lysine) and W (Tryptophan) showing large positive weights38 12 whereas D (Aspartic acid), E (Glutamic acid) and P (Proline) have39 13 large negative weights. The plot of the position dependent 140 14 spectrum features indicates that the importance of different aminc41 15 acids varies with their position in the window. For example, Argi42 16 nine shows large positive weights in the middle of the window, and 43 17 negative weights in the ends; Glutamic acid shows the opposite44 18 behavior. This indicates that the classifier is indeed learning a45 19 position dependent model. 46

20 The results of 5-fold cross validation using the position depend47 21 ent gappy triplet kernel (K^{PDGT}) shown in Table 2 indicate tha**48** 22 this kernel provides comparable performance to other feature rep49 23 resentations using MI-1 SVM. Since the number of dimensions in 50 24 the feature representation of the gappy triplet kernel is much large 51 25 than the number of training examples, MI-1 SVM learning was52 26 performed using the dual formulation for this kernel, which is 53 27 more computationally intensive. That is why we have used 5-fold54 28 cross validation instead of LOPO cross-validation. 55

Next, we ranked the features of the gappy triplet kernel in terms56
 of their weights in MI-1 SVM learning in order to find motifs tha57

are associated with CaM binding. Figure 3c shows the top 100 motifs and their positions. We observe that motifs tend to associate with particular positions, showing that MI-1 SVM uses the flexibility in choosing a representative window to "align" instances of CaM binding sites (for instance, notice the presence of 'R' at positions 10 and 11 across different features). Moreover, it is able to find parts of known CaM binding motifs provided in the CaM Target Database (Yap, Kim, Truong, Sherman, Yuan, & Ikura, 2000). The CaM Target Database classifies CaM binding targets into 5 groups, each characterized by certain motifs: three predominantly calcium dependent motifs (1-10, 1-14 and 1-16, named according to the position of large hydrophobic residues), the IQ motif which is typically not dependent on calcium concentration, and others. As is evident from Figure 3c, IQ, QxxxR, RxxxxR, RGxxxR, RxxL, KxxxxR receive large positive weights. These motifs are components of the IQ subclass of motifs. Other features belonging to different subclasses of motifs that receive large positive weights include: AxxI, IxxxF, LxxV, (from the 1-14 subclass), RR, KK, RxF (from the 1-10 subclass) etc. This clearly illustrates the capabilities of the proposed scheme to learn CaM binding motifs. We also note that most of the top ranking features correspond to a motif with 3 or 4 don't care positions. This is in agreement with the known fact that CaM binding usually occurs via an alpha helix, and this corresponds to the periodicity of the alpha helix.

On the task of CaM binding prediction (Table 3), the performance of discriminant function scoring is only marginally better than that of the 1-spectrum feature representation used in

1 (Hamilton, Reddy, & Ben-Hur, 2011). However, with the cascade 65 2 classification approach with a Gaussian kernel, the results are sig56 3 nificantly better. Even though the AUC for the position independ57 4 ent 1-spectrum features is higher than that of the position dependent 1-spectrum reatures is inglier time ent features, the AUC_{0.1} was higher for position dependent features 585 6 (29.1) in comparison to the simple 1-spectrum features (26.6). 59 In order to obtain a better understanding of what our classifier $\tilde{60}$ 7 8 picks up, we considered the proteins that are not known to bind to61 CaM and ranked that list according to the score provided by $ou \frac{62}{2}$ 9 classifier. We then tested for enrichment of GO terms of segment 63 10 of that list: the first 1000 proteins, proteins 1001 - 2000 etc., using 55 11 12 the GOrilla tool (Eden, Navon, I., Lipson, & Yakhini, 2009). Fo.66 13 the first 1000 we found enriched terms that are in agreement with known functions of CaM binders (Reddy, Ben-Hur, & Day, 2011)69 14 15 In GO molecular function, transcription function activity, and 70 16 CaM-dependent kinase activities were the most highly enriched $\frac{1}{2}$ with adjusted p-values below 10^{-10} . All other enriched terms were

- 17
- related to these except for "inward rectifier potassium channel74 18
- 19 activity" which had an adjusted p-value of 0.02. In GO biologica $\overline{P}5$
- process namespace all the terms except for "response to carbohy-76 20 21
- drate stimulus" (adjusted p-value 0.02) were related to phosphory-78 22 lation and various regulatory processes. In analyzing enrichment79
- 23 for size-1000 chunks we found that the p-values for these function g_{2}
- and processes went down as we went down the ranked list, and for 24
- 25 proteins ranked 5000-6000, no terms showed enrichment.

26 **CONCLUSIONS & FUTURE WORK**

- 27 We have presented a novel MIL algorithm for CaM binding site88 prediction called MI-1 SVM, and shown its performance ad 89 28 ٩N vantages in comparison to the standard MIL SVM and regular 29 30 SVM, which was used in previous work. Our new MIL formula $\tilde{92}$
- 31 tion captures the minimal constraints that a good binding site clas93
- sifier needs to have, and we believe this is the reason for its better 32
- accuracy. Not only that, it also runs more than twice as fast $a\widetilde{96}$ 33
- 34 standard MIL SVM (running time on a dataset of 16,060 windows)7
- was 510.5s for MI-1, 1059.1s for mi-SVM, and 348.3s for vanilla 35 36 SVM). 100
- 37 Expressing binding site prediction as an MIL problem is a natur $\frac{1}{2}$ $\check{D}1$
- 38 way to incorporate uncertainty about binding site location, and oil 02
- results show that this allows the classifier to "align" binding site 10339
- and learn position-dependent motifs that characterize the binding 40
- 41 site. The proposed scheme also shows its efficacy in prediction $\mathbf{406}$ 42 107
- CaM binding proteins. In general, binding sites in proteins or nucleic acids are n_{Pho}^{108} 43
- 44 contiguous in sequence as they are in CaM binding proteins. MIII
- 45 SVM can be extended to solve the generic problem of binding sile11
- prediction by using sequence-based features that capture the no $\frac{112}{13}$ 46
- contiguous nature of binding sites. Currently, MI-1 SVM generida 47
- 48 ates the CaM binding propensity along a protein's length and ca11-15 49 not explicitly identify multiple binding sites. Identifying the num16
- 50 ber of binding sites in a protein remains for future work.

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