

Margin trees for high-dimensional classification

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Abstract

We propose a method for the classification of more than two classes, from high-dimensional features. Our approach is to build a binary decision tree in a top-down manner, using the optimal margin classifier at each split. We implement an exact greedy algorithm for this task, and compare its performance to less greedy procedures based on clustering of the matrix of pairwise margins. We compare the performance of the “margin tree” to the closely related “all-pairs” (one versus one) support vector machine, and nearest centroids on a number of cancer microarray datasets. We also develop a simple method for feature selection. We find that the margin tree has accuracy that is competitive with other methods and offers additional interpretability in its putative grouping of the classes.

Keywords: Maximum margin classifier, support vector machine, decision tree, CART

1 Introduction

We consider the problem of classifying objects into two or more classes, from a set of features. Our main application area is the classification of cancer patient samples from gene expression measurements.

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In these high-dimensional problems the number of features p is larger than the sample size N . The support vector classifier (Boser et al. 1992, Vapnik 1996) has become popular in this setting and generally shows good generalization performance. The classes are generally separable when $p > N$, and in this instance the support vector classifier yields the maximum margin separating hyperplane, when its cost complexity parameter is taken to be large.

When the number of classes K is greater than two, maximum margin classifiers do not generalize easily. Various approaches have been suggested, some based on the two-class classifier (“one-versus-all” and “one-versus one” or “all pairs”), and others modifying the support vector loss function to deal directly with more than two classes (Weston & Watkins 1999, Lee et al. 2004, Rosset et al. 2005). These latter proposals have a nice generalization of the maximum margin property of the two class support vector classifier. Statnikov et al. (2004) contains a comparison of different support vector approaches to classification from microarray gene expression cancer datasets. While these methods can produce accurate predictions, they lack interpretability. In particular, with a large number of classes, the investigator may want not only a classifier but also a meaningful organization of the classes.

In this paper we propose a tree-based maximum margin classifier. Figure 1 illustrates our idea. There are three classes and two features, as shown in the top left panel. We seek the line that partitions the classes into two groups, that has the maximum margin. [The margin is the minimum distance to the decision line among all of the data points.]

The best line is shown in the top right panel, splitting class 1 from classes 2 and 3. We then focus just on classes 2 and 3, and their maximum margin classifier is shown in the bottom left. The overall top-down classifier is summarized by the binary tree shown in the bottom right panel.

We employ strategies like this for larger numbers of classes, producing a binary decision tree with a maximum margin classifier at each junction in the tree.

In section 2 we give details of the margin tree classifier. Section 3 shows the application of the margin tree to a number of cancer microarray datasets. For construction of the tree, all of the classifiers in the margin tree use all of the features (genes). In section 4 we discuss approaches to feature selection. Finally in section 5 we have some further comments and a discussion of related work in the literature.

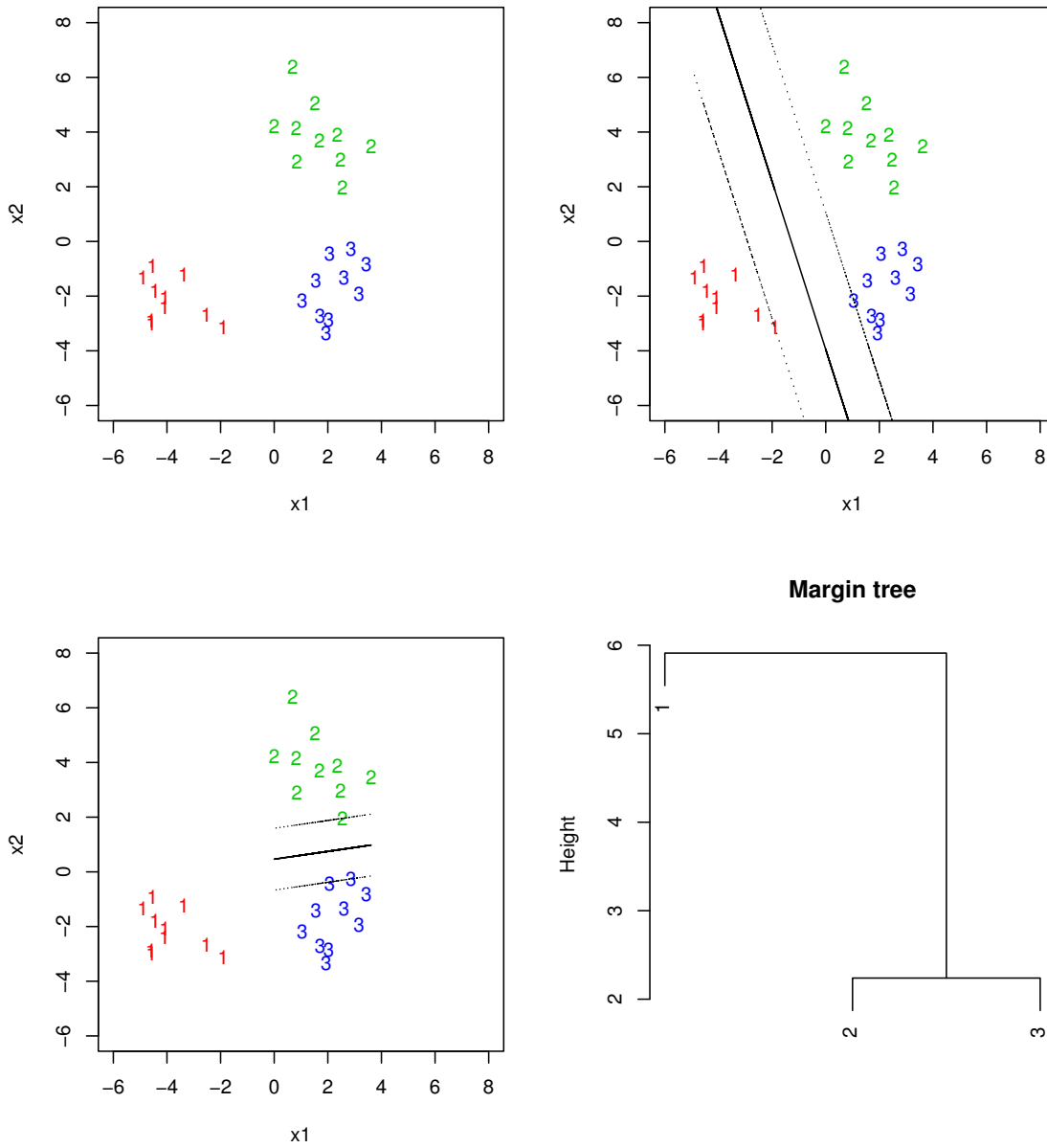


Figure 1: *Simple illustration of a margin tree. There are three classes shown in the top left panel. The largest margin is between class 1 and (2,3), with the optimal classifier shown on the top right. Then we separate class 2 from 3, in the bottom left. These top-down splits are summarized in the margin tree in the bottom right.*

2 The margin tree classifier

Denote the gene expression profiles by $x_j = (x_{1j}, x_{2j}, \dots, x_{pj})$ for $j = 1, 2, \dots, N$ samples falling into one of K classes. The features (genes) are indexed by $i = 1, 2, \dots, p$.

Consider first the case of $K = 2$ classes, C_1 and C_2 . The class outcome is denoted by $y_j = \pm 1$. The maximum margin classifier is defined by the constant β_0 and the weight vector β with components $\sum_i \beta_i^2 = 1$ that maximizes the gap between the classes, or the ‘‘margin’’. Formally,

$$(\beta_0, \beta) = \max(C) \quad \text{where } y_j(\beta_0 + \sum_i \beta_i x_{ij}) \geq C \quad \forall j. \quad (1)$$

The achieved margin $M = 2 \cdot C$. In the examples of this paper, $p > N$ so that all classes are separable and $M > 0$.

Now suppose we have $K > 2$ classes. We consider three different strategies for constructing the tree. These use different criteria for deciding on the best partition of the classes into two groups at each juncture. Having settled on the partition, we use the maximum margin classifier between the two groups of classes, for future predictions.

Let $M(j, k)$ be the maximum margin between classes j and k . Also, let G_1, G_2 be groups of classes, and let $M(G_1, G_2)$ denote the maximum margin between the groups. Finally, denote a partition by $P = \{G_1, G_2\}$.

Then we consider three approaches for splitting a node in the decision tree:

- *Greedy*: maximize $M(G_1, G_2)$ over all partitions P .
- *Single linkage*: Find the partition P yielding the largest margin M_0 so that $\min M(j_1, j_2) \leq M_0$ for $j_1, j_2 \in G_k, k = 1, 2$ and $\min M(j_1, j_2) \geq M_0$ for $j_1 \in G_1, j_2 \in G_2$.
- *Complete linkage*: Find the partition P yielding the largest margin M_0 so that $\max M(j_1, j_2) \leq M_0$ for $j_1, j_2 \in G_k, k = 1, 2$ and $\max M(j_1, j_2) \geq M_0$ for $j_1 \in G_1, j_2 \in G_2$.

The greedy method finds the partition that maximizes the resulting margin overall all possible partitions. Although this may seem prohibitive to compute for a large number of classes, we derive an exact, reasonably fast algorithm for this approach (details below).

The second and third methods require some explanation. They are derived from the *bottom up* (as opposed to top-down) clustering methods. Single linkage clustering successively merges groups based on the minimum distance between any pair of items in each of the group. Complete linkage clustering does the same, but using the maximum distance. Now having built a clustering tree bottom-up, we can interpret each split in the tree in a *top-down* manner, and that is how criteria (2) and (3) above were derived. In particular it is easy to see that the single and complete linkage problems are solved by single and complete linkage agglomerative clustering, respectively, applied to the margin matrix $M(j_1, j_2)$.

The greedy method focusses on the form of the final classifier, and tries to optimize that classification at each stage. Note that the greedy method cares only about the distance between classes in the different partitions, and not about the distance between classes within the same partition. Both the single linkage and complete linkage methods take into account both the between and within partition distances. We will also see in the next section that the complete linkage method can viewed as an approximation to the greedy search.

Figure 2 shows a toy example that illustrates the difference between the greedy and complete linkage algorithms. There are six classes with circular distributions. The greedy algorithm splits off group 1,2, and 3 in succession, and then splits off 4,5,6 as a group. This is summarized in the bottom left panel. The complete linkage algorithm in the bottom right panel instead groups 1,2 and 3 together and 4,5, and 6 together. The complete linkage tree is more balanced and hence may be more useful biologically.

In the experiments in this paper we find that:

- All three methods produce about the same test set accuracy, and about the same as the all-pairs maximum margin classifier.
- The complete linkage approach gives more balanced trees, that may be more interpretable than those from other two methods; the single linkage and greedy methods tend to produce long stringy trees that usually split off one class at a time at each branch. The complete linkage method is also considerably faster to compute than the greedy method.

Thus the complete linkage margin tree emerges as our method of choice.

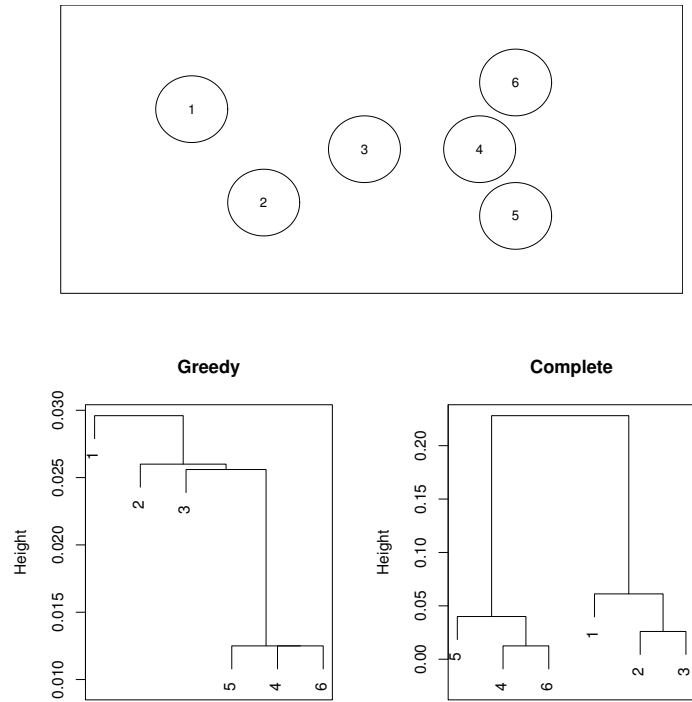


Figure 2: *A toy example illustrating the difference between the greedy and complete linkage algorithms. There are six classes with circular distributions (top panel). The greedy algorithm splits off groups 1,2, and 3 in succession, and then splits off 4,5,6 as a group. This is summarized in the bottom left panel. The complete linkage algorithm (bottom right panel) instead groups 1,2 and 3 together, and 4,5, and 6 together*

2.1 An exact algorithm for the greedy criterion

A key fact is

$$M(G_1, G_2) \leq \min\{M(j_1, j_2), j_1 \in G_1, j_2 \in G_2\}. \quad (2)$$

That is, the margin between two groups of classes is less than or equal to the smallest margin between any pair of classes, one chosen from each group.

Now suppose we seek a partition P with margin M . Rather than enumerate all possible partitions (and their associated maximum margin classifiers), we can speed up the computation by constructing the complete linkage clustering tree, and collapsing all nodes at height M . We know that all classes in any collapsed node must be on the same side of the decision plane, since each class has margin at least M with every other class in that node. Hence we need only consider partitions that keep the collapsed nodes intact.

We summarize the algorithm below:

Exact computation of the best greedy split

1. Construct the complete linkage clustering tree based on the margin matrix $M(j_1, j_2)$.
2. Starting with all classes at the top of the tree, find the partition of each individual class versus the rest, and also the partition that produces two classes in the complete linkage tree. Let M_0 be the largest margin achieved amongst all of these competitors.
3. Cut the complete linkage tree at height M_0 , and collapse all nodes at that height.
4. Consider all partitions of all classes that keep the collapsed nodes intact, and choose the one that gives maximal margin \hat{M} .

This procedure finds the partition of the classes that yields the maximum margin. We then apply this procedure in a top-down recursive manner, until the entire margin tree is grown.

Note that if approximation (2) is an equality, then the complete linkage tree is itself the greedy margin classifier solution. This follows because $\hat{M} = M_0$ in the above algorithm.

2.2 Example: 14 cancer microarray data

As an example, we consider the microarray cancer data of Ramaswamy et al. (2001): there are 16,063 genes and 198 samples in 14 classes. The authors provide training and test sets of size 144 and 54 respectively.

The margin trees are shown in Figure 3. The length of each (non-terminal) arm corresponds to the margin that is achieved by the classifier at that split. The final classifiers yielded 18, 18 and 19 errors, respectively on the test set. By comparison, the all-pairs support-vector classifier yielded 20 errors and the nearest centroid classifier had 35 errors. Later we do a more comprehensive comparison of all of these methods. We note that the greedy and single linkage margin tree are “stringy”, with each partition separating off just one class in most cases. The complete linkage tree is more balanced, producing some potentially useful subgroupings of the cancer classes.

In this example, full enumeration of the partitions at each node would have required computation of 16,382 two class maximum margin classifiers. The exact greedy algorithm required only 485 such classifiers. In general the cost savings can vary, depending on the height M_0 of the initial cut in the complete linkage tree.

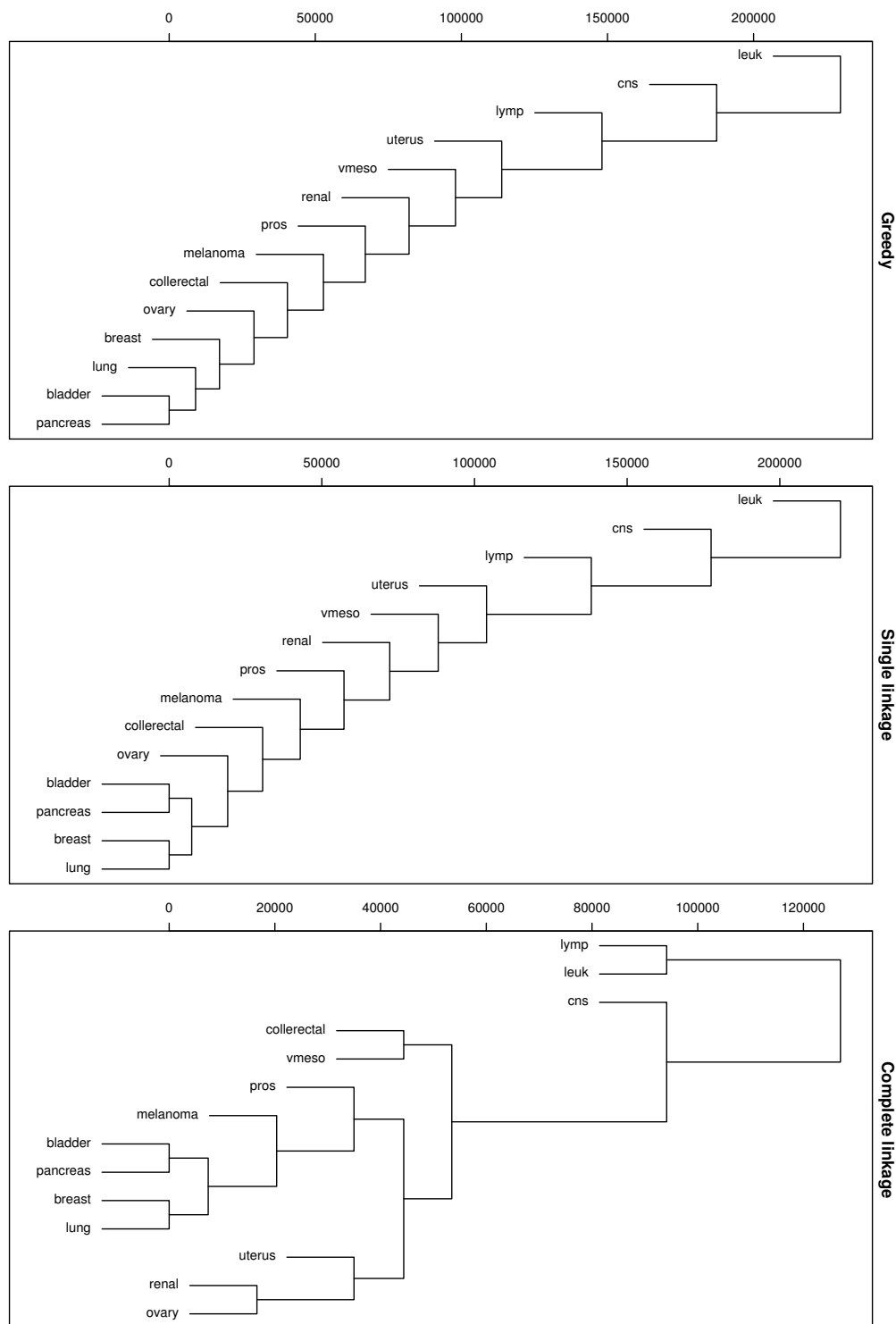
Figure 4 displays the margins that were achieved by each method at their collection of splits. We see that the complete method gives larger margins than the other methods. The largest margin achieved is about 49,000, corresponding to the split between class CNS and Collerectal and so on.. This is larger than the margin between Leukemia and the rest at the top of the greedy tree. This shows that the greediness of the exact algorithm can hurt its overall performance in finding large margin splits.

Table 3 shows the the number of times each classifier disagreed on the test set. The number of disagreements is quite large. However the methods got almost all of the same test cases correct (over 90% overlap), and the disagreements occur almost entirely for test cases in which all methods got the prediction wrong.

3 Application to other cancer microarray datasets

We applied the methods described earlier to the seven microarray cancer datasets shown in Table 2. In each case we randomly sampled 2/3rds of

Figure 3: *Margin trees for the 14-tumor cancer data of Ramaswamy et al. (2001)*



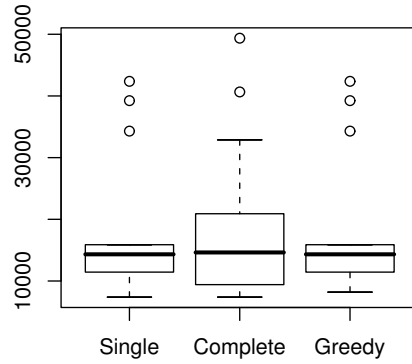


Figure 4: *14 tumor cancer data: margins achieved by each method over the collection of splits.*

	SVM(OP)	MT(Greedy)	MT(single)	MT(Complete)
SVM(OP)	0	10	11	10
MT(Greedy)	10	0	7	2
MT(single)	11	7	0	9
MT(Complete)	10	2	9	0

Table 1: *Number of disagreements on the test set, for different margin tree-building methods.*

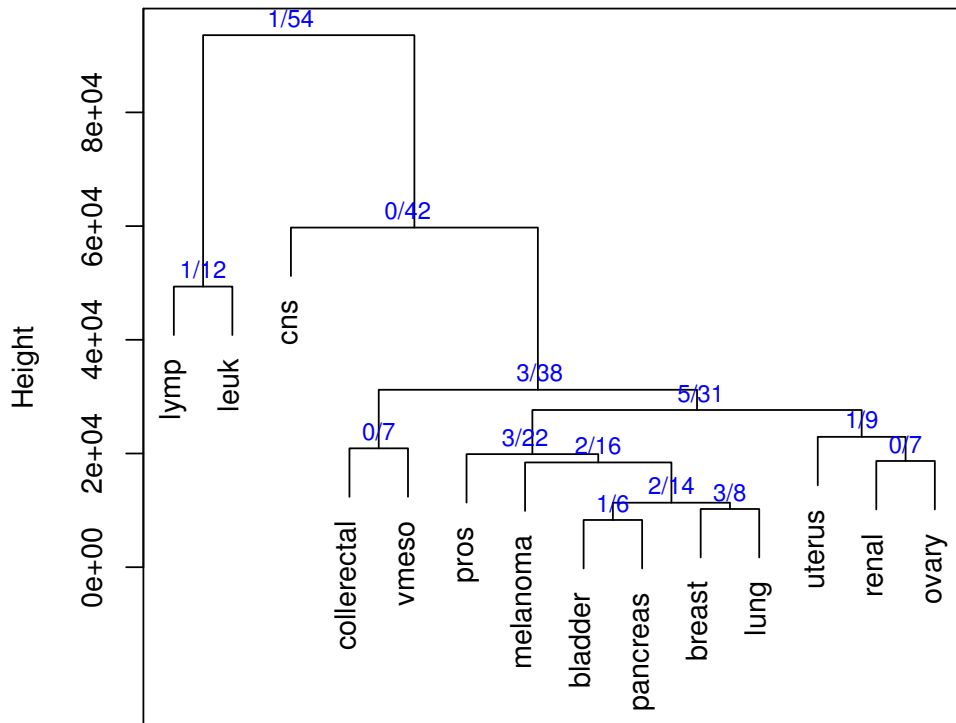


Figure 5: *Test errors for the 14 tumor dataset using the complete linkage approach. Error rates at each decision junction is shown: notice that the errors tend to increase farther down the tree.*

Name	# Classes	# Samples	# Features	Source
Brain	5	22	5597	Pomeroy et al. (2002)
Lymphoma	3	62	4026	Alizadeh et al. (2000)
Small round blue cell tumors	4	63	2308	Khan et al. (2001)
Stanford	14	261	4718	Munagala et al. (2004)
9 tumors	9	60	5726	Staunton et al. (2001)
11 tumors	11	174	12,533	Su et al. (2001)
14 tumors	14	198	16063	Ramaswamy et al. (2001)

Table 2: *Summary of datasets for comparative study*

Dataset	Nearest centroids	SVM (OVO)	MT(Single)	MT(Complete)	MT(Greedy)
Brain	0.236(0.026)	0.207(0.022)	0.229(0.018)	0.229(0.021)	0.221(0.017)
Lymphoma	0.010(0.006)	0.000(0.000)	0.000(0.000)	0.000(0.000)	0.000(0.000)
SRBCT	0.065(0.020)	0.011(0.011)	0.014(0.014)	0.014(0.014)	0.014(0.014)
Stanford	0.075(0.006)	0.063(0.006)	0.070(0.005)	0.079(0.007)	0.072(0.005)
9 tumors	0.478(0.009)	0.507(0.014)	0.526(0.013)	0.522(0.014)	0.545(0.014)
11 tumors	0.139(0.006)	0.110(0.005)	0.106(0.005)	0.106(0.005)	0.110(0.005)
14 tumors	0.493(0.007)	0.345(0.006)	0.318(0.007)	0.322(0.007)	0.315(0.007)

Table 3: *Mean test error rates (standard errors) over 50 simulations, from various cancer microarray datasets. SVM (OVO) is the support vector machine, using the one-versus-one approach; each pairwise classifier uses a large value for the cost parameter, to yield the maximal margin classifier; MT are the margin tree methods, with different tree-building strategies.*

the data to form a training set, and the balance of the data became the test set. The sampling was done in a stratified way to retain balance of the class sizes. This entire process was repeated 50 times, and the mean and standard errors of the test set misclassification rates are shown in Table 3. The nearest centroid method is as described in Tibshirani et al. (2001) and uses no shrinkage for feature selection: we discuss feature selection in section 4. We see that for problems involving more than 4 or 5 classes, the one-versus-one support vector classifier and the margin tree methods sometimes offer an advantage over nearest centroids. The margin tree methods are all very similar to each other and the one-versus-one support vector classifier.

4 Feature selection

The classifiers at each junction of the margin tree each use all of the features (genes). For interpretability it would be clearly beneficial to reduce the set of genes to a smaller set, if one can improve, or at least not significantly worsen, its accuracy. How one does this depends on the goal.

The investigator probably wants to know which genes have the largest contribution in each classifier. For this purpose, we rank each gene by the absolute value of its coefficient $\hat{\beta}_j$. Then to form a reduced classifier, we simply set to zero the first n_k coefficients at split k in the margin tree. We call this “hard-thresholding”.

How do we choose n_k ? It is not all clear that n_k should be the same for each tree split. For example we might be able to use fewer genes near the top of the tree, where the margins between the classes is largest.

Our strategy is as follows. We compute reduced classifiers at each tree split, for a range of values of n_k , and for each, the proportion of the full margin achieved by the classifier. Then we use a common value α for the margin proportion throughout the tree. This strategy allows the classifiers at different parts of the tree to use different number of genes. In real applications, we use tenfold cross-validation to estimate the best value for α .

Figure 6 shows the result of applying hard thresholding to the 14-class cancer data. The plot shows the test error as the margin proportion α is varied. The average number of genes at each of the 13 tree junctions is shown along the horizontal axis. We see that average number of genes can be reduced from about 16,000 to about 2,000 without too much loss of accuracy. But beyond that, the test error increases.

Figure 7 shows a more successful application of the feature selection procedure. The figure shows the result for one training/test split of the 11 class data (12,533 genes) described earlier. With no feature selection the margin tree (left panel) achieves 4/61 errors, the same as the one-versus one support vector machine. Hard-thresholding (middle panel) also yields 4 errors, with an average of just 167 genes per split. The margin proportion is shown at the top of the plot. The right panel shows the number of genes used as a function of the height of the split in the tree, for margin proportion 0.6.

The feature selection procedure described above is simple and computationally fast. Note that having identified a set of features to be removed, we simply set their coefficients $\hat{\beta}_i$ to zero. For reasons of speed and interpretability, we do not recompute the maximum margin classifier in the

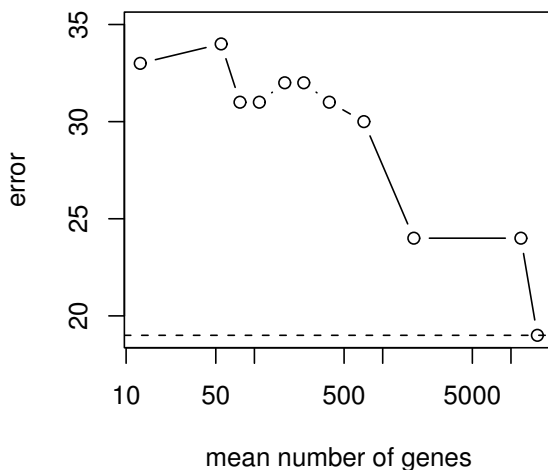


Figure 6: *14 tumor dataset: test errors for reduced numbers of genes.*

subspace of the remaining features (we do however recompute the classifier cutpoint, equal to midpoint between the classes). How much do we lose in this approximation? For the top and bottom splits in the tree of Figure 7, Figure 8 shows the margins achieved by the maximum margin classifier (black points) and the approximation (blue points) as the numbers of genes is reduced. The approximation gives margins remarkably close to the optimal margin until the number of genes drop below 100. Also shown in the figure are the margins achieved by recursive feature elimination (RFE) (Guyon et al. 2002). This is a full backward stepwise procedure, in which successive coefficients are dropped and the optimal margin classifier for the remaining features is recomputed. We see that RFE offers only a small advantage, when the number of genes becomes quite small.

4.1 Results on real datasets

Table 4 shows the results of applying the margin tree classifier (complete linkage) with feature selection, on the datasets described earlier. Tenfold cross-validation was used to choose the margin fraction parameter α , and

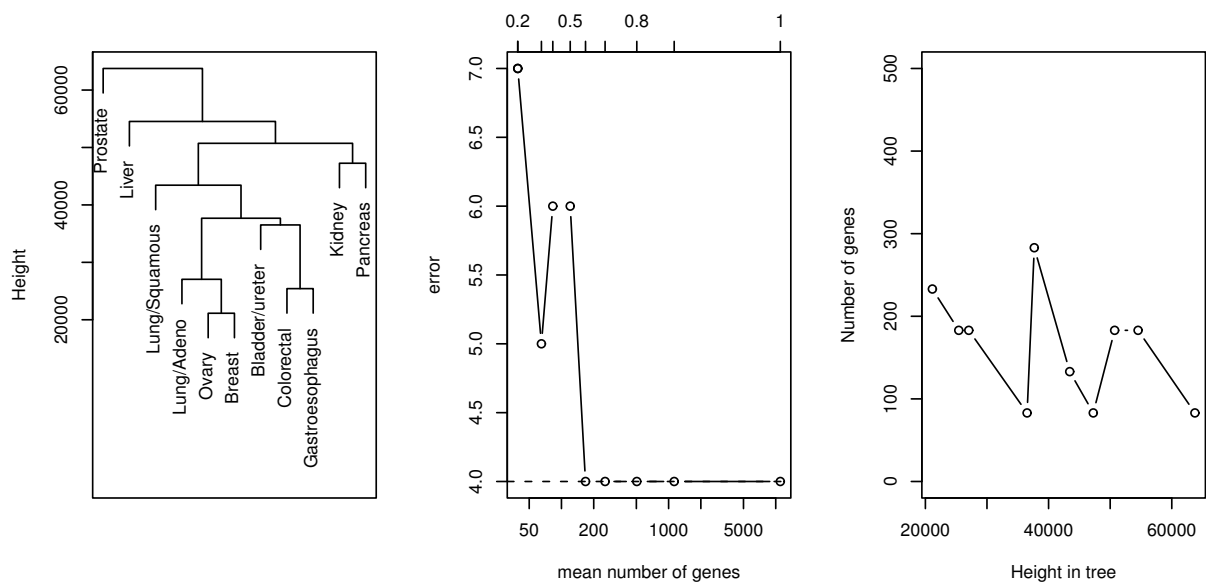


Figure 7: Results for 11 tumor dataset. The left panel shows the margin tree using complete linkage; the test errors from hard-thresholding are shown in the middle, with the margin proportion α indicated along the top of the plot; for the tree using $\alpha = 0.6$, the right panel shows the resulting number of genes at each split in the tree, as a function of the height of that split.

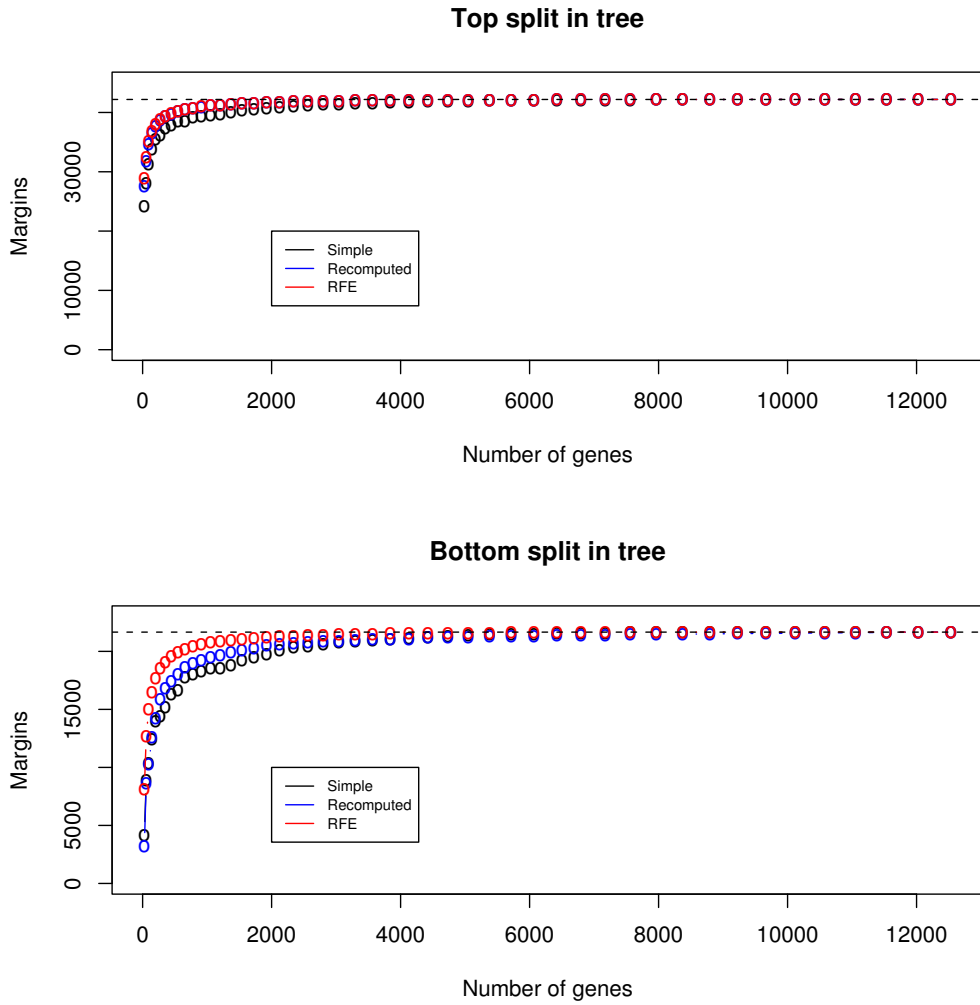


Figure 8: Results for the 11 tumor dataset: margins achieved by the maximum margin classifier using simple hard thresholding without recomputing the weights (black points), with recomputation (blue points) and recursive feature elimination (red points). Top panel refers to the top split in the margin tree; bottom panel refers to the bottom split.

	Nearest shrunken centroids			Margin tree with selection		
	CV errors	Test errors	# genes used	CV errors	Test errors	# genes/split
Brain	0.192(0.024)	0.276(0.04)	260(119.3)	0.264(0.011)	0.176(0.02)	483.3(194.3)
Lymphoma	0.022(0.002)	0.005(0.005)	3463.4(126.4)	0.000(0.000)	0.01(0.006)	408.5(145.0)
SRBCT	0.00(0.00)	0.014(0.009)	44.7(6.9)	0.004(0.002)	0.010(0.007)	5.08(8.6)
Stanford	0.064(0.005)	0.080(0.012)	4420.1(254.6)	0.064(0.006)	0.076(0.007)	1518(532.4)
9 tumors	0.381(0.008)	0.400(0.016)	1163.4(527.2)	0.503(0.019)	0.476(0.022)	2678.1(847.1)
11 tumors	0.125(0.004)	0.156(0.009)	4857.8(1685.9)	0.111(0.008)	0.110(0.012)	2720.5(1523.9)
14 tumors	0.397(0.007)	0.400(0.018)	3928.5(1392.8)	0.327(0.01)	0.311(0.015)	7563.8(1990.1)

Table 4: *CV and test error rates for nearest shrunken centroids and margin trees with feature selection by simple hard thresholding.*

both CV error and test set error are reported in the table. Also shown are results for nearest shrunken centroids (Tibshirani et al. 2001), using cross-validation to choose the shrinkage parameter. We see that (a) hard thresholding generally improves upon the error rate of the full margin tree; (b) margin trees outperform nearest shrunken centroids on the whole, but not in every case. In some cases, the number of gene used has dropped substantially; to get smaller number of genes one could look more closely at the cross-validation curve, to check how quickly it was rising.

If two genes are correlated and both contribute to classifier, they might both remain in the model, under the above scheme. On the other hand, if there is a set of many highly correlated genes that contribute, their coefficients will be diluted and they might all be removed.

Hence it might be desirable to select among the genes in a more aggressive fashion. There are a number of approaches one might try here, for example the recursive feature elimination of Guyon et al. (2002), mentioned above, One could also try the L1-norm support-vector machine (see e.g. Zhu et al. (2003)), but this is also quite slow to compute. Another approach would be to apply the lasso (Tibshirani 1997). All of these methods would be worth trying; however they also suffer from interpretability issues. In particular, the best classifier with say 50 genes might have only a few genes in common with the best classifier with 100 genes. The hard thresholding method described above does not suffer from this drawback. It gives a single ranked list of weights for all genes, for the classifier at each node of the tree.

5 Discussion

The margin-tree method proposed here seems well suited to high-dimensional problems with more than two classes. It has prediction accuracy competitive with multiclass support vector machines and nearest centroid methods, and provides a hierarchical grouping of the classes.

All of the classifiers considered here use a linear kernel, i.e., they use the original input features. The construction of margin tree could also be done using other kernels, using the support vector machine framework.

In addition to the papers on the multiclass support vector classifier mentioned earlier, there is other work related to our paper. The decision tree methods of Breiman et al. (1984) and Quinlan (1993) use top-down splitting to form a binary tree, but use other criteria (different from the margin) for splitting. Probably the closest paper to our work is that of Vural & Dy (2004), who use a top-down binary tree approach. They use K-means clustering of the raw data to divide the points in two groups at each node, before applying a support vector classifier. Bennett & Blue (1997) investigate decision trees with support vectors classifiers at the node, but do not discuss adaptive construction of the tree topology. Park & Hastie (2005) propose hierarchical classification methods using nearest centroid classifiers at each node. They use clustering methods to find the topology of the tree, and their paper has some ideas in common with this one. In fact, the mixture model used for each merged node gives a decision boundary that is similar to the support vector classifier. However the maximum margin classifier used here seems more natural and the overall performance of the margin tree is better.

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