Classification in DNA microarray experiments

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Outline

- General issues in classification
 - feature selection;
 - distance and standardization;
 - loss function;
 - class representation;
 - polychotomous classification;
 - missing data.
- Performance assessment
 - estimation of error rates;
 - bias-variance trade-off.
- Aggregating classifiers: baggig, boosting, random forests.
- Comparison of classifiers on tumor microarray data.



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Classification

Classification is a **prediction** or **learning** problem in which the variable to be predicted assumes one of K unordered values, $\{c_1, c_2, \ldots, c_K\}$, arbitrarily relabeled as $\{1, 2, \ldots, K\}$ or sometimes $\{0, 1, \ldots, K-1\}$.

The K values correspond to K **predefined** classes, e.g., tumor class, bacteria type.

Classification

Associated with each object are

- a response or dependent variable (class label) $Y \in \{1, 2, ..., K\}$, and
- a set of G measurements which form the feature vector or vector of predictor variables $\mathbf{X} = (X_1, \dots, X_G)$.

The feature vector \mathbf{X} belongs to a feature space \mathcal{X} (e.g. the real numbers \Re^G).

The task is to classify an object into one of the K classes on the basis of an observed measurement $\mathbf{X} = \mathbf{x}$, i.e., predict Y from \mathbf{X} .

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Classification in microarray experiments

- Classification is an important question in microarray experiments, for purposes of classifying biological samples and predicting clinical or other outcomes using gene expression data
 - tumor class: ALL vs. AML, classic vs. desmoplastic medulldastoma;
- response to treatment, survival;
- type of bacterial pathogen, etc.
- Although classification is by no means a new subject in the statistical literature, the large and complex multivariate datasets generated by microarray experiments raise new methodological and computational challenges.

Unsupervised vs. supervised learning

Unsupervised learning. The classes are **unknown** a priori and need to be "discovered" from the data.

a.k.a. cluster analysis; class discovery; unsupervised pattern recognition.

Supervised learning. The classes are **predefined** and the task is to understand the basis for the classification from a set of labeled objects (training or learning set). This information is then used to classify future observations.

a.k.a. discriminant analysis; class prediction; supervised pattern recognition.

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Tumor classification using gene expression data

A reliable and precise classification of tumors is essential for successful diagnosis and treatment of cancer.

Current methods for classifying human malignancies rely on a variety of morphological, clinical, and molecular variables.

In spite of recent progress, there are still uncertainties in diagnosis.

Also, it is likely that the existing classes are heterogeneous and comprise diseases which are molecularly distinct and follow different clinical courses.

Tumor classification using gene expression data

DNA microarrays may be used to characterize the molecular variations among tumors by monitoring gene expression profiles on a genomic scale.

This may lead to a finer and more reliable classification of tumors, and to the identification of marker genes that distinguish among these classes.

Eventual clinical implications include an improved ability to understand and predict cancer survival.

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Gene expression data

Features correspond to the expression measures for different genes; classes correspond to different biological outcomes (e.g. ALL vs. AML, survivor vs. non-survivor) and are labeled by $\{1, 2, \ldots, K\}$.

Gene expression data on G genes (features) for n mRNA samples (observations)

 $\mathbf{x}_i = (x_{i1}, x_{i2}, \dots, x_{iG})$

- gene expression profile / feature vector for sample i

 y_i = tumor class / response for sample i,

 $i=1,\ldots,n$.

May have covariates such as age, sex.

Tumor classification using gene expression data

There are three main types of statistical problems associated with tumor classification:

- 1. the identification of new tumor classes using gene expression profiles **unsupervised learning**;
- 2. the classification of malignancies into known classes **supervised learning**;
- 3. the identification of marker genes that characterize the different tumor classes **feature selection**.

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 $x_{gi} =$ expression measure for gene g in mRNA sample i.

It is assumed that the arrays have already been conormalized.

Classifiers

Classifiers

A classifier or predictor C for K tumor classes is a map from the space \mathcal{X} of gene expression profiles into the integers $\{1, 2, \ldots, K\}$.

That is, a classifier **partitions** the space \mathcal{X} of gene expression profiles into K disjoint and exhaustive subsets, A_1, \ldots, A_K , such that for a sample with expression profile $\mathbf{x} = (x_1, \ldots, x_G) \in A_k$ the predicted class is k.

 $\mathcal{C}: \mathcal{X} \to \{1, 2, \dots, K\}.$

Predicted class for an observation $\mathbf{x} \in A_k$, $\hat{y} = \mathcal{C}(\mathbf{x}) = k$.

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Decision theory

Classification can be viewed as a statistical decision theory problem.

Assume observations are independently and identically distributed (i.i.d.) from an unknown multivariate distribution.

The class k **prior**, or proportion of objects of class k in the population, is denoted as $\pi_k = p(Y = k)$.

Objects in class k have feature vectors with class conditional density $p_k(\mathbf{x}) = p(\mathbf{x}|Y = k)$.

Classifiers are built from past experience, i.e., from observations which are known to belong to certain classes. Such observations comprise the **learning (training) set, LS**

$$\mathcal{L} = \{ (\mathbf{x}_1, y_1), \dots, (\mathbf{x}_n, y_n) \}.$$

 $n_k = \#$ of class k observations.

Classifier built from a learning set $\mathcal{L}: \mathcal{C}(\cdot; \mathcal{L}) : \mathcal{X} \to \{1, 2, \dots, K\}$. Predicted class for an observation $\mathbf{x} \in A_k, \hat{y} = \mathcal{C}(\mathbf{x}; \mathcal{L}) = k$.

The random nature of the learning set \mathcal{L} implies that the prediction $\hat{y} = \mathcal{C}(\mathbf{x}; \mathcal{L})$ can also be viewed as *random* for a fixed value of \mathbf{x} , depending on whether one conditions on the learning set or not.

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Decision theory

When (unrealistically) both π_k and $p_k(\mathbf{x})$ are known, the classification problem has a solution – Bayes rule.

This situation also gives upper bounds on the performance of classifiers in the more realistic setting where these quantities are not known – Bayes risk.

Decision theory

It will be useful to introduce the notion of a loss function. The loss function L(h, l) simply elaborates the loss incurred if a class h case is erroneously classified as belonging to class l.

The **risk function** for a classifier is the expected loss when using it to classify, that is,

$$R(\mathcal{C}) = E[L(Y, \mathcal{C}(\mathbf{X}))] = \sum_{k} E[L(k, \mathcal{C}(\mathbf{X}))|Y = k]\pi_{k}$$
$$= \sum_{k} \int L(k, \mathcal{C}(\mathbf{x}))p_{k}(\mathbf{x})\pi_{k}.$$

N.B. Here, the classifier is *fixed*, i.e., probabilities are *conditional* on the learning set. In other circumstances, the classifier may be viewed as random and \mathbf{x} as fixed.

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Bayes rule

In the unlikely situation that the class conditional densities $p_k(\mathbf{x}) = p(\mathbf{x}|Y = k)$ and class priors π_k are known, let

$$p(k \mid \mathbf{x}) = \frac{\pi_k p_k(\mathbf{x})}{\sum_l \pi_l p_l(\mathbf{x})}$$

denote the **posterior probability** of class k given feature vector \mathbf{x} .

The **Bayes rule** predicts the class of an observation \mathbf{x} by that with highest posterior probability

 $\mathcal{C}_B(\mathbf{x}) = \operatorname{argmax}_k p(k \mid \mathbf{x}).$

The Bayes rule minimizes the total risk under a symmetric loss function - **Bayes risk**.

Decision theory

Typically L(h,h) = 0, and in many cases the loss is symmetric, with L(h,l) = 1, $h \neq l$ – making an error of one type is equivalent to making an error of a different type.

Then, the risk is simply the misclassification rate

$$R(\mathcal{C}) = p(\mathcal{C}(\mathbf{X}) \neq Y) = \sum_{k} \int_{\mathcal{C}(\mathbf{x}) \neq k} p_k(\mathbf{x}) \pi_k$$

However, for some important examples such as diagnosis, the loss function is not symmetric.

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Bayes rule

For a general loss function, the classification rule which minimizes the total risk is

$$C_B(\mathbf{x}) = \operatorname{argmin}_l \sum_{h=1}^{K} L(h, l) p(h \mid \mathbf{x}).$$
(1)

Suitable adjustments can be made for specific loss functions and to accommodate the doubt and outlier classes.

Classifiers as estimators of the Bayes rule

Many classifiers can be viewed as versions of this general rule with particular parametric or non-parametric estimates of $p(k \mid \mathbf{x})$. There are two general paradigms (Friedman, 1996).

1. Direct function estimation approach. Cass conditional probabilities $p(k \mid \mathbf{x})$ are estimated directly based on function estimation methodology such as regression.

- Logistic regression;
- Neural networks;
- Classification trees;
- Projection pursuit;
- Nearest neighbor classifiers.

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Maximum likelihood discriminant rule

The frequentist analogue of the Bayes rule is the **maximum likelihood (ML) discriminant rule**. For known class conditional densities $p_k(\mathbf{x}) = p(\mathbf{x}|Y = k)$, the ML rule predicts the class of an observation \mathbf{x} by that which gives the largest likelihood to \mathbf{x} : $C(\mathbf{x}) = \operatorname{argmax}_k p_k(\mathbf{x})$.

In the case of equal class priors π_k , this amounts to maximizing the posterior class probabilities $p(k|\mathbf{x})$, i.e., the Bayes rule.

Classifiers as estimators of the Bayes rule

2. Density estimation approach. Class conditional densities $p_k(\mathbf{x}) = p(\mathbf{x}|Y = k)$ (and priors π_k) are estimated separately for each class and Bayes' Theorem is applied to obtain estimates of $p(k \mid \mathbf{x})$.

- Gaussian maximum likelihood discriminant rules, a.k.a. discriminant analysis;
- Naive Bayes methods, which approximate class conditional densities $p_k(\mathbf{x})$ by the product of their marginal densities on each feature variable;
- Learning vector quantization;
- Bayesian belief networks.

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Fisher linear discriminant analysis

First applied in 1935 by M. Barnard at the suggestion of R. A. Fisher (1936), Fisher linear discriminant analysis (FLDA) consists of

- 1. finding linear combinations $\mathbf{x} \mathbf{a}$ of the gene expression profiles $\mathbf{x} = (x_1, \dots, x_G)$ with large ratios of between-groups to within-groups sums of squares discriminant variables;
- 2. predicting the class of an observation **x** by the class whose mean vector is closest to **x** in terms of the discriminant variables.

Linear and $q^{uadratic}$ discriminant analysis

Linear and quadratic (in the features \mathbf{x}) discriminant rules arise as Bayes rules or maximum likelihood (ML) discriminant rules when features have Gaussian distributions within each class.

For multivariate normal class densities, i.e., for $\mathbf{X}|Y = k \sim N(\mu_k, \Sigma_k)$, the Bayes rule is

 $\mathcal{C}(\mathbf{x}) = \operatorname{argmin}_{k} \left\{ (\mathbf{x} - \mu_{k}) \Sigma_{k}^{-1} (\mathbf{x} - \mu_{k})' + \log |\Sigma_{k}| - 2 \log \pi_{k} \right\}.$

In general, this is a quadratic rule – Quadratic discriminant analysis – $QDA_{.}$

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Linear and $q^{uadratic}$ discriminant analysis

1. Linear discriminant analysis – LDA. When the class densities have the same covariance matrix, $\Sigma_k = \Sigma$, the discriminant rule is based on the square of the Mahalanobis distance and is linear in **x** and given by

 $\mathcal{C}(\mathbf{x}) = \operatorname{argmin}_k (\mathbf{x} - \mu_k) \Sigma^{-1} (\mathbf{x} - \mu_k)'.$

Linear and $q^{uadratic}$ discriminant analysis

The main quantity in the discriminant rule is

 $(\mathbf{x} - \mu_k) \Sigma_k^{-1} (\mathbf{x} - \mu_k)'$, the squared Mahalanobis **distance** from the observation \mathbf{x} to the class k mean vector μ_k .

Thus, intuitively, the predicted class for an observation \mathbf{x} is the class with closest mean vector μ_k , for a suitably defined distance function.

Interesting special cases are described below for homogeneous priors, i.e., π_k constant in k.

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Linear and $q^{uadratic}$ discriminant analysis

2. Diagonal linear discriminant analysis – DLDA. When the class densities have the same diagonal covariance matrix $\Sigma_k = \Delta = \text{diag}(\sigma_1^2, \dots, \sigma_G^2)$, the discriminant rule is linear and given by

$$\mathcal{C}(\mathbf{x}) = \operatorname{argmin}_k \sum_{g=1}^G \frac{(x_g - \mu_{kg})^2}{\sigma_g^2}.$$

Naive Bayes rule for Gaussian class conditional densities.

3. Nearest centroid. In this simplest case, it is assumed that $\Sigma_k = I_G$, the $G \times G$ identity matrix, and observations are classified on the basis of their Euclidean distance from class means μ_k .

Linear and quadratic discriminant analysis

For the

sample Bayes or ML discriminant rules, the population mean vectors and covariance matrices are estimated from a learning set \mathcal{L} , by the sample mean vectors and covariance matrices, respectively: $\hat{\mu}_k = \bar{\mathbf{x}}_k$ and $\hat{\Sigma}_k = S_k$.

For the constant covariance matrix case, the pooled estimate of the common covariance matrix is used:

 $\hat{\Sigma} = S = \sum_{k} (n_k - 1) S_k / (n - K).$

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Weighted gene voting of Golub et al. (1999)

This is a minor variant on **DLDA** for two classes, k = 1, 2.

Sample DLDA classifies an observation \mathbf{x} as 1 iff

$$\sum_{g=1}^{G} \frac{(\bar{x}_{1g} - \bar{x}_{2g})}{\hat{\sigma}_{g}^{2}} \left(x_{g} - \frac{(\bar{x}_{1g} + \bar{x}_{2g})}{2} \right) \ge 0$$

The discriminant function can be rewritten as $\sum_g v_g$, where $v_g = a_g(x_g - b_g)$, $a_g = (\bar{x}_{1g} - \bar{x}_{2g})/\hat{\sigma}_g^2$, and $b_g = (\bar{x}_{1g} + \bar{x}_{2g})/2$.

In Golub et al., $a_g = (\bar{x}_{1g} - \bar{x}_{2g})/(\hat{\sigma}_{1g} + \hat{\sigma}_{2g}) \dots$ (Wrong units).

Linear and quadratic discriminant analysis

The above simple rules may be modified easily to allow for unequal class priors.

Estimates of the priors may, in some cases, be obtained from the sample class proportions $\hat{\pi}_k = n_k/n.$

As with any classifier explicitly estimating the Bayes rule, class posterior probabilities may be used to assess the confidence in the predictions for individual observations.

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Linear and quadratic discriminant analysis

The linear discriminant rules described above are classical and widely used classification tools.

- Simple and intuitive: the predicted class of a test case is the class with the closest mean (using the Mahalanobis metric).
- Estimated Bayes rule: LDA is based on the estimated Bayes rule for Gaussian class conditional densities optimality *when* the model is true.
- Easy to implement: the partition has linear boundaries.
- Good performance in practice: in spite of a possibly high bias, the low variance of the LDA estimates of class posterior probabilities often results in low classification error (see discussion of bias-variance trade-off).

Extensions to LDA

Linear and $q^{uadratic}$ discriminant analysis

However, LDA has a number of obvious limitations which are due in part to its simplicity.

- Linear or even quadratic discriminant boundaries may not be flexible enough.
- Features may have mixture distributions within classes.
- In the case of too many features, performance may degrade rapidly due to over-parameterization and high variance of parameter estimates.

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Extensions to LDA

Mixture discriminant analysis, MDA. Gass conditional densities are modeled as mixtures of Gaussian densities, with different mean vectors but the same covariance matrices. The EM algorithm may be used for maximum likelihood estimation of the parameters of the Gaussian components.

Automatic feature selection. Shrinkage methods can be applied to perform automatic feature selection. For example, the class mean vectors can be shrunken using matrices of between-groups and within-groups sums of squares. Parameters controlling the amount of shrinking can be selected by cross-validation. Ripley (1996) and Hastie et al. (2001) discuss various extensions to LDA to address these issues.

Flexible discriminant analysis, FDA. Viewing LDA as a linear regression problem, suggests considering more general non-parametric forms of regression for building a more flexible classifier. FDA amounts to performing LDA on transformed responses and/or features. See logistic discrimination.

Penalized discriminant analysis, PDA. A penalized Mahalanobis distance is used to enforce smoothness of the within-class covariance matrix of the features (possibly transformed as in FDA).

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Logistic discrimination

For Gaussian class conditional densities with common covariance matrix (and other models)

$\log p(k|\mathbf{x}) - \log p(1|\mathbf{x}) = \alpha_k + \mathbf{x}\beta_k.$

This suggests modeling $\log p(k|\mathbf{x}) - \log p(1|\mathbf{x})$ more generally by some parametric family of functions, say $g_k(\mathbf{x};\theta)$ (with $g_1(\mathbf{x};\theta) \equiv 0$). Estimates of the class posterior probabilities are then given by

$$\hat{p}(k|\mathbf{x}) = rac{\exp g_k(\mathbf{x};\hat{ heta})}{\sum_l \exp g_l(\mathbf{x};\hat{ heta})}.$$

Classification is done by the (estimated) Bayes rule, i.e., $C(\mathbf{x}; \mathcal{L}) = \operatorname{argmax}_k \hat{p}(k|\mathbf{x}).$

Logistic discrimination

In the machine learning literature the function $\exp a_k / \sum_l \exp a_l$ is known as the **softmax** function, in the earlier statistical literature, it is known as the **multiple logit** function.

In the linear case, one takes $g_k(\mathbf{x}; \theta) = \alpha_k + \mathbf{x}\beta_k$.

Logistic regression models are typically fit by maximum likelihood, using a Newton–Raphson algorithm known as **iteratively reweighted least squares** (IRLS).

Logistic discrimination provides a more direct way of estimating posterior probabilities and is easier to generalize than classical linear discriminant analysis (LDA and QDA), e.g. neural networks.

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Nearest neighbor classifiers

Note that, for a large enough number of neighbors k, k-nearest neighbor classifiers suggest simple estimates of the class posterior probabilities: the proportion of votes for each class.

The class posterior probability estimates $\hat{p}(l|\mathbf{x})$ may be used to measure confidence for individual predictions.

For k = 1, the nearest neighbor partition of the feature space \mathcal{X} corresponds to the **Dirichlet tessellation** of the learning set \mathcal{L} .

Classifiers with k = 1 are generally quite successful.

Nearest neighbor classifiers

Nearest neighbor methods are based on a measure of **distance** between observations, such as the Euclidean distance or one minus the correlation between two gene expression profiles.

The k-nearest neighbor rule, k-NN, due to Fix and Hodges (1951), classifies an observation x as follows

- 1. find the k observations in the learning set that are closest to \mathbf{x} ;
- 2. predict the class of \mathbf{x} by majority **vote**, i.e., choose the class that is most common among those k neighbors.

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Nearest neighbor classifiers: which k?

In general, the number of neighbors k can be chosen by cross-validation.

Each observation in the learning set is treated in turn as if it were in the test set: its distance to all of the other learning set cases (except itself) is computed and it is classified by the nearest neighbor rule.

The classification for each learning set observation is then compared to the truth to produce the cross–validation error rate.

This is done for a number of k's and the k for which the cross-validation error rate is smallest is retained.

Extensions of nearest neighbor rule

Nearest neighbor classifiers

Nearest neighbor classifiers were initially proposed by Fix and Hodges (1951) as consistent non–parametric estimates of maximum likelihood discriminant rules.

Non-parametric estimates of the class conditional densities $p_k(\mathbf{x})$ are obtained by first reducing the dimension of the feature space \mathcal{X} from G to one using a distance function.

The proportions of neighbors in each class are then used in place of the corresponding class conditional densities in the maximum likelihood discriminant rule.

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Extensions of nearest neighbor rule

Distance weights. The standard k-nearest neighbor rule equally weights all k neighbors, regardless of their distance from the test case. A more sensitive rule may be obtained by assigning weights to the neighbors that are inversely proportional to their distance from the test sample – somewhat controversial.

Differential misclassification costs. The minimum vote majority may be adjusted based on the class to be called. This corresponds to weighted voting by the neighbors, where weights are suggested by equation (1).

E.g. When $L(h, l) = L_h I(h \neq l)$, the weights are given by $w_l = L_l$.

The nearest neighbor rule can be refined to deal with issues of unequal class priors, differential misclassification costs, and feature selection. Many of these refinements involve some form of **weighted voting** for the neighbors.

Class priors. Suppose the class prior probabilities π_l are known and the learning set class proportions $\hat{\pi}_l = n_l/n$ are different from the π_l .

- Votes may be weighted according to neighbor class, $w_l = \pi_l/n_l$.
- The distance to class l cases may be weighted by $(n_l/\pi_l)^G$ for a G-dimensional feature space (Brown & Koplowitz, 1979).

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Extensions of nearest neighbor rule

Feature selection. One of the most important issues in k-nearest neighbor classification is the choice of a distance function and the selection of relevant features.

For instance, in the context of gene expression data, a large number of genes are constantly expressed across classes, thus resulting in a large number of irrelevant or noise variables. Inclusion of features with little or no relevance can substantially degrade the performance of the classifier.

Feature selection may be implemented automatically, by modifying the distance function used by the classifier, or by using classification trees.

Extensions of nearest neighbor rule

Friedman (1994). Flexible metric nearest neighbor classification approach in which the relevance of each feature (or linear combination of features) is estimated locally for each test case. The proposed procedure, the machette, is a hybrid between classical nearest neighbor classifiers and tree–structured recursive partitioning techniques.

Hastie & Tibshirani (1996). In the discriminant adaptive nearest neighbor procedure, DANN, the distance function is based on local discriminant information.

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Classification trees

Binary tree structured classifiers are constructed by repeated splits of subsets (nodes) of the measurement space \mathcal{X} into two descendant subsets, starting with \mathcal{X} itself. Each terminal subset is assigned a class label and the resulting partition of \mathcal{X} corresponds to the classifier.

Three main aspects of tree construction: (i) the selection of the splits; (ii) the decision to declare a node terminal or to contin^{ue} splitting; (iii) the assignment of each terminal node to a class.

Different tree classifiers use different approaches to deal with these three issues. Here, we use **CART – Classification And Regression Trees** – of Breiman et al. (1984).

Extensions of nearest neighbor rule

Buttrey & Karo (2002). Hybrid or composite classifier, k-NN-in-leaf, which partitions the feature space using a classification tree and classifies test set cases using a standard k-nearest neighbor rule using only training cases in the same leaf as the test case.

Each k-NN classifier may have a different number of neighbors k, a different set of feature variables, and a different choice of scaling.

The k-NN-in-leaf classifier differs from Friedman's machette in that only one tree is ever grown.

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Classification trees

1. **Splitting rule.** At each node, choose the split that maximizes the decrease in impurity.

E.g. of impurity functions:

Gini index $\phi(p_1, \ldots, p_K) = \sum_{k \neq l} p_k p_l = 1 - \sum_k p_k^2$, entropy, and twoing rule.

- 2. **Split-stopping rule.** Grow a large tree, selectively **prune** the tree upward, getting a decreasing sequence of subtrees. Use **cross-validation** to identify the subtree having the lowest estimated misclassification rate.
- 3. Class assignment rule. For each terminal node, choose the class that minimizes the resubstitution estimate of the misclassification probability, given that a case falls into this node.

Refinements: class priors, loss function, splits based on linear combinations of variables, missing values (surrogate splits).

Support vector machines

When the optimization problem is reformulated in terms of Lagrangians, the feature vectors ${\bf x}$ only appear in the objective function via dot products ${\bf x_i}\cdot {\bf x_j}.$

For a cost parameter C, building the classifier involves maximizing the dual

$$L_D = \sum_i \alpha_i - \frac{1}{2} \sum_{i,j} \alpha_i \alpha_j y_i y_j \mathbf{x_i} \cdot \mathbf{x_j}$$

subject to $0 \le \alpha_i \le C$ and $\sum_i \alpha_i y_i = 0, \forall i$. The solution is given by

$$\mathbf{w} = \sum_{i=1}^{N_S} \alpha_i y_i \mathbf{x_i},$$

where N_S is the number of support vectors.

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Support vector machines

User-specified parameters for the SVM C-classification method: the type of kernel K, i.e., non-linear transformation of the original data, and the cost parameter C for misclassifications when the training data are non-separable in the transformed space.

| Classifier | Kernel |
|--------------------------------------|--|
| Polynomial of degree p | $K(\mathbf{x}, \mathbf{y}) = (\mathbf{x} \cdot \mathbf{y} + 1)^p$ |
| Gaussian radial basis function (RBF) | $K(\mathbf{x}, \mathbf{y}) = \exp(- \mathbf{x} - \mathbf{y} ^2 / 2\sigma^2)$ |
| Two–layer sigmoidal neural network | $K(\mathbf{x}, \mathbf{y}) = \arctan(\kappa^{\mathbf{x} \cdot \mathbf{y}} - \delta)$ |

SVMs are designed for binary outcomes. They can be generalized to multiclass problems by solving several binary problems simultaneously (see below).

Support vector machines

Support vector machines, SVM, were introduced by Vapnik (1979, 1998).

For binary classification (-1 vs. 1), the main idea is to find the *best* hyperplane $\mathbf{w} \cdot \mathbf{x} + b = 0$ separating the classes in the learning set.

What is best? Maximize the **margin**, i.e., the sum of the distances from the hyperplane to the closest positive and negative correctly classified samples, while penalizing for the number of misclassifications.

One can search for the hyperplane in the original space, linear **SVMs**, or in a higher–dimensional space, **non–linear SVMs**.

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Support vector machines

In many situations it is useful to consider **non-linear** (in \mathbf{x}) decision functions or class boundaries. Mapping the data into higher dimensional spaces and then reducing the problem to the linear case leads to a simple solution.

Suppose the data are mapped from R^G to \mathcal{H} using $\Phi(\cdot)$, then the training algorithm depends on the data only through the dot product $\Phi(\mathbf{x_i}) \cdot \Phi(\mathbf{x_j})$.

If there is a **kernel function** K such that $K(\mathbf{x_i}, \mathbf{x_j}) = \Phi(\mathbf{x_i}) \cdot \Phi(\mathbf{x_j})$, then one does not even need to know $\Phi(\cdot)$ explicitly.

Classifier partitions

The following figures compare the partitions produced by different classifiers: LDA, QDA, k-NN, and CART.

The classifiers were applied to the brain tumor MD survival dataset of Pomeroy et al. (2002) using only the two genes with the largest absolute t-statistics.

Predicted responses "survivor" and "non–survivor" are indicated by shades of red and blue, respectively.

The entire learning set of n = 60 samples was used to build the classifiers, the resubstitution error rate is shown below the plots.

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Figure 2: *Brain tumor MD survival dataset.* QDA partition for the two genes with the largest absolute *t*-statistics.



Example: Linear discriminant analysis

Figure 1: Brain tumor MD survival dataset. LDA partition for the two genes with the largest absolute t-statistics.

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Figure 3: Brain tumor MD survival dataset. k-NN partitions (k = 1, 3, 5, 11) for the two genes with the largest absolute t-statistics



Figure 4: Brain tumor MD survival dataset. CART (10-fold CV) partition for the two genes with the largest absolute t-statistics.

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Other classifiers

- Neural networks;
- Learning vector quantization (LVQ);
- Bayesian belief networks;
- etc.



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In contrast, standard DA and nearest neighbor classifiers do not perform feature selection; all variables, whether relevant or not, are used in building the classifier.

Explicit feature selection

Multivariate approaches. More refined feature selection procedures consider the *joint* distribution of the expression measures, in order to detect genes with weak main effects but possibly strong interactions.

Bo & Jonassen (2002): **subset selection** procedures for screening gene pairs to be used in classification.

Breiman (1999): select genes according ^{to} importance statistics which are defined in terms of prediction accuracy (see random forests below).

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Feature selection and performance assessment

The importance of taking feature selection into account when assessing the performance of the classifier cannot be stressed enough.

Feature selection *is* an aspect of building the classifier, whether done explicitly or implicitly.

Thus, when using for example cross-validation to estimate generalization error, feature selection should be done *not* on the entire learning set, but separately for each cross-validation sample used to build the classifier.

Leaving out feature selection from cross-validation or other resampling-based performance assessment method results in overly optimistic error rates.

Explicit feature selection

One–gene–at–a–time approaches. Genes are ranked based on the value of a univariate test statistic such as: t– or F-statistic; non–parametric Wilcoxon or Kruskal–Wallis statistic; p–value.

Possible meta-parameters include the number of genes G or a p-value cut-off. A formal choice of these parameters may be achieved by cross-validation or bootstrap procedu^{res}.

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Implicit feature selection

Feature selection may also be performed *implicitly* by the classification rule itself.

In classification trees, features are selected at each step based on reduction in impurity and the number of features used (or size of the tree) is determined by pruning the tree using cross-validation. Thus, feature selection is an inherent part of tree building and pruning deals with the issue of over-fitting.

Shrinkage methods and adaptive distance functions may be used for linear discriminant analysis and nearest neighbor classification.

Distance and standardization

Classification trees. Trees are invariant under monotone transformations of individual features (genes), e.g. standardization. They are not invariant to standardization of the observations (normalization in the microarray context).

Linear and quadratic discriminant analysis. Based on the Mahalanobis distance of the observations from the class means. Thus, the classifiers are invariant to standardization of the variables (genes), but not the observations (arrays).

Nearest neighbor classifiers. One must explicitly decide on an appropriate standardization and distance function for the problem under consideration. These classifiers are in general affected by standardization of both features and observations.

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Unequal sample class representation

In many situations, such as medical diagnosis, the representation of the classes in the learning set does not reflect their importance in the problem.

For example, in a binary classification problem with a rare disease class (d) and a common healthy class (h), a learning set obtained by random sampling from the population would contain a vast majority of healthy cases.

Unequal class sample sizes could possibly lead to serious biases in the estimation of posterior probabilities $p(\mathbf{d} \mid \mathbf{x})$ and $p(\mathbf{h} \mid \mathbf{x})$.

Distance and standardization

See lecture "Distances and expression measures" for more detail.

Both the **distance** function and the **scale** the observations are measured in can have a large impact on the performance of the classifier, however, the effect varies depending on the classifier.

The choice of transformation and distance function should thus be made jointly and in conjunction with the choice of classifier.

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Loss function

In many diagnosis settings, the loss incurred from misclassifying a diseased (d) person as healthy (h) far outweighs the loss incurred by making the error of classifying a healthy person as diseased.

These differential misclassification costs should be reflected in the loss function.

Suppose the loss from the first error is e > 1 times higher than that from the second. In the case of the Bayes rule, one could modify the posterior probability cut-offs and classify a patient as diseased if $p(\mathbf{d} \mid \mathbf{x}) > c = 1/(1+e)$. Such a classifier would minimize the risk for the loss function $L(\mathbf{d}, \mathbf{h}) = eL(\mathbf{h}, \mathbf{d})$.

Unequal sample class representation

Consider the case of linear discriminant analysis which assumes a common covariance matrix estimated using both samples. The pooled estimate of the covariance matrix will be dominated by the more abundant sample.

This is fine if the covariance matrix is really the same for both classes. However, in the case of unequal covariance matrices, the bias in estimating class posterior probabilities $p(k \mid \mathbf{x})$ is more severe when the classes are unequally represented in the learning set.

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Biased sampling of classes

Downweighting and subsampling are helpful in dealing with the estimation bias for densities $p_k(\mathbf{x})$, however they effectively make the sample proportions differ from the population proportions. This leads to biased estimates of the class posterior probabilities $p(k|\mathbf{x})$.

The plug-in estimators of class posterior probabilities are in fact estimating quantities proportional to $p(k \mid \mathbf{x})n_k/\pi_k$.

Adjustment is thus required to ensure that estimators of the class posterior probabilities $p(k \mid \mathbf{x})$ are approximately unbiased.

This can be done by specifying appropriate priors for DA and CART, and by using weighted voting for nearest neighbors.

Biased sampling of classes

Approaches to alleviate estimation biases

- Subsample the abundant population so that both classes are on an equal footing in the parameter estimation. Problem: this would be wasteful of training data.
- Downweight cases from the abundant class so that the sum of the weights is equal to the number of cases in the less abundant clas.
- Obtain less biased estimates of the class posterior probabilities, either by theory (as can be done for LDA) or via bias reduction techniques such as the jackknife.

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Polychotomous classification

It may be advantageous to convert a polychotomous or K-class classification problem into a series of binary or two-class problems (e.g. for a large number of classes with unequal representation in the learning set).

- All pairwise binary classification problems. Consider all $\binom{K}{2}$ binary classification problems; the final predicted class is the class that is selected most often as the predicted class or winner in the $\binom{K}{2}$ decisions.
- One-against-all binary classification problems. Application of K binary rules to a test case x yields estimates of class probabilities; the final K-class decision rule selects the class withlargest estimated posterior probability.

Performance assessment

Missing data

Some classifiers are able to handle missing values by performing automatic imputation (e.g. trees), while others either ignore missing data or require imputedata (e.g. LDA).

Classification trees can readily handle missing values through the use of

surrogate splits, i.e., splits that are most similar to the best split at a particular node.

For microarray data, simple weighted **nearest neighbor imputation** was found to provide accurate and robust estimates of missing values (Troyanskaya et al., 2001).

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Performance assessment

1. Resubstitution estimation. Error rate on the learning set.

Problem: can be severely biased downward.

2. Test set estimation. In the absence of a genuine test set, cases in the learning set \mathcal{L} may be repeatedly randomly divided into two sets, \mathcal{L}_1 and \mathcal{L}_2 ; the classifier is built using \mathcal{L}_1 and the error rate is computed for \mathcal{L}_2 .

Problem: reduces effective sample size; no widely accepted guidelines for choosing the relative size of these artificial learning sets and test sets. Virtually every application of classification methods to microarray data includes some discussion of the performance of the classifier(s).

The ability to predict biological outcomes using gene expression measures is usually evaluated based on estimates of the **classification error**.

The reported error rates are often biased downward and give an overly optimistic view of the predictive power of expression data.

Obtaining accurate or *honest* estimates of classification error is a very important and delicate problem.

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Performance assessment

3. V-fold cross-validation (CV) estimation.

Cases in the learning set \mathcal{L} are randomly divided into Vsubsets \mathcal{L}_v , $v = 1, \ldots, V$, of as nearly equal size as possible. Classifiers are built on learning sets $\mathcal{L} - \mathcal{L}_v$, test set error rates are computed for \mathcal{L}_v , and averaged over v.

Bias-variance trade-off: small Vs typically give a larger bias, but a smaller variance and mean squared error.

Performance assessment

Performance assessment

- 4. Leave-one-out cross-validation (LOOCV). Special case for V = n.
 - In general low bias but high variance.
 - For stable (low variance) classifiers such as k–NN, LOOCV provides good estimates of generalization error rates.
 - Computationally intensive for large n.
- 5. **Out–of–bag estimation.** See discussion of random forests below.

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Performance assessment

However, in most cases, the important features are unknown and the intended inference includes feature selection.

Then, CV estimates as above tend to suffer from a **downward bias** and inference is not warranted.

Features should be selected only on the basis of the samples in $\mathcal{L} - \mathcal{L}_v$ for CV estimation.

The use of cross-validation (or any other estimation method) is intended to provide accurate estimates of classification error rates.

It is important to note that these estimates relate **only** to the experiment that was (cross–) validated.

There is a common practice in microarray classification of doing feature selection using *all* of the learning set and then using cross–validation only on the classifier building portion of the process.

In that case, inference can only be applied to the latter portion of the process.

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Performance assessment

The above remarks apply to other error rate estimation methods. E.g. test set and out–of–bag estimation.

They also apply to other aspects of the classifier training process. E.g. choice of the number of neighbors k for k-NN, choice of kernel K and cost parameter C in SVMs.

Performance assessment

In the case of unequal representation of the classes, some form of stratified sampling may be needed to ensure balance across important classes in all subsamples.

In addition, for complex experimental designs, such as factorial or time–course designs, the resampling mechanisms used for computational inference should reflect the design of the experiment.

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Bias, variance, and error rates

For simplicity, consider binary classification, i.e., K = 2 and $Y \in \{-1,1\}$, and let $f(\mathbf{x}) = p(1|\mathbf{x}) = p(Y = 1|\mathbf{X} = \mathbf{x})$ denote the posterior probability for class 1.

Consider predicted classes $C(\mathbf{x}; \mathcal{L})$ based on estimates $\hat{f}(\mathbf{x}; \mathcal{L})$ of the class posterior probabilities $f(\mathbf{x})$.

Denote the **mean** and **variance** of the estimators $\hat{f}(\mathbf{x}; \mathcal{L})$ by

$$\mu_{\mathbf{x}} = E[f(\mathbf{x}; \mathcal{L})],$$

$$\sigma_{\mathbf{x}}^2 = E[(\hat{f}(\mathbf{x}; \mathcal{L}) - \mu_{\mathbf{x}})^2]$$

Bias, variance, and error rates

The random nature of the learning set \mathcal{L} implies that for any realization \mathbf{x} of the feature vector, the predicted class $\hat{Y} = \mathcal{C}(\mathbf{x}; \mathcal{L})$ is a random variable.

Intuitively, for a fixed value of the feature vector \mathbf{x} , as the learning set varies, so will the predicted class $\hat{Y} = \mathcal{C}(\mathbf{x}; \mathcal{L})$.

It is thus meaningful and instructive to consider distributional properties (e.g. bias and variance) of classifiers when assessing and comparing the performance of different classifiers.

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Bias and variance – Estimation error

The mean squared estimation error, averaged over learning sets \mathcal{L} , is

$$MSE_{\mathbf{x}} = E[(\hat{f}(\mathbf{x};\mathcal{L}) - f(\mathbf{x}))^2] = \sigma_{\mathbf{x}}^2 + (\mu_{\mathbf{x}} - f(\mathbf{x}))^2$$

= Variance + Bias². (2)

Recall again that here \mathbf{x} is fixed and the randomness comes from the learning set \mathcal{L} .

Bias and variance – Classification error

Now consider the effect of $\mu_{\mathbf{x}}$ and $\sigma_{\mathbf{x}}^2$ on classification error (Friedman, 1996). The **Bayes rule** is fixed and independent of \mathcal{L} , and for binary classification problems it is given by

$$y_B = \begin{cases} 1, & \text{if } f(\mathbf{x}) \ge 1/2 \\ -1, & \text{if } f(\mathbf{x}) < 1/2 \end{cases}.$$

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Bias and variance – Classification error

Following Friedman (1996), use a normal approximation for the distribution of $\hat{f}.$ Then

$$p(\hat{Y} \neq y_B | \mathbf{X} = \mathbf{x}) = \Phi\left(\operatorname{sign}(f(\mathbf{x}) - 1/2) \; \frac{1/2 - \mu_{\mathbf{x}}}{\sigma_{\mathbf{x}}}\right)$$

where $\Phi(\cdot)$ is the standard normal cdf. Hence, the classification error rate is

$$p(\hat{Y} \neq Y | \mathbf{X} = \mathbf{x}) = |2f(\mathbf{x}) - 1| \Phi \left(\operatorname{sign}(f(\mathbf{x}) - 1/2) \frac{1/2 - \mu_{\mathbf{x}}}{\sigma_{\mathbf{x}}} \right) + p(Y \neq y_B | \mathbf{X} = \mathbf{x}).$$
(4)

Bias and variance – Classification error The classification error is $p(\hat{Y} \neq Y | \mathbf{X} = \mathbf{x}) = p(\hat{Y} = -1 | \mathbf{X} = \mathbf{x}) p(Y = 1 | \mathbf{X} = \mathbf{x}) + p(\hat{Y} = 1 | \mathbf{X} = \mathbf{x}) p(Y = -1 | \mathbf{X} = \mathbf{x})$ $= p(\hat{Y} = -1 | \mathbf{X} = \mathbf{x}) f(\mathbf{x}) + p(\hat{Y} = 1 | \mathbf{X} = \mathbf{x}) (1 - f(\mathbf{x}))$ $= |2f(\mathbf{x}) - 1| p(\hat{Y} \neq y_B | \mathbf{X} = \mathbf{x}) + p(Y \neq y_B | \mathbf{X} = \mathbf{x})$ = Estimation error + Bayes error rate.(3)

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Bias, variance, and error rates

Comparison of equations (2) and (4) shows that moments $\mu_{\mathbf{x}}$ and $\sigma_{\mathbf{x}}^2$ of the distribution of $\hat{f}(\mathbf{x}; \mathcal{L})$ have a very different impact on estimation error (for $f(\mathbf{x})$) and on classification error.

The **bias-variance trade-off** is very different for these two types of error: for **estimation error**, the dependence on bias and variance is **additive**; for **classification error**, there is an **interaction** effect.

Variance tends to dominate bias for classification error. This suggests that certain methods with high bias for function estimation may nonetheless perform well for classification because of their low variance. E.g. Naive Bayes (DLDA) and nearest neighbors.

Aggregating classifiers

Breiman (1996, 1998) found that gains in accuracy could be obtained by **aggregating predictors** built from perturbed versions of the learning set. In classification, the multiple versions of the predictor are aggregated by **voting**.

Let $\mathcal{C}(\cdot; \mathcal{L}_b)$ denote the classifier built from the *b*th perturbed learning set \mathcal{L}_b and let w_b denote the weight given to predictions made by this classifier. The predicted class for an observation **x** is given by

 $\operatorname{argmax}_k \sum_{b} w_b I(\mathcal{C}(\mathbf{x}; \mathcal{L}_b) = k).$

Key to improved accuracy: instability of the classifier, i.e., variance.

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Random forests

A **random forest** is a collection of tree classifiers, where each tree depends on the value of a random vector, i.i.d. for all trees in the forest (Breiman, 1999).

- Random learning set bagging. Each tree is formed from a bootstrap sample of the learning set the random vector consists of the outcomes of n draws at random with replacement from $\{1, \ldots, n\}$.

Bagging

Standard bagging. In the simplest form of **bagging** – bootstrap **agg**regating – perturbed learning sets are non–parametric bootstrap replicates of the learning set, i.e., are drawn at random with replacement from the learning set.

Predictors are built for each perturbed dataset and aggregated by

plurality voting $(w_b = 1)$.

Parametric bootstrap. Perturbed learning sets can be generated according to a mixture of multivariate Gaussian distributions.

Convex pseudo-data. Breiman (1996).

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Random forests

The above two sources of randomness are combined.

A maximal exploratory tree is grown (pure terminal nodes) for each bootstrap learning set, and the forest obtains a classification by plurality voting.

Note that the main ideas in random forests are applicable to other types of classifiers than trees.

By–products of aggregation

I. Out-of-bag estimate of error rate. No need for CV or test set estimation of error rate; an unbiased estimate is obtained as a by-product of the bootstrap.

For each bootstrap sample, about 1/3 of the cases are left out and not used in the construction of the tree \Rightarrow **out-of-bag** test set.

For the *b*th bootstrap sample, put the out–of–bag cases down the *b*th tree to get a test set classification. Let the final test set classification of the forest be the class having the most votes. Compare this classification to the class labels of the learning set to get the out–of–bag estimate of the error rate.

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By–products of aggregation

III. Variable importance statistics. Here, variable **importance** is defined in terms of the contribution to **prediction accuracy**, i.e., predictive power.

For each tree, randomly permute the values of the gth variable for the out–of–bag cases, put these new covariates down the tree, and get new classifications for the forest.

By–products of aggregation

- II. Case-wise information.
 - Prediction votes. The proportions of votes for each class can be viewed as estimates of the class posterior probabilities (∈ [0,1]).

The prediction vote for the winning class gives a measure of confidence for the prediction of individual observations.

• Vote margins. Vote margins are defined as the proportion of votes for the true class minus the maximum of the proportions of votes for each of the other classes ($\in [-1,1]$). Low vote margins may indicate that the corresponding samples are hard to predict or, in some cases, they may reflect mislabeled learning set cases.

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By–products of aggregation

The importance of the gth variable can be defined in a number of ways.

- *Importance measure 1:* the difference between the out–of–bag error rate for randomly permuted *g*th variable and the original out–of–bag error rate.
- Importance measure 2: the average across all cases of the differences between the margins for the randomly permuted gth variable and for the original data.
- *Importance measure 3:* the number of lowered margins minus the number of raised margins.
- *Importance measure 4:* the sum of all decreases in impurity in the forest due to the *g*th variable, normalized by number of trees.

A da Boost

- **Initialization.** The resampling probabilities $\{p_1^{(0)}, \ldots, p_n^{(0)}\}$ are initialized to $p_i^{(0)} = 1/n$.
 - 2. AdaBoost iterations. For $1 \le b \le B$

- Clustering:

same class).

- Multidimensional scaling;

- (a) Using the current resampling probabilities $\{p_1^{(b-1)}, \ldots, p_n^{(b-1)}\}$, sample with replacement from \mathcal{L} to get a learning set \mathcal{L}_b of size n.
 - (b) Build a classifier $C(\cdot; \mathcal{L}_b)$ based on \mathcal{L}_b .
- (c) Run the learning set L through the classifier C(: L_b) and let d_i = 1 if the *i*th case is classified incorrectly and d_i = 0 otherwise.

(d) Define

$$\epsilon_b = \sum_i p_i^{(b-1)} d_i$$
 and $\beta_b = (1 - \epsilon_b)/\epsilon_b$,

By-products of aggregation

• IV. Intrinsic proximities between cases. Proportion of trees for which cases *i* and *j* are in the same terminal node.

- Outlier detection: 1/(sum of squared proximities of cases in

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and update the resampling probabilities for the (b+1)st step by

$$= \frac{p_{i}^{(b-1)}\beta_{b}^{d_{i}}}{\sum_{i}p_{i}^{(b-1)}\beta_{b}^{d_{i}}}.$$

 $p_i^{(b)}$

- (e) In the event that $\epsilon_b \ge 1/2$ or $\epsilon_b = 0$, the resampling probabilities are reset to be equal.
- 3. Final classification. After B steps, the classifiers $C(: \mathcal{L}_1), \ldots, C(: \mathcal{L}_B)$ are aggregated by weighted voting, with $C(: \mathcal{L}_b)$ having weight $w_b = \log(\beta_b)$.

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Boosting

The data are **resampled adaptively** so that the weights in the resampling are increased for those cases most often misclassified.

The aggregation of predictors is done by weighted voting.

AdaBoost – Freund and Schapire (1997), Breiman (1998).

LogitBoost – Friedman et al. (2000), Dettling & Buhlmann (2002).

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Comparison study – classifiers

- Linear and quadratic discriminant analysis
 - Fisher linear discriminant analysis (FLDA);
 - Diagonal linear discriminant analysis (DLDA)
 - \rightarrow gene voting scheme of Golub et al. is a variant of DLDA;
 - Diagonal quadratic discriminant analysis (DQDA).
- Nearest neighbor classifiers.
- Classification trees (CART).
- SVMs.
- Baggig, boosting, random forests.

N.B. DLDA and DQDA are a.k.a. naive Bayes classifiers.

Ref. Dudoit & Fridlyand (2002), Dudoit et al. (2002).

Comparison study – datasets

- Lymphoma. Alizadeh et al. (2000) (cDNA arrays). n = 81 samples, G = 4,682 genes, 3 classes (B-CLL, FL, DLBCL).
- Leukemia. Golub et al. (1999) (Affymetrix chips). n = 72 samples, G = 3,571 genes, 3 classes (B–cell ALL, T–cell ALL, AML).
- NCI 60. Ross et al. (2000) (cDNA arrays). *n* = 64 samples, *G* = 5, 244 genes, 8 classes.

discussion

Comparison study

- Brain cancer. Pomeroy et al. (2002) (Affymetrix chips). Classic vs. desmoplastic MD: n = 34 samples, G = 5,893 genes. Survivor vs. non-survivor: n = 60 samples, G = 4,459 genes.
- Breast cancer. West et al. (2001) (Affymetrix chips).
 ER +ve vs. ER -ve: n = 49 samples, G = 7, 129 genes.
 Nodal +ve vs. nodal -ve: n = 49 samples, G = 7, 129 genes.

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aggregated classifiers: error "correlated" LDA: ignoring Unlike classification trees and nearest neighbors, intuitive classifiers, their main limitation is that Classification trees are capable of handling and improvement in the performance of aggregated they give very little insight into mechanisms classifiers and to gain more insight into the rates, prediction votes, variable importance Although nearest neighbors are simple and relationship between tumor class and gene LDA is unable to take into account gene With larger training sets, we expect an revealing interactions between features. correlation between genes helped here. underlying the class distinctions. Useful by–product of vs."Diagonal" LDA expression levels. interactions. statistics.

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Comparison study – results

- Simple classifiers such as naive Bayes (DLDA) and *k*–NN performed remarkably well compared to more complex ones.
- Aggregation improved the performance of unstable classifiers such as CART.
- The impact of gene screening varied depending on the dataset and classifier. Some screening of the genes down to G = 10 100 seems advisable.
- It is essential to take into account gene screening and other training decisions in error rate estimation procedures.

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Discussion

Discussion

There are two main goals

- **Prediction.** Predict biological outcomes for future samples (tumor class, survival) using a collection of predictor variables (gene expression measures).
- Information. Extract information about the underlying data generating mechanism, i.e., on the relationship between responses and predictor variables.

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R classification software

- class: knn and lvq functions.
- e1071: svm function.
- ipred: baggig, resampling based estimation of prediction error.
- LogitBoost: boosting for tree stumps.
- MASS: lda, polr, and qda functions.
- mlbench: Machine Learning Benchmark Problems.
- RanForests: http:/zoberkeley.edu/users/breiman/.
- rpart: classification and regression trees.

Data models: High–dimensionalata, unknown distribution, many models will provide similar predictive accuracy.

Accuracy vs. simplicity (interpretability): a predictor does not have to be simple to provide accurate prediction or reliable information about the relationship between responses and predictor variables (e.g. random forests).

Dimensionality: newer classification procedures thrive on the number of variables, the more the better.

Ref. Breiman (2001)

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