An introduction into the setup, definitions and use of PRTools is given. PRTools4 is extended and enhanced with respect to version 3 and thereby not fully compatible with it. This manual includes the description of a further upgrade: PRTools4.1. Still not all possibilities are fully exploited on the user level, or not at all. See release notes on page 58. Readers are assumed to be familiar with Matlab and should have a basic understanding of field of statistical pattern recognition.

Delft Pattern Recognition Research  tel: +31 15 2786143
Faculty EWI - ICT fax: +31 15 2781843
Delft University of Technology email: prtools@prtools.org
P.O. Box 5046, 2600 GA Delft http://prtools.org/
The Netherlands
**Availability, licences, copyright, reference**

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1. Motivation

In statistical pattern recognition one studies techniques for the generalization of examples to decision rules to be used for the detection and recognition of patterns in experimental data. This area of research has a strong computational character, demanding a flexible use of numerical programs for data analysis as well as for the evaluation of the procedures. As still new methods are being proposed in the literature a programming platform is needed that enables a fast and flexible implementation. Pattern recognition is studied in almost all areas of applied science. Thereby the use of a widely available numerical toolset like Matlab may be profitable for both, the use of existing techniques, as well as for the study of new algorithms. Moreover, because of its general nature in comparison with more specialized statistical environments, it offers an easy integration with the preprocessing of data of any nature. This may certainly be facilitated by the large set of toolboxes available in Matlab.

The about 200 pattern recognition routines and the additional 200 support routines offered by PRTools in its present state represent a basic set covering largely the area of statistical pattern recognition. Many methods and proposals, however, are not yet implemented. Some choices are accidental as the routines were programmed by the developers for their own research, sometimes in a way that was good for their private purposes. The important field of neural networks has partially been skipped as Matlab already includes a very good toolbox in that area. Just an interface to some basic routines is offered by PRTools to facilitate a comparison with traditional techniques.

PRTools has a few limitations. Due to the heavy memory demands of Matlab very large problems with learning sets of tens of thousands of objects cannot always be handled directly. In version 4.1 of the toolbox some tools to use large sets of files on disk are included. In the present version, PRTools4, the handling of missing data has been prepared, but hardly any routine has been are implemented. The use of symbolic data is not supported. Recently the possibility of soft (and thereby also fuzzy) labels has been added, as well as the usage of multiple labels. Just a few routines make use of them now. Also multi-dimensional target fields are allowed, but at this moment no procedure makes use of this possibility. Finally, support for misclassification costs has been implemented, but this is still on an experimental level.

In section 2 we present the basic philosophy about mappings and datasets. Section 3 presents the actual implementation, which is illustrated by examples in section 4. In section 5 further details are given, focussing on defining and using datasets and mappings. Section 7 lists the most important procedures of the toolbox. The examples included in the distribution of PRTools are listed in section 8, together with their expected results. Finally release notes of the versions 4.0 and 4.1 are given in sections 9 and 10. Here a summary of changes can be found that may be important for experienced users of PRTools.
2. Essential concepts

For the automatic recognition of the classes of objects, first some measurements have to be collected, e.g. using sensors, then they have to be represented, e.g. in a feature space and after some possible feature reduction steps they can be finally mapped by a classifier on the set of class labels. Between the initial representation in the feature space and this final mapping on the set of class labels the representation may be changed several times: simplified feature spaces (feature selection), normalization of features (e.g. by scaling), linear or nonlinear mappings (feature extraction), classification by a possible set of classifiers, combining classifiers and the final labelling. In each of these steps the data is transformed by some mapping.

Based on this observation the following two basic concepts of PRTools are defined:

- **datasets**: matrices in which the rows represent the objects and the columns the features, class memberships, or other fixed sets of properties (e.g. distances to a fixed set of other objects). In PRTool4.1 an extension of the dataset concept has been defined: **datafiles**. These refer to datasets to be created from directories of files.

- **mappings**: transformations operating on datasets.

As pattern recognition has two stages, *training* and *execution*, mappings have also two types: *untrained* and *trained*.

An *untrained mapping* refers just to the concept of a method, e.g. forward feature selection, PCA, or Fisher’s linear discriminant. It may have some parameters that are needed for training, e.g. the desired number of features or some regularization parameters. If an untrained mapping is applied to a dataset it will be trained (*training*).

A *trained mapping* is specific for the training set used to train the mapping. This dataset thereby determines the input dimensionality (e.g. the number of input features) as well as the output dimensionality (e.g. the number of output features or the number of classes). When a trained mapping is applied to a dataset it will transform the dataset according to its definition (*execution*).

In addition *fixed mappings* are used. They are almost identical to trained mappings, except that they don’t result from a training step, but are directly defined by the user: e.g. the transformation of distances by a sigmoid function to the [0,1] interval.

PRTools deals with sets of *labeled* or *unlabeled objects* and offers routines for the generalization of such sets into functions for *mapping* and *classification*. A *classifier* is thereby a special case of a *mapping* as it maps objects on class labels or on [0,1] intervals that may be interpreted as *class memberships*, *soft labels*, or *posterior probabilities*. An *object* is a k-dimensional vector of *feature values*, *distances*, *(dis)similarities* or *class memberships*. Within PRTools they are usually just called features. It is assumed that for all objects in a problem all values of the same set of features are given. The space defined by the actual set of features is called the *feature space*. Objects are represented as points or vectors in this space. New objects in a feature space are usually gradually converted to labels by a series of *mappings* followed by a final *classifier*. 
Sets of objects may be given externally or may be generated by one of the data generation routines of PRTools. Their labels may also be given externally or may be the result of a cluster analysis. By this technique similar objects within a larger set are grouped (clustered). The similarity measure is defined by the cluster technique in combination with the object representation in the feature space. Some clustering procedures do not just generate labels, but also a classifier that classifies new objects in the same way.

A fundamental problem is to find a good distance measure that agrees with the dissimilarity of the objects represented by the feature vectors. Throughout PRTools the Euclidean distance is used as default. However, scaling the features and transforming the feature spaces by different types of mappings effectively changes the distance measure.

The dimensionality of the feature space may be reduced by the selection of subsets of good features. Several strategies and criteria are possible for searching good subsets. Feature selection is important because it decreases the amount of features that have to be measured and processed. In addition to the improved computational speed in lower dimensional feature spaces there might also be an increase in the accuracy of the classification algorithms.

Another way to reduce the dimensionality is to map the data on a linear or nonlinear subspace. This is called linear or nonlinear feature extraction. It does not necessarily reduce the number of features to be measured, but the advantage of an increased accuracy may still be gained. Moreover, as lower dimensional representations yield less complex classifiers better generalizations can be obtained.

Using a training set a classifier can be trained such that it generalizes this set of examples of labeled objects into a classification rule. Such a classifier can be linear or nonlinear and can be based on two different kinds of strategies. The first strategy minimizes the expected classification error by using estimates of the probability density functions. In the second strategy this error is minimized directly by optimizing the classification function over its performance over the learning set or a separate evaluation set. In this approach it has to be avoided that the classifier becomes entirely adapted to the training set, including its noise. This decreases its generalization capability. This 'overtraining' can be circumvented by several types of regularization (often used in neural network training). Another technique is to simplify the classification function afterwards (e.g. the pruning of decision trees).

In PRTools4.1 the possibility of an automatic optimisation has been introduced for parameters controlling the complexity or the regularization of the training procedures of mappings and classifiers. This is based on a cross validation (see below) over the training set and roughly increases the time needed for training by a factor 100.

Constructed classification functions may be evaluated by independent test sets of labeled objects. These objects have to be excluded from the training set, otherwise the evaluation becomes optimistically biased. If they are added to the training set, however, better classification functions can be expected. A solution to this dilemma is the use of cross validation and rotation methods by which a small fraction of objects is excluded from training and used for testing. This fraction is rotated over the available set of objects and results are averaged. The extreme case is the leave-one-out method for which the excluded fraction is as large as one object.
The performance of classification functions can be improved by the following methods:

1. A *reject* option in which the objects close to the decision boundary are not classified. They are rejected and might be classified by hand or by another classifier.

2. The selection or averaging of classifiers.

3. A multi-stage classifier for *combining* classification results of several other classifiers.

For all these methods it is profitable or necessary that a classifier yields some distance measure, confidence or posterior probability in addition to the hard, unambiguous assignment of labels.
3. Implementation

PRTools makes use of the possibility offered by Matlab to define “Classes” and “Objects”. These programming concepts should not be confused with the classes and objects as defined in Pattern Recognition. The two main “Classes” defined in PRTools are: dataset and mapping. As a child of dataset also datafile has been defined, inheriting most properties of dataset. A large number of operators (like * or []) and Matlab commands have been overloaded and have thereby a special meaning when applied to a dataset and/or a mapping.

The central data structure of PRTools is the dataset. It primarily consists of a set of objects represented by a matrix of feature vectors. Attached to this matrix is a set of labels, one for each object and a set of feature names, also called feature labels. Labels can be integer numbers or character strings. Moreover, a set of prior probabilities, one for each class, is stored. In most help files of PRTools, a dataset is denoted by A. In almost any routine this is one of the inputs. Almost all routines can handle multi-class object sets. It is possible that for some objects no label is specified (a NaN is used, or an empty string). Such objects are, unless otherwise mentioned, skipped during training. It is possible to define more than one set of labels in a dataset. For instance, when the objects are pixels in an image, then they may be labelled according their image segment, but also according to the image, or to the sensor used, or the place the image has been measured.

Data structures of the “Classes” mapping store data transformations (‘mappings’), classifiers, feature extracting results, data scaling definitions, nonlinear projections, etcetera. They are usually denoted by W.

The easiest way to apply a mapping W to a dataset A is by A*W. The matrix multiplication symbol * is overloaded to this purpose. It is similar to the pipe (‘|’) command in Unix. This operation may also be written as map(A,W). Like everywhere else in Matlab, concatenations of operations are possible, e.g. A*W1*W2*W3 and are executed from left to right.

A typical example is given below:

A = gendath([50 50]); % Generate Highleyman’s classes, 50 objects / class
% Training set C (20 objects / class)
% Test set D (30 objects / class)
[C,D] = gendat(A,[20 20]);

W1 = ldc(C); % linear
W2 = qdc(C); % quadratic
W3 = parzenc(C); % Parzen
W4 = bpxnc(C,3); % Neural net with 3 hidden units

disp([testc(D*W1),testc(D*W2),testc(D*W3),testc(D*W4)]);

scatterd(A); % scatter plot
plotc({W1,W2,W3,W4});
This command file first generates by `gendath` two sets of labeled objects, both containing 50 two-dimensional object vectors, and stores them, their labels and prior probabilities in the dataset $A$. The distribution follows the so-called 'Highleyman classes'. The next call to `gendat` takes this dataset and splits it at random into a dataset $C$, further on used for training, and a dataset $D$, used for testing. This training set $C$ contains 20 objects from both classes. The remaining $2 \times 30$ objects are collected in $D$.

In the next lines four classification functions (discriminants) are computed, called $W_1$, $W_2$, $W_3$ and $W_4$. The first three are in fact density estimators based on various assumptions (class priors stored in $C$ are taken into account). Formally they are just mappings, as $E = D \ast W_1$ computes the class densities for the objects stored in $D$. $E$ has as many columns as there are classes in the training set for $W_1$ (in this case two). As the test routine `testc` (test classifier) assigns objects (represented by the rows in $E$) to the class corresponding with the highest density (times prior probability) the mappings $W_1$, ..., $W_4$ can be used as classifiers. The linear classifier $W_1$ (ldc) and quadratic classifier $W_2$ (qdc) are both based on the assumption of normally distributed classes. The first assumes equal class covariance matrices. The Parzen classifier estimates the class densities by the Parzen density estimation and has a built-in optimization for the smoothing parameter. The fourth classifier uses a feed forward neural network with three hidden units. It is trained by the back propagation rule using a varying stepsize.

Below the results are displayed and plotted. The test dataset $D$ is used in `testc` on each of the four discriminants. They are combined in a cell array, but individual calls are possible as well. The estimated probabilities of error are displayed in the Matlab command window and may look like:

$$0.1500 \quad 0.0333 \quad 0.1333 \quad 0.0833$$

Finally the classes are plotted in a scatter diagram together with the discriminants, see below. The plot routine `plotc` draws a vectorized straight line for the linear classifiers and computes the discriminant function values in all points of the plot grid (default $30 \times 30$) for the nonlinear discriminants. After that, the zero discriminant values are computed by interpolation and plotted.
4. Advanced example

The following, more advanced example is one of the standard examples that comes with PRTools. It defines a set of base classifiers and combines them in several ways. They are trained and evaluated on a 10-dimensional 2-class problem consisting of just two normal distributions with high correlations. This example shows various constructs of PRTools that facilitate the handling of sets of classifiers, often desirable for comparative studies:

- The definition of a sequence of untrained mapping like feature selection procedure and a classifier (e.g. \( w_2 = \text{featself}([],'NN',3)*\text{ldc} \)).
- The simultaneous training of a set of untrained classifiers stored in a cell array \( (W) \) by the same training set \( (B) \) in a single call \( (V = B*W) \), resulting in a cell array of trained classifiers \( (V) \).
- The construction of a set of combined classifiers stored in a cell array \( (V\text{ALL}) \) from the combined set of base classifiers \( (V\text{ALL}) \) and a set of possible combining rules stored in a cell array \( (W\text{C}) \) by a single statement \( (V\text{C} = V\text{ALL} * W\text{C}) \).
- The simultaneous evaluation of a cell array of trained classifiers \( (V \text{ or } V\text{C}) \) by the same test set \( C \) in a single call \( (\text{testc}(C,V) \text{ or } \text{testc}(C,C\text{V})) \).

PREX_COMBINING  PRTools example on classifier combining

Presents the use of various fixed combiners for some classifiers on the 'difficult data'.

```matlab
% Generate 10-dimensional data
A = gendatd([100,100],10);

% Select the training set of 40 = 2x20 objects
% and the test set of 160 = 2x80 objects
[B,C] = gendat(A,0.2);

% Define 5 untrained classifiers, (re)set their names
% w1 is a linear discriminant (LDC) in the space reduced by PCA
w1 = klm([],0.95)*ldc;
w1 = setname(w1,'klm - ldc');
% w2 is an LDC on the best (1-NN leave-one-out error) 3 features
w2 = featself([],'NN',3)*ldc;
w2 = setname(w2,'NN-FFS - ldc');
% w3 is an LDC on the best (LDC apparent error) 3 features
w3 = featself([],ldc,3)*ldc;
w3 = setname(w3,'LDC-FFS - ldc');
% w4 is an LDC
w4 = ldc;
w4 = setname(w4,'ldc');
% w5 is a 1-NN
w5 = knnc([],1);
w5 = setname(w5,'1-NN');

% Store classifiers in a cell
W = \{w_1,w_2,w_3,w_4,w_5\};

% Train them all
V = B*W;

% Test them all
disp([newline 'Errors for individual classifiers'])
testc(C,V);

% Construct combined classifier
VALL = [V(:)];

% Define combiners
WC = \{prod\_c,mean\_c,median\_c,max\_c,min\_c,vote\_c\};

% Combine (result is cell array of combined classifiers)
VC = VALL * WC;

% Test them all
disp([newline 'Errors for combining rules'])
testc(C,VC)

This script generates the below output. Note that testc, if called with a cell array of classifiers, lists the names of the classifiers and generates a table.

Errors for individual classifiers

Test results result for

clsf_1 : klm - ldc
clsf_2 : NN-FFS - ldc
clsf_3 : LDC-FFS - ldc
clsf_4 : ldc
clsf_5 : 1-NN

<table>
<thead>
<tr>
<th></th>
<th>clsf_1</th>
<th>clsf_2</th>
<th>clsf_3</th>
<th>clsf_4</th>
<th>clsf_5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Difficult Dataset</td>
<td>0.094</td>
<td>0.475</td>
<td>0.081</td>
<td>0.081</td>
<td>0.163</td>
</tr>
</tbody>
</table>

Errors for combining rules

Test results result for

clsf_1 : Product combiner
clsf_2 : Mean combiner
clsf_3 : Median combiner
clsf_4 : Maximum combiner
clsf_5 : Minimum combiner
clsf_6 : Voting combiner

<table>
<thead>
<tr>
<th></th>
<th>clsf_1</th>
<th>clsf_2</th>
<th>clsf_3</th>
<th>clsf_4</th>
<th>clsf_5</th>
<th>clsf_6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Difficult Dataset</td>
<td>0.094</td>
<td>0.169</td>
<td>0.094</td>
<td>0.163</td>
<td>0.081</td>
<td>0.081</td>
</tr>
</tbody>
</table>
5. Some Details

The command help files and the examples given below should give sufficient information to use the toolbox with a few exceptions. These are discussed in the following sections. They deal with the ways classifiers and mappings are represented. As these are the constituting elements of a pattern recognition analysis, it is important that the user understands these issues.

5.1 Datasets

A dataset consists of a set of \( m \) objects, each given by \( k \) features. In PRTools such a dataset is represented by a \( m \times k \) matrix: \( m \) rows, each containing an object vector of \( k \) features. Usually a dataset is labeled. An example of a definition is:

```matlab
> A = dataset([1 2 3; 2 3 4; 3 4 5; 4 5 6],[3 3 5 5])
> 4 by 3 dataset with 2 classes: [2 2]
```

The 4 by 3 data matrix (4 objects given by 3 features) is accompanied by a label list of 4 labels, connecting each of the objects to one of the two classes, 3 and 5. Class labels can be numbers or strings and should always be given as rows in the label list. It is possible that some, or all objects are unlabeled. If the label list is not given all objects are unlabeled. In addition it is possible to assign labels to the columns (features) of a dataset:

```matlab
> A = dataset(rand(100,3),genlab([50 50],[3 5]'))
> A = setfeatlab(A,['r1';'r2';'r3'])
> 100 by 3 dataset with 2 classes: [50 50]
```

The routine `genlab` generates 50 labels with value 3, followed by 50 labels with value 5. By `setfeatlab` the labels (‘r1’, ‘r2’, ‘r3’) for the three features are set. These are just feature names. Various other fields can be set as well. One of the ways to see these fields is by converting the dataset to a structure:

```matlab
> struct(A)
ans =
    data: [100x3 double]
    lablist: {2x4 cell}
    nlab: [100x1 double]
    labtype: 'crisp'
    targets: []
    featlab: [3x2 char]
    featdom: {{[] {[]} {[]}}}
    prior: []
    cost: []
    objsize: 100
    featsize: 3
    ident: [100x1 struct]
    version: {{1x1 struct} '21-Jul-2007 15:16:57'}
    name: []
    user: []
```

They can be inspected individually by various get commands defined for datasets, e.g.
> getfeatlab(A)
ans =
r1
r2
r3

Important is the possibility to set prior probabilities for each of the classes by the command
setprior(A,prob,lablist). The prior values in prob should sum to one. If prob is empty or if
it is not supplied the prior probabilities are computed from the dataset label frequencies. If prob
equals zero then equal class probabilities are assumed.

More than a single set of labels can be defined for a dataset. E.g.
> A = addlabels(A,char('apple','pear','apple','banana'),'fruitnames')
creates a second set of labels for the same objects. The active one can be selected by the
changelablist command. The nlab field points into the active label list and is used by PRTools
to find the real labels:
>> A = changelablist(A,1)
4 by 3 dataset with 2 classes: [2  2]
>> getnlab(A)
ans =
  1
  1
  2
  2
>> getlablist(A)
ans =
    3
    5
>> A = changelablist(A,'fruitnames')
4 by 3 dataset with 3 classes: [2 1 1]
>> getnlab(A)
ans =
  1
  3
  1
  2
>> getlablist(A)
ans =
apple
banana
pear
> getlabels(A)
ans =
apple
pear
apple
banana
This last command, getlabels, does not refer to a field in the dataset, but retrieves the labels using the indices stored in nlab to get the corresponding elements of lablist. So getlabels(A) is equivalent to
> nlab = getnlab(A);
> labels = lablist(nlab,:);

Various others items stored in a dataset can be retrieved by commands like getdata and getsize.

The size of the dataset can be found by
> [m,k] = size(A);
> [m,k,c] = getsize(A);

in which \( m \) is the number of objects, \( k \) the number of features and \( c \) the number of classes (equal to \( \max(nlab) \)). Datasets can be combined by \([A;B]\) if \( A \) and \( B \) have equal numbers of features and by \([A \ B]\) if they have equal numbers of objects. Creating subsets of datasets can be done by \( A(I,J) \) in which \( I \) is a set of indices defining the desired objects and \( J \) is a set of indices defining the desired features.

The original data matrix can be retrieved by getdata(A), double(A) or by +A.

Be aware that the order of classes returned by getprob and getlablist is the standard order used in PRTools and may differ from the one used in the definition of \( A \).

For more information, type help datasets.

5.2 Datasets help information

Datasets in PRTools are in the MATLAB language defined as objects of the class DATASET. Below, the words 'object' and 'class' are used in the pattern recognition sense.

A dataset is a set consisting of \( M \) objects, each described by \( K \) features. In PRTools, such a dataset is represented by a \( M \times K \) matrix: \( M \) rows, each containing an object vector of \( K \) elements. Usually, a dataset is labeled. An example of a definition is:

\[
\begin{align*}
\text{DATA} & = [\text{RAND}(3,2) \ ; \ \text{RAND}(3,2)+0.5]; \\
\text{LABS} & = ['A';'A';'A';'B';'B';'B'];
\end{align*}
\]

\( A = \text{DATASET(DATA,LABS)}; \)

which defines a \([6 \times 2]\) dataset with 2 classes.

The \([6 \times 2]\) data matrix (6 objects given by 2 features) is accompanied by labels, assigning each of the objects to one of the two classes A and B. Class labels can be numbers or strings and should always be given as rows in the label list. A label may also have the value NaN or may be an empty string, indicating an unlabeled object. If the label list is not given, all objects are marked as unlabeled.

Various other types of information can be stored in a dataset. The most simple way to get an overview is by typing:
STRUCT(A)

which for the above example displays the following:

- **DATA**: [6x2 double]
- **LABLIST**: (2x4 cell)
- **NLAB**: [6x1 double]
- **LABTYPE**: 'crisp'
- **TARGETS**: []
- **FEATLAB**: [2x1 double]
- **FEATDOM**: {1x2 cell}
- **PRIOR**: []
- **COST**: []
- **OBJSIZE**: 6
- **FEATSIZE**: 2
- **IDENT**: [6x1 struct]
- **VERSION**: {{[1x1 struct] '21-Jul-2007 15:16:57'}
- **NAME**: []
- **USER**: []

These fields have the following meaning:

- **DATA**: an array containing the objects (the rows) represented by features (the columns). In the software and help-files, the number of objects is usually denoted by M and the number of features is denoted by K. So, DATA has the size of [M,K]. This is also defined as the size of the entire dataset.

- **LABLIST**: The names of the classes, can be strings stored in a character array. If they are numeric they are stored in a column vector. Mixtures of these are not supported. The LABLIST field is a structure in which more than a single label list and the corresponding priors and costs are stored. PRTools keeps automatically track of this. See the MULTI_LABELING help file for more details.

- **NLAB**: an [M x 1] vector of integers between 1 and C, defining for each of the M objects its class.

- **LABTYPE**: 'CRISP', 'SOFT' or 'TARGETS' are the three possible label types. In case of 'CRISP' labels, a unique class, defined by NLAB, is assigned to each object, pointing to the class names given in LABLIST.

  For 'SOFT' labels, each object has a corresponding vector of C numbers between 0 and 1 indicating its membership (or confidence or posterior probability) of each of the C classes. These numbers are stored in TARGETS of the size M x C. They don't necessarily sum to one for individual row vectors. Labels of type 'TARGETS' are in fact no labels, but merely target vectors of length C. The values are again stored in TARGETS and are not restricted in value.

- **TARGETS**: [M,C] array storing the values of the soft labels or targets.
FEATLAB : A label list (like LABLIST) of K rows storing the names of the features.
FEATDOM : A cell array describing for each feature its domain.
PRIOR : Vector of length C storing the class prior probabilities. They should sum to one. If PRIOR is empty ([]) it is assumed that the class prior probabilities correspond to the class frequencies.
COST : Classification cost matrix. COST(I,J) are the costs of classifying an object from class I as class J. Column C+1 generates an alternative reject class and may be omitted, yielding a size of [C,C]. An empty cost matrix, COST = [] (default) is interpreted as COST = ONES(C) - EYE(C) (identical costs of misclassification).
OBJSIZE : The number of objects, M. In case the objects are related to a n-dimensional structure, OBJSIZE is a vector of length n, storing the size of this structure. For instance, if the objects are pixels in a [20 x 16] image, then OBJSIZE = [20,16] and M = 320.
FEATSIZE : The number of features, K. In case the features are related to an n-dimensional structure, FEATSIZE is a vector of length n, storing the size of this structure. For instance, if the features are pixels in a [20 x 16] image, then FEATSIZE = [20,16] and K = 320.
IDENT : A structure array of M elements storing user defined fields giving additional information on each of the objects. See SETIDENT.
VERSION : Some information related to the version of PRTools used for defining the dataset.
NAME : A character string naming the dataset, possibly used to annotate related graphics.
USER : A structure with user defined fields not used by PRTools. See DATASET/SETUSER

The fields can be set in the following ways:

1. In the DATASET construction command after DATA and LABELS using the form of (field name, value pairs), e.g.
   A = DATASET(DATA,LABELS,'PRIOR',[0.4 0.6],'FEATLIST',['AA';'BB']);
   Note that the elements in PRIOR refer to classes as they are ordered in LABLIST.
2. For a given dataset A, the fields may be changed similarly by the SET command:
   A = SET(A,'PRIOR',[0.4 0.6],'FEATLIST',['AA';'BB']);
3. By the commands
   SETDATA, SETFEATLAB, SETFEATDOM, SETFEATSIZE, SETIDENT, SETLABELS, SETLABLIST, SETLABTYPE, SETNAME, SETNLAB, SETOBJSIZE, SETPRIOR, SETTARGETS, SETUSER.
4. By using the dot extension as for structures, e.g.
   A.PRIOR = [0.4 0.6];
A.FEATLIST = ['AA';'BB'];

Note that there is no field LABELS in the DATASET definition. Labels are converted to NLAB and LABLIST. Commands like SETLABELS and A.LABELS, however, exist and take care of the conversion.

The data and information stored in a dataset can be retrieved as follows:

1. By DOUBLE(A) and by +A, the content of the A.DATA is returned.
   
   \[ N,LABLIST \] = CLASSSIZES(A);

   It returns the numbers of objects per class and the class names stored in LABLIST.

   By DISPLAY(A), it writes the size of the dataset, the number of classes and the label type on the terminal screen.

   By SIZE(A), it returns the size of A.DATA: numbers of objects and features.

   By SCATTERD(A), it makes a scatter plot of a dataset.

   By SHOW(A), it may be used to display images that are stored as features or as objects in a dataset.

2. By the GET command, e.g:
   
   \[ PRIOR,FEATLIST \] = GET(A,'PRIOR','FEATLIST');

3. By the commands:
   
   GETDATA, GETFEATLAB, GETFEATSIZE, GETIDENT, GETLABELS, GETLABLIST, GETLABTYPE, GETNAME, GETNLAB, GETOBJSIZE, GETPRIOR, GETCOST, GETSIZE, GETTARGETS, GETTARGETS, GETUSER, GETVERSION.

   Note that GETSIZE(A) does not refer to a single field, but it returns \[ M,K,C \].

   The following commands do not return the data itself, instead they return indices to objects that have specific identifiers, labels or class indices:

   FINDIDENT, FINDLABELS, FINDNLAB.

4. Using the dot extension as for structures, e.g.

   PRIOR = A.PRIOR;
   FEATLIST = A.FEATLIST;

Many standard MATLAB operations and a number of general MATLAB commands have been overloaded for variables of the DATASET type.

### 5.3 Datafiles

Datafiles are constructed to solve the memory problem connected with datasets. The latter are always in core and their size is thereby restricted to the size of the computer memory. As in processing datasets often (temporarily) copies are created, it is in practice advisable to keep datasets under 10 million elements (objectsize x featuresize). A number of operations handle datasets sequentially, or can be written like that, e.g. fixed mappings, testing and the training of some simple classifiers. So there is no problem to have such data stored on disk and have it read when needed. The datafile construct enables this is in fact an administration to keep track of the data and to have all additional information ready to reshape the desired pieces of data into a dataset when it has to be processed.
In understanding the handling of datafiles it is important to keep this characteristic in mind: they are just administration and the real processing is postponed until it is needed. PRTools makes use of this characteristic to integrate in the datafile concept raw preprocessing of, for instance images and signals. Arbitrary preprocessing of images can be defined for datafiles. It is the responsibility of the user that a preprocessing sequence ends in a format that can be straightforwardly transformed into a dataset. E.g., after preprocessing all images have to be of the same size or generate feature vectors of the same length.

Datafiles are formally children of the `class` dataset. Consequently they inherit the properties of datasets and ideally every operation that can be performed on a dataset can also be performed on a datafile. As can be understood from the above, this cannot always be true due to memory restrictions. There is a general routine, prmemory, by which the user can define what the maximum dataset sizes are that can be allowed. PRTools makes use of this to split datafiles into datasets of feasible sizes. An error is generated when this is not possible.

Routines that don’t accept datafiles should return an understandable error messages if called with a datafile. In most help text is included when routines accept datafiles.

The use of datafiles starts by the datafile construct that points to a directory in which each file is interpreted as an object. There is a savedatafile command that executes all processing defined for a datafile and stores the data on disk, ready to be used for defining a new datafile. This is the only place where PRTools writes data to disk. For more details, read the next section.

### 5.4 Datafiles help information

Datafiles in PRTools are in the MATLAB language defined as objects of the class DATAPLLE. They inherit most of their properties of the class DATASET. They are a generalisation of this class allowing for large datasets distributed over a set of files. Before conversion to a dataset preprocessing can be defined. There are four types of datafiles:

- **raw**: Every file is interpreted as a single object in the dataset. These files may, for instance, be images of different size.
- **pre-cooked**: In this case the user should supply a command that reads a file and converts it to a dataset.
- **half-baked**: All files should be mat-files, containing a single dataset.
- **mature**: This is a datafile by PRTools, using the SAVEDATAFILE command after execution of all preprocessing defined for the datafile.

A datafile is, like a dataset, a set consisting of M objects, each described by K features. K might be unknown, in which case it is set to zero, K=0. Datafiles store an administration about the files or directories in which the objects are stored. In addition they can store commands to preprocess the files before they are converted to a dataset and postprocessing commands, to be executed after conversion to a dataset.

Datafiles are mainly an administration. Operations on datafiles are possible as long as they can be stored (e.g. filtering of images for raw datafiles, or object selection by GENDAT). Commands that are able to
process objects sequentially, like NMC and TESTC can be executed on datafiles.

Whenever a raw datafile is sufficiently defined by pre- and postprocessing it can be converted into a dataset. If this is still a large dataset, not suitable for the available memory, it should be stored by the SAVEDATAFILE command and is ready for later use. If the dataset is sufficiently small it can be directly converted into a dataset by DATASET.

The main commands specific for datafiles are:

- **DATAFILE** - constructor. It defines a datafile on a directory.
- **ADDPREPROC** - adds preprocessing commands (low level command)
- **ADDPOSTPROC** - adds postprocessing commands (low level command)
- **FILTM** - user interface to add preprocessing to a datafile.
- **SAVEDATAFILE** - executes all defined pre- and postprocessing and stores the result as a dataset in a set of matfiles.
- **DATASET** - conversion to dataset

Datafiles have the following fields, in addition to all dataset fields.

- **ROOTPATH** - Absolute path of the datafile
- **FILES** - names of directories (for raw datafiles) or mat-files (for converted datafiles)
- **TYPE** - datafile type
- **PREPROC** - preprocessing commands in a struct array
- **POSTPROC** - postprocessing commands as mappings
- **DATASET** - stores all dataset fields. Note that the DATA field as well as the target field are empty and that the IDENT.FILE_INDEX field is used to store for every object a pointer to a file or directory in FILES.

Almost all operations defined for datasets are also defined for datafiles, with a few exceptions. Also fixed and trained mappings can handle datafiles, as they process objects sequentially. The use of untrained mappings in combination with datafiles is a problem, as they have to be adapted to the sequential use of the objects. Mappings that can handle datafiles are indicated in the Contents file.

The possibility to define preprocessing of objects (e.g. images) with different sizes makes datafiles useful for handling raw data and measurements of features.

### 5.5 Classifiers and mappings

There are many commands to train and use mappings between spaces of different (or equal) dimensionalities. For example:
if $A$ is a $m$ by $k$ dataset ($m$ objects in a $k$-dimensional space) and $W$ is a $k$ by $n$ mapping (map from $k$ to $n$ dimensions) then $A*W$ is a $m$ by $n$ dataset ($m$ objects in a $n$-dimensional space)

Mappings can be linear or affine (e.g. a rotation and a shift) as well as nonlinear (e.g. a neural network). Typically they can be used as classifiers. In that case a $k$ by $n$ mapping maps a $k$-feature data vector on the output space of a $n$-class classifier (exception: 2-class classifiers like discriminant functions may be implemented by a mapping to a 1-dimensional space like the distance to the discriminant, $n = 1$).

Mappings are of the data type 'mapping' (class($W$) is 'mapping'), have a size of $[k,n]$ if they map from $k$ to $n$ dimensions. Mappings can be instructed to assign labels to the output columns, e.g. the class names. These labels can be retrieved by

```
labels = getlabels(W); before the mapping, or
labels = getlabels(A*W); after the dataset $A$ is mapped by $W$.
```

Mappings can be learned from examples, (labeled) objects stored in a dataset $A$, for instance by training a classifier:

```
W1 = ldc(A); the normal densities based linear classifier
W2 = knnc(A,3); the 3-nearest neighbor rule
W3 = svc(A,'p',2); the support vector classifier based on a 2-nd order polynomial kernel
```

Untrained or empty mappings are supported. They may be very useful. In this case the dataset is replaced by an empty set or entirely skipped:

```
V1 = ldc; V2 = knnc([],a); V3 = svc([],'p',2);
```

Such mappings can be trained later by

```
W1 = A*V1; W2 = A*V2; W3 = A*V3;
```

(which is equivalent to the statements a few lines above) or by using cell arrays

```
V = {ldc, knnc([],a), svc([],'p',2)}; W = A*V;
```

The mapping of a test set $B$ by $B*W1$ is now equivalent to $B*(A*V1)$. Note that expressions are evaluated from left to right, so $B*A*V1$ will result in an error as the multiplication of the two datasets ($B*A$) is executed first.

Some trainable mappings do not depend on class labels and can be interpreted as finding a feature space that approximates as good as possible the original dataset given some conditions and measures. Examples are the Karhunen-Loève Mapping ($klm$), principle component analysis ($pca$) and kernel mapping ($kernelm$) by which nonlinear, kernel PCA mappings can be computed.

In addition to trainable mappings, there are fixed mappings, which operation is not computed from a training set but defined by just a few parameters. A number of them can be set by cmapm. Other ones are $sigm$ and $invsigm$. 


The result $D$ of mapping a test set on a trained classifier, $D = B*W_1$ is again a dataset, storing for each object in $B$ the output values of the classifier. For discriminants they are sigmoids of distances, mapped on the $[0,1]$ interval, for neural networks their unnormalized outputs and for density based classifiers the densities. For all of them holds: the larger, the more similar with the corresponding class. The values in a single row (object) don’t necessarily sum to one. This can be achieved by the fixed mapping $\text{classc}$:

$$D = B*W_1*\text{classc}$$

The values in $D$ can be interpreted as posterior probability estimates or classification confidences. Such a classification dataset has column labels (feature labels) for the classes and row labels for the objects. The class labels of the maximum values in each object row can be retrieved by

$$\text{labels} = D*\text{labeld}; \text{ or } \text{labels} = \text{labeld}(D);$$

A global classification error follows from

$$e = D*\text{testc}; \text{ or } e = \text{testc}(D);$$

Mappings can be combined in the following ways:

- **sequential:** $W = W_1 * W_2 * W_3$ (equal inner dimensions)
- **stacked:** $W = [W_1, W_2, W_3]$ (equal numbers of 'rows' (input dimensions))
- **parallel:** $W = [W_1; W_2; W_3]$ (unrestricted)

The output size of the parallel mapping is irregularly equal to $(k_1+k_2+k_3)$ by $(n_1+n_2+n_3)$ as the output combining of columns is undefined. In a stacked or parallel mapping columns having the same label can be combined by various combiners like $\text{maxc}$, $\text{meanc}$ and $\text{prodc}$. If the classifiers $W_1$, $W_2$ and $W_3$ are trained for the same $n$ classes, their output labels are the same and may be combined by $W = \text{prodc}([W_1; W_2; W_3])$ into a $(k_1+k_2+k_3)$ by $n$ classifier.

The above combinations can also be defined for untrained mappings and can be trained afterwards. This may be useful if they have to be trained for a series of datasets.

$W$ for itself, or $\text{display}(W)$ lists the size and type of a classifier as well as the routine used for computing a mapping $A*W$. The construction of a combined mapping may be inspected by $\text{parsc}(W)$.

Affine mappings (e.g. constructed by $\text{klm}$) may be transposed. This is useful for back projection of data into the original space. For instance:

$$W = \text{klm}(A,3); \text{ % computes 3-dimensional KL transform}$$

$$B = A*W; \quad \text{% maps A on W, resulting in B.}$$

$$C = B*W'; \quad \text{% back-projection of B in the original space.}$$

A mapping may be given an output selection by $W = W(:,J)$, in which $J$ is a set of indices pointing to the desired classes.

$$B = A*W(:,J); \text{ is equivalent to } B = A*W; B = B(:,J);$$

Input selection is not possible for a mapping.
For more information, type \texttt{help mappings}.

5.6 Mappings help information

Mappings in PRTools are in the MATLAB language defined as objects of the class \texttt{MAPPING}. In the text below, the words ‘object’ and ‘class’ are used in the pattern recognition sense.

In the Pattern Recognition Toolbox PRTools, there are many commands to define, train and use mappings between spaces of different (or equal) dimensionalities. Mappings operate mainly on datasets, i.e. variables of the type \texttt{DATASET} (see also \texttt{DATASETS}) and generate datasets and/or other mappings. For example:

\begin{verbatim}
if A is an M x K dataset (M objects in a K-dimensional space) and W is a K x N mapping (a map from K to N dimensions) then A*W is an M x N dataset (M objects in a N-dimensional space)
\end{verbatim}

This is enabled by overloading the \texttt{*}-operator for the \texttt{MAPPING} variables. \(A*W\) is executed by \texttt{MAP(A,W)} and may also be called as such.

Mappings can be linear (e.g. a rotation) as well as nonlinear (e.g. a neural network). Typically they are used to represent classifiers. In that case, a \(K \times C\) mapping maps a \(K\)-feature data vector on the output space of a \(C\)-class classifier (an exception: some 2-class classifiers, like the discriminant functions may be implemented by a mapping onto a 1-dimensional space determined by the distance to the discriminant).

Mappings are of the data-type \texttt{MAPPING} (\texttt{CLASS(W)} is a \texttt{MAPPING}), have a size of \(K \times C\) if they map from \(K\) to \(C\) dimensions. Four types of mapping are defined:

- untrained, \(V = A*W\)

Trains the untrained mapping \(W\), resulting in the trained mapping \(V\). \(W\) has to be defined by \(W = \texttt{MAPPING(MAPPING\_FILE,{PAR1, PAR2})}\), in which \texttt{MAPPING\_FILE} is the name of the routine that executes the training and \texttt{PAR1, PAR2} are two parameters that have to be included into the call to \texttt{THE MAPPING\_FILE}. Consequently, \(A*W\) is executed by \texttt{PRTools as MAPPING\_FILE(A,PAR1,PAR2)}.

Example: train the 3-NN classifier on the generated data
\begin{verbatim}
W = knnc([],3); % untrained classifier
V = gendatd([50 50])*W; % trained classifier
\end{verbatim}

- trained, \(D = B*V\)

Maps the dataset \(B\) on the trained mapping or classifier \(V\), e.g. as trained above. The resulting dataset \(D\) has as many objects (rows) as \(A\), but its feature size is now \(C\) if \(V\) is a \(K \times C\) mapping. Typically, \(C\)
is the number of classes in the training set A or a reduced number of
features determined by the training of V. V is defined by
\[ V = MAPPING(MAPPING_FILE, 'trained', DATA, LABELS, SIZE_IN, SIZE_OUT) \]
where MAPPING_FILE is the name of the routine that executes the mapping,
DATA is a field in which the parameters are stored (e.g. weights) for
the mapping execution, LABELS are the feature labels to be assigned to
the resulting dataset \[ D = B^*V \] (e.g. the class names) and SIZE_IN and
SIZE_OUT are the dimensionalities of the input and output spaces. They
are used for error checking only. D = B^*V is executed by PRTools as
MAPPING_FILE(B,W).

Example:
\[
A = gendatd([50 50], 10); \quad \% \text{generate random 10D datasets} \\
B = gendatd([50 50], 10); \\
W = klm([], 0.9); \quad \% \text{untrained mapping, Karhunen-Loeve projection} \\
V = A^*W; \quad \% \text{trained mapping V} \\
D = B^*V; \quad \% \text{the result of the projection of B onto V}
\]

- **fixed**, \[ D = A^*W \]

Maps the dataset A by the fixed mapping W, resulting into a transformed
dataset D. Examples are scaling and normalization, e.g.
\[ W = MAPPING('SIGM', 'fixed', S) \]
defines a fixed mapping by the sigmoid function SIGM a scaling parameter S. A^*W is executed by PRTools as
SIGM(A,S).

Example: normalize the distances of all objects in A such that their
city block distances to the origin are one.
\[
A = gendatb([50 50]); \\
W = normm; \\
D = A^*W;
\]

- **combiner**, \[ U = V^*W \]

Combines two mappings. The mapping W is able to combine itself with V
and produces a single mapping U. A combiner is defined by
\[ W = MAPPING(MAPPING_FILE, 'combiner', {PAR1, PAR2}) \]
in which MAPPING_FILE is the name of the routine that executes the combining and PAR1, and PAR2
are the parameters that have to be included into the call to the
MAPPING_FILE. Consequently, V^*W is executed by PRTools as
MAPPING_FILE(V, PAR1, PAR2). In a call as \[ D = A^*V^*W \], first \[ B = A^*V \]
is resolved and may result in a dataset B. Consequently, W should be able
to handle datasets, and MAPPING_FILE is now called by
MAPPING_FILE(B, PAR1, PAR2) Remark: the combiner construction is not
necessary, since PRTools stores \[ U = V^*W \] as a SEQUENTIAL mapping (see
below) if W is not a combiner. The construction of combiners, however,
may increase the transparency for the user and efficiency in
computations.

Example:
A = gendatd([50 50],10); % generate random 10D datasets
B = gendatd([50 50],10);
V = klm([],0.9); % untrained Karhunen-Loeve (KL) projection
W = ldc;  % untrained linear classifier LDC
U = V*W;  % untrained combiner
T = A*U;  % trained combiner
D = B*T;  % apply the combiner (first KL projection, then LDC) to B

Differences between the four types of mappings are now summarized for a dataset A and a mapping W:
A*W   - untrained : results in a mapping
       - trained   : results in a dataset, size checking
       - fixed     : results in a dataset, no size checking
       - combiner  : treated as fixed

Suppose V is a fixed mapping, then for the various possibilities of the mapping W, the following holds:
A*(V*W) - untrained : evaluated as V*(A*V*W), resulting in a mapping
       - trained   : evaluated as A*V*W, resulting in a dataset
       - fixed     : evaluated as A*V*W, resulting in a dataset
       - combiner  : evaluated as A*V*W, resulting in a dataset

Suppose V is an untrained mapping, then for the various possibilities of the mapping W holds:
A*(V*W) - untrained : evaluated as A*V*(A*(A*V)*W), results in mapping
       - trained   : evaluated as A*V*W, resulting in a mapping
       - fixed     : evaluated as A*V*W, resulting in a mapping
       - combiner  : evaluated as A*(V*W), resulting in a mapping

Suppose V is a trained mapping, then for the various possibilities of the mapping W holds:
A*(V*W) - untrained : evaluated as V*(A*V*W), resulting in a mapping
       - trained   : evaluated as A*V*W, resulting in a dataset
       - fixed     : evaluated as A*V*W, resulting in a dataset
       - combiner  : evaluated as A*(V*W), resulting in a dataset

The data fields stored in the MAPPING W = A*QDC can be found by STRUCT(W) which may display:
MAPPING_FILE: 'normal_map'
MAPPING_TYPE: 'trained'
DATA        : [1x1 struct]
LABELS      : [2x1 double]
SIZE_IN     : 2
SIZE_OUT    : 2
SCALE       : 1
COST        : []
OUT_CONV    : 0
NAME        : []
These fields have the following meaning:

MAPPING_FILE: Name of the m-file that executes the mapping.
MAPPING_TYPE: Type of mapping: 'untrained','trained','fixed' or 'combiner'.
DATA : Parameters or data for handling or executing the mapping.
LABELS : Label list used as FEATLAB for labeling the features of the output DATASET.
SIZE_IN : Expected input dimensionality of the data to be mapped. If not set, it is neglected, otherwise it is used for the error checking and display of the mapping size on the command line.
SIZE_OUT : Dimensionality of the output space. It should correspond to the size of LABLIST. SIZE_OUT may be size vector, e.g. describing the size of an image. See also the FEATSIZE field of DATASET.
SCALE : Output multiplication factor. If SCALE is a scalar all multiplied by it. SCALE may also be a vector with size as defined by SIZE_OUT to set separate scalings for each output.
COST : Classification costs in case the mapping defines a classifier.
OUT_CONV : Defines for trained and fixed mappings the output conversion:
0 - no conversion (to be used for mappings that output confidences or densities;
1 - sigmoid (for discriminants that output distances);
2 - normalization (for converting densities and confidences into posterior probability estimates;
3 - for performing sigmoid as well as normalization.
NAME : Name of the mapping, used for informing the user on the command line, as well as for annotating plots.
USER : User field, not used by PRTools.
VERSION : Some information related to the version of PRTools used for the mapping definition.

The fields can be set in the following ways:
1. At the end of the MAPPING construction command by a set of { fieldname, value pairs }, e.g.
   W = MAPPING('affine','trained',DATA,LABELS,5,2,'NAME','PCA Mapping')
2. For a given mapping W fields may be changed similarly by the SET command: W = SET(W,'NAME','PCA Mapping');
3. By the commands SETMAPPING_FILE, SETMAPPING_TYPE, SETDATA, SETLABELS,
   SETSIZE, SETSIZE_IN, SETSIZE_OUT, SETSCALE, SETOUT_CONV, SETNAME and
   SETUSER.
4. Using the dot extension as for structures, e.g. A.NAME = 'PCA MAPPING'

The information stored in a mapping can be retrieved as follows:
1. By DOUBLE(W) and by +W the content of the W.DATA is returned.
   DISPLAY(W) writes the size of the mapping, the number of classes and
the label type on the terminal screen.

SIZE(W) returns dimensionalities of input space and output space.

SCATTERD(A) makes a scatter-plot of a dataset.

SHOW(W) may be used to display images that are stored in mappings with
the MAPPING_FILE ‘affine’.

2. By the GET command, e.g: [name,user] = GET(W,’NAME’,’USER’);
3. By the commands GETMAPPING_FILE, GETMAPPING_TYPE, GETDATA, GETLABELS,
   SIZE, GETSIZE, GETSIZE_IN, GETSIZE_OUT, GETSCALE, GETCOST,
   GETOUT_CONV, GETNAME and GETUSER.
4. Using the dot extension as for structures, e.g. NAME = W.NAME;
5. The routines ISAFFINE, ISCLASSIFIER, ISCOMBINER, ISEMPTY, ISFIXED,
   ISTRAINED and ISUNTRAINED test on some mapping types and states.

Some standard MATLAB operations have been overloaded for variables of the
type MAPPING. They are defined as follows:

W'       Defined for affine mappings only. It returns a transposed
         mapping.

[W V]    Builds a combined classifier (see STACKED) operating in the
         same feature space. A * [W V] = [A*W A*V].

[W;V]    Builds a combined classifier (see PARALLEL) operating in
         different feature spaces: [A B] * [W;V] = [A*W B*V]. W and V
         should be mappings that correspond to the feature sizes of
         A and B.

A*W      Maps a DATASET A by the MAPPING W. This is executed by
         MAP(A,W).

V*W      Combines the mappings V and W sequentially. This is executed
         by SEQUENTIAL(V,W).

W+c      Defined for affine mappings only.

W(:,K)   Output selection. If W is a trained mapping, just the features
         listed in K are returned.

5.7 How to write your own mapping

Users can add new mappings or classifiers by a single routine that should support the following type
of calls:

W = mymapm([], par1, par2, ...); Defines the untrained, empty mapping.
W = mymapm(A, par1, par2, ...); Defines the map based on the training dataset A.
B = mymapm(A, W); Defines the mapping of dataset A on W, resulting in a dataset B.

To see some examples list the routines kernelm or subsc.

Below the subspace classifier subsc is listed. This classifier approximates each class by a linear
subspace and assigns new objects to the class of the closest subspace found in the training set. The
dimensionalities of the subspaces can be directly set by W = subsc(A,N), in which the integer N
determines the dimensionality of all class subspaces, or by \( W = \text{subsc}(A,\alpha) \), in which \( \alpha \) is the desired fraction of the retained variance, e.g. \( \alpha = 0.95 \). In both cases the class subspaces \( V \) are determined by a principle component analysis of the single class datasets.

The three possible types of calls, listed above are handled in the three main parts of the routine. If no input parameters are given (\( \text{nargin} < 1 \)) or no input dataset is found (\( A \) is empty) an untrained classifier is returned. This is useful for calls like \( W = \text{subsc}([],N) \), defining an untrained classifier that can be used in routines like \( \text{cleval}(A,W,...) \) that operate on arbitrary untrained classifiers, but also to facilitate training by constructions as \( W = A\ast\text{subsc} \) or \( W = A\ast\text{subsc}([],N) \).

The training section of the routine is accessed if \( A \) is not empty and \( N \) is either not supplied or set by the user as a double (i.e. the subspace dimensionality or the fraction of the retained variance). PRTools takes care that calls like \( W = A\ast\text{subsc}([],N) \) are executed as \( W = \text{subsc}(A,N) \). The first parameter in the mapping definitions \( W = \text{mapping}(mfilename,...) \) is substituted by Matlab as ‘\text{subsc}’ (\( mfilename \) is a function that returns the name of the calling file). This string is stored by PRTools in the \text{mapping_file} field of the mapping \( W \) and used to call \text{subsc} whenever it has to be applied to a dataset.

The trained mapping \( W \) can be applied to a test dataset \( B \) by \( D = B\ast W \) or by \( D = \text{map}(B,W) \). Such a call is converted by PRTools to \( D = \text{subsc}(B,W) \). Consequently, the second parameter of \text{subsc()} \( N \) is now substituted by the mapping \( W \). This is executed in the final part of the routine. Here, the data stored in the data field of \( W \) during training is retrieved (class mean, rotation matrix and mean square distances of the training objects) and used to find normalized distances of the test objects to the various subspaces. Finally they are converted to a density, assuming a normal distribution of distances. These values are returned in a dataset using the \text{setdata} routine. This dataset is thereby similar to the input dataset: it contains the same object labels, object identifiers, etcetera. Just the data itself is changed and the columns refer now to classes instead of features.

%SUBSC Subspace Classifier
% % \( W = \text{SUBSC}(A,N) \)
% \( W = \text{SUBSC}(A,\text{FRAC}) \)
% % INPUT
% \( A \) Dataset
% \( N \) or \( \text{FRAC} \) Desired model dimensionality or fraction of retained
% \( \text{variance per class} \)
% % OUTPUT
% \( W \) Subspace classifier
% % DESCRIPTION
% Each class in the trainingset \( A \) is described by linear subspace of
% dimensionality \( N \), or such that at least a fraction \( \text{FRAC} \) of its variance
% is retained. This is realised by calling \( \text{PCA}(A\text{I},N) \) or \( \text{PCA}(A\text{I},\text{FRAC}) \) for
% each subset \( A\text{I} \) of \( A \) (objects of class \( I \)). For each class a model is
% built that assumes that the distances of the objects to the class
% subspaces follow a one-dimensional distribution.
% New objects are assigned to the class of the nearest subspace.
% Classification by \( D = B^*W \), in which \( W \) is a trained subspace classifier
% and \( B \) is a testset, returns a dataset \( D \) with one-dimensional densities
% for each of the classes in its columns.
% If \( N \) (ALF) is NaN it is optimised by REGOPTC.

% REFERENCE
% See DATASETS, MAPPINGS, PCA, FISHERC, FISHERM, GAUSSM, REGOPTC

function W = subsc(A,N)

    name = 'Subspace classf.';

    % handle default
    if nargin < 2, N = 1; end

    % handle untrained calls like subsc([],3);
    if nargin < 1 | isempty(A)
        W = mapping(mfilename,{N});
        W = setname(W,name);
        return
    end

    if isa(N,'double') & isnan(N)    % optimize regularisation parameter
        defs = {1};
        parmin_max = [1,size(A,2)];
        W = regoptc(A,mfilename,{N},defs,[1],parmin_max,testc([],'soft'),0);
    elseif isa(N,'double')

        % handle training like A*subsc, A*subsc([],3), subsc(A)
        % PRTools takes care that they are all converted to subsc(A,N)

        islabtype(A,'crisp');    % allow crisp labels only
        isvaldfile(A,1,2);        % at least one object per class, two objects
        % allow for datafiles
        A = testdatasize(A,'features');    % test whether they fit
        A = setprior(A,getprior(A));        % avoid many warnings
        [m,k,c] = getsize(A);        % size of the training set
        for j = 1:c
            % run over all classes

    end
B = seldat(A,j); % get the objects of a single class only
u = mean(B); % compute its mean
B = B - repmat(u,size(B,1),1); % subtract mean
v = pca(B,N); % compute PCA for this class
v = v*v'; % trick: affine mappings in original space
B = B - B*v; % differences of objects and their mappings
s = mean(sum(B.*B,2)); % mean square error w.r.t. the subspace
data(j).u = u; % store mean
data(j).w = v; % store mapping
data(j).s = s; % store mean square distance
end
% define trained mapping,
% store class labels and size
W = mapping(mfilename,'trained',data,getlablist(A),k,c);
W = setname(W,name);

elseif isa(N,'mapping')

% handle evaluation of a trained subspace classifier W for a dataset A.
% The command D = A*W is by PRTools translated into D = subsc(A,W)
% Such a call is detected by the fact that N appears to be a mapping.
W = N; % avoid confusion: call the mapping W
m = size(A,1); % number of test objects
[k,c] = size(W); % mappingsize: from K features to C classes
d = zeros(m,c); % output: C class densities for M objects
for j=1:c % run over all classes
u = W.data(j).u; % class mean in training set
v = W.data(j).w; % mapping to subspace in original space
s = W.data(j).s; % mean square distance
B = A - repmat(u,m,1); % substract mean from test set
B = B - B*v; % differences objects and their mappings
d(:,j) = sum(B.*B,2)/s; % convert to distance and normalise
end
d = exp(-d/2)/sqrt(2*pi); % convert to normal density
A = dataset(A); % make sure A is a dataset
d = setdata(A,d,getlabels(W)); % take data from D and use
% class labels as given in W
% other information in A is preserved
W = d; % return result in output variable W
else
error('Illegal call') % this should not happen
end

return
6. References


7. A review of the toolbox

We will now shortly discuss the PRTools commands group by group. The two basic structures of the toolbox can be defined by the constructors dataset and mapping. In a dataset the feature values of a set of objects are stored, together with their class labels, feature names, prior probabilities, classifications costs and various types of user annotation. There are many commands to store data in and retrieve data from a dataset. These commands can also be used to retrieve or redefine the data. It is thereby not necessary to use the general Matlab converter struct() for decomposing the structures. By getlabels and getfeatlab the labels assigned to the objects and features can be found. The generation and handling of data is further facilitated by genlab for the generation of labels and renumlab for the parsing of labels and coding them into natural numbers between one and the number of classes. These numerical labels can be retrieved by getnlab. They point into a list of class labels called lablist, which can be retrieved by getlablist.

<table>
<thead>
<tr>
<th>Datasets and Mappings</th>
</tr>
</thead>
<tbody>
<tr>
<td>dataset</td>
</tr>
<tr>
<td>datasets</td>
</tr>
<tr>
<td>classsizes</td>
</tr>
<tr>
<td>get</td>
</tr>
<tr>
<td>getlabels</td>
</tr>
<tr>
<td>getnlab</td>
</tr>
<tr>
<td>getfeat</td>
</tr>
<tr>
<td>getfeatlab</td>
</tr>
<tr>
<td>getlablist</td>
</tr>
<tr>
<td>genclass</td>
</tr>
<tr>
<td>genlab</td>
</tr>
<tr>
<td>remclass</td>
</tr>
<tr>
<td>seldat</td>
</tr>
<tr>
<td>setdata</td>
</tr>
<tr>
<td>addlabels</td>
</tr>
<tr>
<td>setlabels</td>
</tr>
<tr>
<td>changelablist</td>
</tr>
<tr>
<td>multi_labeling</td>
</tr>
<tr>
<td>matchlab</td>
</tr>
<tr>
<td>renumlab</td>
</tr>
<tr>
<td>primport</td>
</tr>
<tr>
<td>mapping</td>
</tr>
<tr>
<td>mappings</td>
</tr>
<tr>
<td>getlab</td>
</tr>
</tbody>
</table>
There is a large set of routines for the generation of arbitrary normally distributed classes (gauss), and for various specific problems (gendatc, gendatd, gendath, gendatm and gendats).

There are two commands for enriching classes by noise injection (gendatk and gendatp). These are used for the general test set generator gendatt. A given dataset can be split into a training set and a test set gendat. The routine gendat splits the dataset at random into two sets. Subsets of datasets can be created by seldat. A total overview of all commands to generate datasets and to read datasets from disk (provided they are available) is given by prdatasets.

<table>
<thead>
<tr>
<th>Data Generation</th>
</tr>
</thead>
<tbody>
<tr>
<td>circles3d  Create a dataset containing 2 circles in 3 dimensions</td>
</tr>
<tr>
<td>lines5d   Create a dataset containing 3 lines in 5 dimensions</td>
</tr>
<tr>
<td>gauss     Generation of multivariate Gaussian distributed data</td>
</tr>
<tr>
<td>gencirc   Generation of a one-class circular dataset</td>
</tr>
<tr>
<td>gendatb   Generation of banana shaped classes</td>
</tr>
<tr>
<td>gendatc   Generation of circular classes</td>
</tr>
<tr>
<td>gendatd   Generation of two difficult classes</td>
</tr>
<tr>
<td>gendath   Generation of Highleyman classes</td>
</tr>
<tr>
<td>gendati   Generation of random windows from images</td>
</tr>
<tr>
<td>gendatk   Nearest neighbor data generation</td>
</tr>
<tr>
<td>gendatl   Generation of Lithuanian classes</td>
</tr>
<tr>
<td>gendatm   Generation of many Gaussian distributed classes</td>
</tr>
<tr>
<td>gendatp   Parzen density data generation</td>
</tr>
<tr>
<td>gendats   Generation of two Gaussian distributed classes</td>
</tr>
<tr>
<td>prdata    Read data from file and convert into a dataset</td>
</tr>
<tr>
<td>seldat    Select classes / features / objects from dataset</td>
</tr>
<tr>
<td>prdataset Read existing dataset from file</td>
</tr>
<tr>
<td>prdatasets Overview of all datasets and data generators</td>
</tr>
</tbody>
</table>

These are the main specific datafile commands needed for the user: the definition, saving datafiles and operating on datafiles. Many mappings for datasets can be applied on datafiles as well. They are however just stored internally in the datafile administration and only executed when the datafile is converted to a dataset, OR when it is stored by savedatafile.

<table>
<thead>
<tr>
<th>Datafiles</th>
</tr>
</thead>
<tbody>
<tr>
<td>datafile             Define datafile from set of files in directory</td>
</tr>
<tr>
<td>savedatafile         Save datafile, store intermediate result</td>
</tr>
<tr>
<td>filtm                Mapping for arbitrary processing of datafile</td>
</tr>
</tbody>
</table>

- 34 -
All routines operate in multi-class problems. `labeld` and `testc` are the general classification and testing routines. They can handle any classifier from any routine, including the ones to follow.

Classifiers and mappings can be trained by a dataset using commands like $W = \text{fisherc}(A)$, or $W = \text{knncc}(A,3)$. Such commands may also be written as $W = A*\text{fisherc}$, and $W = A*\text{polyc}([],[],3)$. The possibility to assign an untrained classifier to a variable like $V = \text{polyc}([],[],3)$ allows for routines that have untrained classifiers as input, e.g. the general classifier evaluation routine `cleval` (see below).

Some more examples, also showing the use of cell arrays of classifiers and datasets:

```
A = \text{gendatb}([100,100]); \% Generate 2 classes of 100 objects each
\% Generate 50% for training
[Train,Test] = \text{gendat}(A,0.5); \% and 50% for testing
\% Define set of untrained classifiers
W = {\text{fisherc}, \text{loglc, nmc, polyc}([],[],3)};
V = Train*W; \% Train them all and construct classifiers
D = Test*V; \% Test them by C
E = D*testc; \% Store classification errors
```
Classifiers for normal distributed classes can be trained by ldc, qdc and udc, while nbayesc assumes known densities. The all follow the Bayes rule using the priors stored in the datasets. The special purpose test routine testn can be used if the parameters of the normal distribution (means and covariances) are known or estimated by meancov.

### Normal Density Based Classification

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>distmaha</td>
<td>Mahalanobis distance</td>
</tr>
<tr>
<td>meancov</td>
<td>Estimation of means and covariance matrices</td>
</tr>
<tr>
<td>nbayesc</td>
<td>Bayes classifier for given normal densities</td>
</tr>
<tr>
<td>ldc</td>
<td>Normal densities based linear classifier</td>
</tr>
<tr>
<td>qdc</td>
<td>Normal densities based quadratic classifier</td>
</tr>
<tr>
<td>udc</td>
<td>Normal densities based classifier (independent features)</td>
</tr>
<tr>
<td>mogc</td>
<td>Mixture of gaussians classification</td>
</tr>
<tr>
<td>testn</td>
<td>Error estimate of discriminant on normal distributions</td>
</tr>
</tbody>
</table>

### Nonlinear Classification

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>knnc</td>
<td>k-nearest neighbor classifier</td>
</tr>
<tr>
<td>testk</td>
<td>Error estimation for k-nearest neighbor rule</td>
</tr>
<tr>
<td>edicon</td>
<td>Edit and condense training sets</td>
</tr>
<tr>
<td>parzen</td>
<td>Parzen classifier</td>
</tr>
<tr>
<td>parzendc</td>
<td>Parzen density based classifier</td>
</tr>
<tr>
<td>testp</td>
<td>Error estimation for Parzen classifier</td>
</tr>
<tr>
<td>treec</td>
<td>Construct binary decision tree classifier</td>
</tr>
<tr>
<td>naivebc</td>
<td>Naive Bayes classifier</td>
</tr>
<tr>
<td>bpxnc</td>
<td>Train neural network classifier by back-propagation</td>
</tr>
<tr>
<td>lmnc</td>
<td>Train neural network by Levenberg-Marquardt rule</td>
</tr>
<tr>
<td>perlc</td>
<td>Linear perceptron</td>
</tr>
<tr>
<td>rbnc</td>
<td>Train radial basis neural network classifier</td>
</tr>
<tr>
<td>neurc</td>
<td>Automatic neural network classifier</td>
</tr>
<tr>
<td>rncnc</td>
<td>Random neural network classifier</td>
</tr>
<tr>
<td>svc</td>
<td>Support vector classifier</td>
</tr>
<tr>
<td>nusvc</td>
<td>Support vector classifier</td>
</tr>
<tr>
<td>rbsvc</td>
<td>Radial basis SV classifier</td>
</tr>
<tr>
<td>kernelc</td>
<td>General kernel/dissimilarity based classification</td>
</tr>
</tbody>
</table>

knnc and parzen are similar in the sense that the classifiers they build still include all training objects and that their parameter (the number of neighbors or the smoothing parameter) can be user supplied or can be optimized over the training set using a leave-one-out error estimation. For the Parzen classifier the smoothing parameter can also be estimated by parzenml using an optimization of the density estimation. The special purpose testing routines testk and testp are useful for obtaining leave-one-out error estimations. parzendc is based on a optimization of each of the class densities separately by parzenml.
Decision trees can be constructed by `treec`, using various criterion functions, stopping rules or pruning techniques. The resulting classifier can be used in `labeld, testc` and `plotc`.

PRTools offers three neural network classifiers (`bpxnc`, `lmnn` and `rbnn`) based on Matlab’s Neural Network Toolbox, which should be available to run these routines. The resulting classifiers are ready to use by `labeld, testc` and `plotc`. The automatic neural network classifier `neurc` builds a network without any parameter setting by the user. Random neural network classifiers can be generated by `rnnc`. Its first layer is totally random, the second layer is optimized by a linear classifier.

The Support Vector Classifier (`svc`) can be called for various kernels as defined by `kernelm`. `nusvm` is a slightly different version in which the regularisation parameter can be defined in terms of the expected error. `rbsvm` optimizes parameter settings internally. The support vector classifiers are optimized by a quadratic programming procedure.

<table>
<thead>
<tr>
<th>Feature Selection</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>feateval</strong></td>
</tr>
<tr>
<td><strong>featrank</strong></td>
</tr>
<tr>
<td><strong>featselb</strong></td>
</tr>
<tr>
<td><strong>featself</strong></td>
</tr>
<tr>
<td><strong>featsellr</strong></td>
</tr>
<tr>
<td><strong>featseli</strong></td>
</tr>
<tr>
<td><strong>featselo</strong></td>
</tr>
<tr>
<td><strong>featselp</strong></td>
</tr>
<tr>
<td><strong>featselm</strong></td>
</tr>
</tbody>
</table>

The feature selection routines `featselb`, `featself`, `featseli`, `featselo` and `featselp` generate subsets of features, calling `feateval` for evaluating the feature set. `featselm` offers a general entry for feature selection, calling one of the other methods. All routines produce a mapping \( W \) (e.g. \( W = \text{featself}(A, [], k) \)). So the reduction of a dataset \( A \) to \( B \) is done by \( B = A \times W \).
A classifier maps, after training, objects from the feature space into its output space. For two-class discriminants these are sigmoids of distances, for neural networks their unnormalized outputs (i.e. they don’t necessarily sum to one) and for density based classifiers the densities. Discriminants are normalized such that their sigmoid outputs are optimal posterior probability estimates. The dimensionality of the classifier output space equals the number of classes (an exception is possible for two-class classifiers, that may have a one-dimensional output space). This output space may be mapped on posterior probability for other classifiers than discriminants by \( \text{classc} \), which takes care of normalization. Classification (determining the class with maximum output) is done by \( \text{labeld} \), which generates the labels of that class.

A general Bayes plug-in classification if offered by \( \text{bayesc} \). This routine expects as inputs proper density estimating routine. Suppose we have one-class datasets \( A \), \( B \) and \( C \) for which the densities are estimators are determined by \( WA = \text{gaussm}(A,3) \), \( WB = \text{knnm}(B,5) \) and \( WC = \text{parzenm}(C) \), then a Bayes classifier using class priors \( P = [0.3 \ 0.3 \ 0.4] \), can be built by \( \text{W} = \text{bayesc}(WA,WB,WC,[0.3 \ 0.3 \ 0.4],\text{char}('\text{apple}','\text{banana}','\text{coco}')) \).

In order to make various density based classifiers like \( \text{ldc} \), \( \text{udc} \), \( \text{qdc} \), \( \text{mogc} \), \( \text{parzenc} \), \( \text{parzendc} \) and \( \text{knnm} \) comparable, they output the proper densities (e.g. \( D = B*\text{qdc}(A) \)). For high-dimensional spaces this causes that in the tails of the distributions an exact zero density is returned, due to the finite numerical accuracy. This may even be the case for all classes, by which the posterior probabilities, computed after applying \( \text{classc} \) (\( D = B*\text{qdc}(A) * \text{classc} \)), become undefined. The routine \( \text{logdens} \) may be used to solve this problem. It forces the density based classifiers based on normal distributions and Parzen estimators (\( \text{ldc}, \text{udc}, \text{qdc}, \text{mogc}, \text{parzenc}, \text{parzendc} \)) to a direct computation of log-densities, followed by an appropriate rescaling and an immediate

---

### Classifiers and Tests (general)

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{bayesc} )</td>
<td>Bayes classifier by combining density estimates</td>
</tr>
<tr>
<td>( \text{classim} )</td>
<td>Classify image using a given classifier</td>
</tr>
<tr>
<td>( \text{classc} )</td>
<td>Convert mapping to classifier</td>
</tr>
<tr>
<td>( \text{labeld} )</td>
<td>Find labels of objects by classification</td>
</tr>
<tr>
<td>( \text{cleveld} )</td>
<td>Classifier evaluation (learning curve)</td>
</tr>
<tr>
<td>( \text{clevelbd} )</td>
<td>Classifier evaluation, bootstrap version</td>
</tr>
<tr>
<td>( \text{clevalf} )</td>
<td>Classifier evaluation (feature size curve)</td>
</tr>
<tr>
<td>( \text{confmat} )</td>
<td>Computation of confusion matrix</td>
</tr>
<tr>
<td>( \text{costm} )</td>
<td>Cost mapping, classification using costs</td>
</tr>
<tr>
<td>( \text{crossval} )</td>
<td>Error estimation by cross-validation</td>
</tr>
<tr>
<td>( \text{cnormc} )</td>
<td>Normalization of discriminants</td>
</tr>
<tr>
<td>( \text{dispererror} )</td>
<td>Error matrix, information on classifiers and datasets</td>
</tr>
<tr>
<td>( \text{logdens} )</td>
<td>Convert density estimates to log-densities</td>
</tr>
<tr>
<td>( \text{labelim} )</td>
<td>Construct image of labeled pixels</td>
</tr>
<tr>
<td>( \text{mclassc} )</td>
<td>Multi-class classifier from 2-class discriminants</td>
</tr>
<tr>
<td>( \text{reject} )</td>
<td>Compute error-reject curve</td>
</tr>
<tr>
<td>( \text{roc} )</td>
<td>Compute receiver-operator curve</td>
</tr>
<tr>
<td>( \text{testc} )</td>
<td>Error estimation routine for trained classifiers</td>
</tr>
<tr>
<td>( \text{testauc} )</td>
<td>Estimate error as area under the ROC</td>
</tr>
</tbody>
</table>
normalization. Consequently $W = qdc(A)$; $D = B\log\text{dens}(W)$ computes better posterior probabilities in the tails of the distribution. This applies for lcd, udc, qdc, mogc, parzenc and parzendc.

Error estimates for test data are made by testc and confmat. More advanced techniques like rotating datasets over test sets and training sets, are offered by crossval, cleval and clevalb.

### Mappings

<table>
<thead>
<tr>
<th>Mapping</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>affine</td>
<td>Construct affine (linear) mapping from parameters</td>
</tr>
<tr>
<td>bhatm</td>
<td>Two-class Bhattacharyya mapping</td>
</tr>
<tr>
<td>cmapm</td>
<td>Compute some special maps</td>
</tr>
<tr>
<td>featselm</td>
<td>Feature selection map, general routine</td>
</tr>
<tr>
<td>fisherm</td>
<td>Fisher mapping</td>
</tr>
<tr>
<td>invsigm</td>
<td>Inverse sigmoid map</td>
</tr>
<tr>
<td>gaussm</td>
<td>Mixture of Gaussians density estimation</td>
</tr>
<tr>
<td>kernelm</td>
<td>PCA based kernel mapping</td>
</tr>
<tr>
<td>klm</td>
<td>Decorrelation and Karhunen Loève mapping (PCA)</td>
</tr>
<tr>
<td>klnms</td>
<td>Scaled version of klm, useful for pre-whitening</td>
</tr>
<tr>
<td>knnm</td>
<td>k-Nearest neighbor density estimation</td>
</tr>
<tr>
<td>map</td>
<td>General routine for computing and executing mappings</td>
</tr>
<tr>
<td>mclassm</td>
<td>Computation of mapping from multi-class dataset</td>
</tr>
<tr>
<td>nlfisher</td>
<td>Nonlinear Fisher mapping</td>
</tr>
<tr>
<td>normm</td>
<td>Object normalization map</td>
</tr>
<tr>
<td>parzentm</td>
<td>Parzen density estimation</td>
</tr>
<tr>
<td>parzenml</td>
<td>ML estimation of Parzen smoothing parameter.</td>
</tr>
<tr>
<td>pca</td>
<td>Principle Component Analysis</td>
</tr>
<tr>
<td>proxm</td>
<td>Proximity mapping and kernel construction</td>
</tr>
<tr>
<td>reducm</td>
<td>Reduce to minimal space mapping</td>
</tr>
<tr>
<td>rejectm</td>
<td>Creates rejecting mapping</td>
</tr>
<tr>
<td>remoutl</td>
<td>Remove outliers</td>
</tr>
<tr>
<td>scalem</td>
<td>Compute scaling data</td>
</tr>
<tr>
<td>sigm</td>
<td>Sigmoid mapping</td>
</tr>
<tr>
<td>spatm</td>
<td>Augment image dataset with spatial label information</td>
</tr>
<tr>
<td>kernelm</td>
<td>Kernel mapping, dissimilarity representation</td>
</tr>
<tr>
<td>userkernel</td>
<td>User supplied kernel definition</td>
</tr>
<tr>
<td>gtm</td>
<td>Fit a Generative Topographic Mapping (GTM) by EM</td>
</tr>
<tr>
<td>plotgtm</td>
<td>Plot a Generative Topographic Mapping in 2D</td>
</tr>
<tr>
<td>som</td>
<td>Simple routine computing a Self-Organizing Map (SOM)</td>
</tr>
<tr>
<td>plotsom</td>
<td>Plot a Self-Organizing Map in 2D</td>
</tr>
<tr>
<td>mds</td>
<td>Non-linear mapping by multi-dimensional scaling</td>
</tr>
<tr>
<td>mds_cs</td>
<td>Linear mapping by classical scaling</td>
</tr>
<tr>
<td>mds_init</td>
<td>Initialization of multi-dimensional scaling</td>
</tr>
<tr>
<td>mds_stress</td>
<td>Dissimilarity of distance matrices</td>
</tr>
</tbody>
</table>

Classifiers are a special type of mapping, as their output spaces are related to class membership. In general a mapping converts data from one space to another. This may be done by a fixed procedure, not depending on a dataset, but controlled by at most some parameters. Most of these mappings that
don’t need training are collected by cmapm (e.g. shifting, rotation, deletion of particular features), another example is the sigmoidal mapping sigm. Some of the mappings that need training don’t depend on the object labels, e.g. the principal component analysis (PCA) by pca, klm and klms, object normalization by normm and scaling by scalem, and nonlinear PCA or kernel PCA by kernelm. The other routines depend on object labels as they define the mapping such that the class separability is maximized in one way or another. The Fisher criterion is optimized by fisherm, the scatter by klm (if called by labelled data), density separability for normal distributions by nlfisherm and general class separability by lmnm.

<table>
<thead>
<tr>
<th>Combining classification rules</th>
</tr>
</thead>
<tbody>
<tr>
<td>averagec</td>
</tr>
<tr>
<td>baggingc</td>
</tr>
<tr>
<td>votec</td>
</tr>
<tr>
<td>maxc</td>
</tr>
<tr>
<td>minc</td>
</tr>
<tr>
<td>meanc</td>
</tr>
<tr>
<td>medianc</td>
</tr>
<tr>
<td>prodc</td>
</tr>
<tr>
<td>traincc</td>
</tr>
<tr>
<td>parsc</td>
</tr>
<tr>
<td>parallel</td>
</tr>
<tr>
<td>stacked</td>
</tr>
<tr>
<td>sequential</td>
</tr>
</tbody>
</table>

Classifiers can be combined by horizontal and vertical concatenation, see section 5.5, e.g. $W = [W_1, W_2, W_3]$. Such a set of classifiers can be combined by several rules, like majority voting (majorc), combining the posterior probabilities in several ways (maxc, minc, meanc, medianc and prodc), or by training an output classifier (traincc). The way classifiers are combined can be inspected by parsc.

<table>
<thead>
<tr>
<th>Regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>linearr</td>
</tr>
<tr>
<td>ridger</td>
</tr>
<tr>
<td>lassor</td>
</tr>
<tr>
<td>svmr</td>
</tr>
<tr>
<td>ksmoothr</td>
</tr>
<tr>
<td>knnr</td>
</tr>
<tr>
<td>pinvr</td>
</tr>
<tr>
<td>plsr</td>
</tr>
<tr>
<td>plsm</td>
</tr>
<tr>
<td>testr</td>
</tr>
<tr>
<td>rsquared</td>
</tr>
</tbody>
</table>
Images can be stored, either as features (im2feat), or as objects (im2obj) in a dataset. The first possibility is useful for segmenting images using a vector of values for each pixels (e.g. in case of multi-color images, or as a result of a filter bank). The second possibility enables the classification of entire images using their pixels as features. Such datasets can be displayed by `show`. The relation with image processing is established by `dataim`, enabling arbitrary image operations. Simple filtering can be sped up by `datfilt`, `datgauss` and `datunif`.

<table>
<thead>
<tr>
<th>Command</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>classim</code></td>
<td>Classify image using a given classifier</td>
</tr>
<tr>
<td><code>dataim</code></td>
<td>Image operation on dataset images.</td>
</tr>
<tr>
<td><code>data2im</code></td>
<td>Convert dataset to image</td>
</tr>
<tr>
<td><code>getobjsize</code></td>
<td>Retrieve image size of feature images in datasets</td>
</tr>
<tr>
<td><code>getfeatsize</code></td>
<td>Retrieve image size of object images in datasets</td>
</tr>
<tr>
<td><code>datfilt</code></td>
<td>Filter dataset image</td>
</tr>
<tr>
<td><code>datgauss</code></td>
<td>Filter dataset image by Gaussian filter</td>
</tr>
<tr>
<td><code>datunif</code></td>
<td>Filter dataset image by uniform filter</td>
</tr>
<tr>
<td><code>im2obj</code></td>
<td>Convert image to object in dataset</td>
</tr>
<tr>
<td><code>im2feat</code></td>
<td>Convert image to feature in dataset</td>
</tr>
<tr>
<td><code>spatm</code></td>
<td>Augment image dataset with spatial label information</td>
</tr>
<tr>
<td><code>show</code></td>
<td>Display images stored in dataset</td>
</tr>
</tbody>
</table>
For many of these operations the DipImage toolbox is needed, like for the following ones.

### Operations on images in datasets and datafiles

<table>
<thead>
<tr>
<th>Command</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>classim</td>
<td>Classify image using a given classifier</td>
</tr>
<tr>
<td>dataim</td>
<td>Image operation on dataset images (features or objects)</td>
</tr>
<tr>
<td>doublem</td>
<td>Convert datafile images into double</td>
</tr>
<tr>
<td>filtrim</td>
<td>Image operation on objects in datafiles/datasets</td>
</tr>
<tr>
<td>datfilt</td>
<td>Filter dataset image (outdated)</td>
</tr>
<tr>
<td>datgauss</td>
<td>Filter dataset image by Gaussian filter</td>
</tr>
<tr>
<td>datunif</td>
<td>Filter dataset image by uniform filter</td>
</tr>
<tr>
<td>hsv2rgb</td>
<td>Convert HSV to RGB images</td>
</tr>
<tr>
<td>rgb2hsv</td>
<td>Convert RGB to HSV images</td>
</tr>
<tr>
<td>spatm</td>
<td>Augment image dataset with spatial label information</td>
</tr>
<tr>
<td>im_bdilation</td>
<td>Binary dilation</td>
</tr>
<tr>
<td>im_berosion</td>
<td>Binary erosion</td>
</tr>
<tr>
<td>im_box</td>
<td>Bounding box</td>
</tr>
<tr>
<td>im_bpropagation</td>
<td>Binary propagation</td>
</tr>
<tr>
<td>im_center</td>
<td>Center image</td>
</tr>
<tr>
<td>im_fft</td>
<td>FFT transform (and more)</td>
</tr>
<tr>
<td>im_gaussf</td>
<td>Gaussian filtering</td>
</tr>
<tr>
<td>im_gray</td>
<td>Multi-band to gray-value conversion</td>
</tr>
<tr>
<td>im_histEqualize</td>
<td>Histogram equalization</td>
</tr>
<tr>
<td>im_invert</td>
<td>Invert image</td>
</tr>
<tr>
<td>im_label</td>
<td>Labeling binary images</td>
</tr>
<tr>
<td>im_maxf</td>
<td>Maximum filter</td>
</tr>
<tr>
<td>im_minf</td>
<td>Minimum filter</td>
</tr>
<tr>
<td>im_resize</td>
<td>Resize images</td>
</tr>
<tr>
<td>im_rotate</td>
<td>Rotate images</td>
</tr>
<tr>
<td>im_scale</td>
<td>Scale images</td>
</tr>
<tr>
<td>im_select_blob</td>
<td>Select largest blob</td>
</tr>
<tr>
<td>im_stretch</td>
<td>Contrast stretching of images</td>
</tr>
<tr>
<td>im_threshold</td>
<td>Threshold images</td>
</tr>
</tbody>
</table>

### Feature extraction from images in datasets and datafiles

<table>
<thead>
<tr>
<th>Command</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>histm</td>
<td>Convert images to histograms</td>
</tr>
<tr>
<td>im_moments</td>
<td>Computes moments as features from object images</td>
</tr>
<tr>
<td>im_mean</td>
<td>Computes center of gravity</td>
</tr>
<tr>
<td>im_measure</td>
<td>Computes some measurements</td>
</tr>
<tr>
<td>im_profile</td>
<td>Computes image profiles</td>
</tr>
<tr>
<td>im_stat</td>
<td>Compute some simple statistics</td>
</tr>
</tbody>
</table>
### Clustering and Distances

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>distm</td>
<td>Distance matrix between two data sets.</td>
</tr>
<tr>
<td>emclust</td>
<td>Expectation - maximization clustering</td>
</tr>
<tr>
<td>proxm</td>
<td>Proximity mapping and kernel construction</td>
</tr>
<tr>
<td>hclust</td>
<td>Hierarchical clustering</td>
</tr>
<tr>
<td>kcentres</td>
<td>K-centers clustering</td>
</tr>
<tr>
<td>kmeans</td>
<td>K-means clustering</td>
</tr>
<tr>
<td>modeseek</td>
<td>Clustering by mode seeking</td>
</tr>
<tr>
<td>mds</td>
<td>Non-linear mapping by multi-dimensional scaling (Sammon)</td>
</tr>
<tr>
<td>mds_cs</td>
<td>Linear mapping by classical scaling</td>
</tr>
<tr>
<td>mds_init</td>
<td>Initialisation of multi-dimensional scaling</td>
</tr>
<tr>
<td>mds_stress</td>
<td>Dissimilarity of distance matrices</td>
</tr>
</tbody>
</table>

### Plotting

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>gridsize</td>
<td>Set gridsize of scatterd, plotd and plotm plots</td>
</tr>
<tr>
<td>plotc</td>
<td>Plot discriminant function in scatterplot</td>
</tr>
<tr>
<td>plote</td>
<td>Plot error curves</td>
</tr>
<tr>
<td>plotf</td>
<td>Plot feature distribution</td>
</tr>
<tr>
<td>plotm</td>
<td>Plot mapping in scatterplot</td>
</tr>
<tr>
<td>ploto</td>
<td>Plot object functions</td>
</tr>
<tr>
<td>plotr</td>
<td>Plot error curves</td>
</tr>
<tr>
<td>plotdg</td>
<td>Plot dendrogram (see hclust)</td>
</tr>
<tr>
<td>scatterd</td>
<td>Scatterplot</td>
</tr>
<tr>
<td>scatterdui</td>
<td>Scatterplot scatterplot with feature selection</td>
</tr>
<tr>
<td>scatterr</td>
<td>Scatter regression dataset</td>
</tr>
</tbody>
</table>

### Examples

<table>
<thead>
<tr>
<th>Example</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>prex_cleval</td>
<td>Learning curves</td>
</tr>
<tr>
<td>prex_combining</td>
<td>Classifier combining</td>
</tr>
<tr>
<td>prex_confmat</td>
<td>Confusion matrix, scatterplot and gridsize</td>
</tr>
<tr>
<td>prex_datafile</td>
<td>Datafile usage</td>
</tr>
<tr>
<td>prex_datasets</td>
<td>Show scatter plots of standard datasets</td>
</tr>
<tr>
<td>prex_density</td>
<td>Various density plots</td>
</tr>
<tr>
<td>prex_eigenfaces</td>
<td>Use of images and eigenfaces</td>
</tr>
<tr>
<td>prex_matchlab</td>
<td>Clustering the Iris dataset</td>
</tr>
<tr>
<td>prex_mcplot</td>
<td>Multi-class classifier plot</td>
</tr>
<tr>
<td>prex_plotc</td>
<td>Dataset scatter and classifier plot</td>
</tr>
<tr>
<td>prex_som</td>
<td>Self-organizing map</td>
</tr>
<tr>
<td>prex_spatm</td>
<td>Spatial smoothing of image classification</td>
</tr>
<tr>
<td>prex_cost</td>
<td>Cost matrices and rejection</td>
</tr>
<tr>
<td>prex_logdens</td>
<td>Density based classifier improvement</td>
</tr>
<tr>
<td>prex_regr</td>
<td>Regression example</td>
</tr>
<tr>
<td>Function</td>
<td>Description</td>
</tr>
<tr>
<td>------------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>cdats</td>
<td>Support routine for checking datasets</td>
</tr>
<tr>
<td>iscolumn</td>
<td>Test on column array</td>
</tr>
<tr>
<td>iscomdset</td>
<td>Test on compatible datasets</td>
</tr>
<tr>
<td>isdataim</td>
<td>Test on image dataset</td>
</tr>
<tr>
<td>isdataset</td>
<td>Test on dataset</td>
</tr>
<tr>
<td>isfeatim</td>
<td>Test on feature image dataset</td>
</tr>
<tr>
<td>ismapping</td>
<td>Test on mapping</td>
</tr>
<tr>
<td>isobjim</td>
<td>Test on object image dataset</td>
</tr>
<tr>
<td>isparallel</td>
<td>Test on parallel mapping</td>
</tr>
<tr>
<td>isstacked</td>
<td>Test on stacked mapping</td>
</tr>
<tr>
<td>issym</td>
<td>Test on symmetric matrix</td>
</tr>
<tr>
<td>isvaldset</td>
<td>Test on valid dataset</td>
</tr>
<tr>
<td>matchlablist</td>
<td>Match entries of label lists</td>
</tr>
<tr>
<td>newfig</td>
<td>Control of figures on the screen</td>
</tr>
<tr>
<td>newline</td>
<td>Generate a new line in the command window</td>
</tr>
<tr>
<td>nlabcmp</td>
<td>Compare two label lists and count the differences</td>
</tr>
<tr>
<td>prversion</td>
<td>returns version information on PRTools</td>
</tr>
<tr>
<td>showfigs</td>
<td>Distribute all open figures over screen</td>
</tr>
<tr>
<td>delfigs</td>
<td>Delete all figures</td>
</tr>
</tbody>
</table>
8. Examples

The following examples are available under PRTools. We present here the source codes and the output they generate.

8.1 PREX_CLEVAL Learning curves

help prex_cleval
echo on
    % set desired learning sizes
learnsize = [3 5 10 15 20 30];
    % generate Highleyman's classes
A = gendath([100,100]);
    % define classifiers (untrained)
W = {ldc,qdc,knnc([],1),treec};
    % average error over 10 repetitions
    % test set is complement of training set
E = cleval(A,W,learnsize,10);
    % output E is a structure, specially designed for plotr
plotr(E)

echo off
% Generate 10-dimensional data
A = gendatd([100,100],10);
% Select the training set of 40 = 2x20 objects
% and the test set of 160 = 2x80 objects
[B,C] = gendat(A,0.2);
% Define 5 untrained classifiers, (re)set their names
% w1 is a linear discriminant (LDC) in the space reduced by PCA
w1 = klm([],0.95)*ldc;
  w1 = setname(w1,'klm - ldc');
  % w2 is an LDC on the best (1-NN leave-one-out error) 3 features
w2 = featself([],'NN',3)*ldc;
  w2 = setname(w2,'NN-FFS - ldc');
  % w3 is an LDC on the best (LDC leave-one-out error) 3 features
w3 = featself([],ldc,3)*ldc;
  w3 = setname(w3,'LDC-FFS - ldc');
% w4 is an LDC
w4 = ldc;
  w4 = setname(w4,'ldc');
% w5 is a 1-NN
w5 = knnc([],1);
  w5 = setname(w5,'1-NN');
% Store classifiers in a cell
W = {w1,w2,w3,w4,w5};
% Train them all
V = B*W;
% Test them all
disp([newline 'Errors for individual classifiers'])
testc(C,V);
% Construct combined classifier
VALL = [V(:)];
% Define combiners
WC = {prodc,meanc,medianc,maxc,minc,votec};
% Combine (result is cell array of combined classifiers)
VC = VALL * WC;
% Test them all
disp([newline 'Errors for combining rules'])
testc(C,VC)

Errors for individual classifiers

<table>
<thead>
<tr>
<th>Classifier</th>
<th>Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>klm - ldc</td>
<td>0.125</td>
</tr>
<tr>
<td>NN-FFS - ldc</td>
<td>0.506</td>
</tr>
<tr>
<td>LDC-FFS - ldc</td>
<td>0.100</td>
</tr>
<tr>
<td>ldc</td>
<td>0.094</td>
</tr>
</tbody>
</table>

Errors for combining rules

<table>
<thead>
<tr>
<th>Combiner</th>
<th>Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>Product combiner</td>
<td>0.075</td>
</tr>
<tr>
<td>Mean combiner</td>
<td>0.275</td>
</tr>
<tr>
<td>Median combiner</td>
<td>0.113</td>
</tr>
<tr>
<td>Maximum combiner</td>
<td>0.275</td>
</tr>
<tr>
<td>Minimum combiner</td>
<td>0.094</td>
</tr>
<tr>
<td>Voting combiner</td>
<td>0.088</td>
</tr>
</tbody>
</table>
8.3 PREX_CONFMAT Confusion matrix, scatterplot and gridsize

%PREX_CONFMAT PrTools example confusion matrix, scatterplot and gridsize
% Prtools example code to show the use of confusion matrix,
% scatterplot and gridsize.
help prex_confmat; echo on

% Load 8-class 2D problem
randn('state',1); rand('state',1); a = gendatm;
% Compute the Nearest Mean Classifier
w = nmc(a);
% Scatterplot
figure; gridsize(30); scatterd(a,'legend');
% Plot the classifier
plotc(w);
title([getname(a) ', Gridsize 30']);

gridsize(100);
% Set higher gridsize
figure; scatterd(a,'legend');
plotc(w);
title([getname(a) ', Gridsize 100']);
% Classify training set
d = a*w;
% Look at the confusion matrix and compare it to the scatterplot
confmat(d);

echo off
c = num2str(gridsize);
disp(' ')
disp('Classifier plots are inaccurate for small gridsizes. The standard');
disp('value of 30 is chosen because of the speed, but it is too low to');
disp('ensure good plots. Other gridsizes may be set by gridsize(n).')
disp('Compare the two figures and appreciate the difference.')

Multi–Class Problem, Gridsize 30

Multi–Class Problem, Gridsize 100
8.4 PREX_DENSITY Various density plots

help prex_density
figure
echo on
    % Generate one-class data
    a = gencirc(200);

    % Parzen density estimation
    w = parzendc(a);
    subplot(2,2,1);
    scatterd(a,[10,5]);
    plotm(w);
    title('Parzen Density')
    % 3D density plot
    subplot(2,2,2);
    scatterd(a,[10,5]);
    plotm(w,3);

    % Mixture of Gaussians (5)
    w = mogc(a,5);
    subplot(2,2,3);
    scatterd(a,[10,5]);
    plotm(w);
    title ...  
       ('Mixture of 5 Gaussians')
    % 3D density plot
    subplot(2,2,4);
    scatterd(a,[10,5]);
    plotm(w,3);
    drawnow

disp('Study figure at full screen, shrink and hit return')
pause

figure
% Store four density estimators
W = {qdc udc parzendc mogc};
% generate data
a = +gendath;
% plot densities and estimator name
for j=1:4
    subplot(2,2,j)
    scatterd(a,[10,5])
    plotm(a*W{j})
    title([getname(W{j}) ' density estimation'])
end
echo on
8.5 PREX_EIGENFACES Use of images and eigenfaces

help prex_eigenfaces
echo on
    % load one image for each subject (takes a while)
a = faces([1:40],1);
    % compute eigenfaces
w = pca(a);
    % show them
newfig(1,3); show(w); drawnow
    % project all faces on eigenface space
b = [];
for j = 1:40
    a = faces(j,[1:10]);
b = [b;a*w];
    % don't echo loops
echo off
end
echo on
    % show scatterplot of first two eigenfaces
newfig(2,3)
scatterd(b)
title('Scatterplot on first two eigenfaces')
    % compute leave-one-out error curve
featsizes = [1 2 3 5 7 10 15 20 30 39];
e = zeros(1,length(featsizes));
for j = 1:length(featsizes)
k = featsizes(j);
e(j) = testk(b(:,1:k),1);
echo off
end
echo on
    % plot error curve
newfig(3,3)
plot(featsizes,e)
xlabel('Number of eigenfaces')
ylabel('Error')
echo off

Scatterplot on first two eigenfaces
help prex_matchlab
echo on
rand('state',5);
a = iris;
% Find clusters in Iris dataset.
J1 = kmeans(a,3);
% Finds about the same clusters but labels them
differently due to random initialization.
J2 = kmeans(a,3);
conflam(J1,J2);
% 'best' rotation of label names as
[J3,C] = matchlab(J1,J2);
% confusion matrix is now almost diagonal.
conflam(J1,J3);
% Conversion from J2 to J3: J3 = C(J2,:);
C
echo off

<table>
<thead>
<tr>
<th>Estimated Labels</th>
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<tr>
<td>Totals</td>
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</tbody>
</table>
8.7 PREX-MC PLOT Multi-class classifier plot

help prex_mcplot
echo on
gridsize(100) % generate 2 x 2 normal distributed classes
a = +gendath([20,20]); % data only
b = +gendath([20,20]); % data only
A = [a; b + 5]; % shift 2 over [5,5]
% generate 4-class labels
lab = genlab([20 20 20 20],[1 2 3 4]');
A = dataset(A,lab); % construct dataset
A = setname(A,'4-class dataset')
figure
scatterd(A,'.'); drawnow; % make scatter plot for right size
w = qdc(A); % compute normal densities based quadratic classifier
plotc(w,'col'); drawnow; % plot filled classification regions
hold on;
scatterd(A); % redraw scatter plot
hold off

echo off
help prex_plotc
echo on

% generate Highleyman data
A = gendath([100 100]);
% split in training and test set
[C,D] = gendat(A,[20 20]);
% compute classifiers
w1 = ldc(C);            % linear
w2 = qdc(C);            % quadratic
w3 = parzenc(C);        % Parzen
w4 = lmnc(C,3);         % neural net
% compute and display errors
W = {w1,w2,w3,w4};      % store classifiers in cell
disp(D*W*testc);        % plot errors
% plot data and classifiers
figure
scatterd(A);            % scatterplot
plotc({w1,w2,w3,w4});   % plot classifiers

echo off
help prex_spatm
echo on
    % load EM image
    a = emim31;
    % extract small training set
    b = gendat(a,500);
    % use it for finding 3 clusters
    [d,w] = emclust(b,nmc,3);
    % classify entire image and show it
    c = a*w;
    classim(c);
    title('Original classification')
    % smooth image,
    % combine spectral and spatial classifier, show it
    e = spatm(c)*maxc;
    figure
    classim(e);
    title('Smoothed classification')

% echo off
8.10 PREX_COSTM PRTools example on cost matrices and rejection

Prttools example code to show the use of cost matrices and how to introduce a reject class.

% Generate a three class problem
randn('state',1);
rand('state',1);
n = 30;
class_labels = char('apple','pear','banana');
a = [gendatb([n,n]);  gauss(n,[-2 6])];
laba = genlab([n n n],class_labels);
a = setlabels(a,laba);
% Compute a simple ldc
w = ldc(a);
% Scatterplot and classifier
figure;
gridsize(30);
scatterd(a,'legend');
plotc(w);
% Define a classifier with a new cost matrix,
% which puts a high cost on misclassifying
% pears to apples
cost = [0.0 1.0 1.0;
       9.0 0.0 1.0;
       1.0 1.0 0.0];
w = w*classc*costm([],cost,class_labels);
plotc(w,'b');
% Define a classifier with a cost matrix where
% an outlier class is introduced. For this an
% extra column in the cost matrix has to be defined.
% Furthermore, the class labels have to be supplied
% to give the new class a name.
cost = [0.0 1.0 1.0 0.2;
       9.0 0.0 1.0 0.2;
       1.0 1.0 0.0 0.2];
class_labels = char('apple','pear','banana','reject');
w = w*classc*costm([],cost,class_labels);
plotc(w,'--')

The black decision boundary shows the standard ldc classifier for this data. When the misclassification cost of a pear to an apple is increased, we obtain the blue classifier. When on top of that a rejection class is introduced, we get the blue dashed classifier. In that case, all objects between the dashed lines are rejected.

Cost of basic classifier = 0.51
Cost of cost classifier = 0.24
Cost of reject classifier = 0.10
8.11 PREX_LOGDENS Improving density based classifiers

This example shows the use and results of LOGDENS for improving the classification in the tail of the distributions

```matlab
% Generate a small two-class problem
randn('state',1);
rand('state',1);
a = gendatb([20 20]);
% Compute two classifiers: Mixture of Gaussians and Parzen
w_mogc = mogc(a); w_mogc = setname(w_mogc,'MoG');
w_parz = parzenc(a); w_parz = setname(w_parz,'Parzen');
% Scatterplot with MoG classifier
subplot(3,2,1);
scatter(a);
plotc(w_mogc); xlabel(''); ylabel('');
set(gca,'xtick',[],'ytick',[])
title('MoG density classifier','fontsize',12)
drawnow
% Scatterplot with Parzen classifier
subplot(3,2,2);
scatter(a);
plotc(w_parz); xlabel(''); ylabel('');
set(gca,'xtick',[],'ytick',[])
title('Parzen density classifier','fontsize',12)
drawnow
% Scatterplot from a distance :
% far away points are inaccurately classified
subplot(3,2,3);
scatterd([a; [150 100]; [-150 -100]]);
plotc(w_mogc); xlabel(''); ylabel('');
set(gca,'xtick',[],'ytick',[])
title('MoG: bad for remote points','fontsize',12)
drawnow
% Scatterplot from a distance :
% far away points are inaccurately classified
subplot(3,2,4);
scatterd([a; [20 12]; [-20 -12]]);
plotc(w_parz); xlabel(''); ylabel('');
set(gca,'xtick',[],'ytick',[])
title('Parzen: bad for remote points','fontsize',12)
drawnow
% Improvement of MOGC by LOGDENS
subplot(3,2,5);
scatterd([a; [150 100]; [-150 -100]]);
plotc([w_mogc,logdens(w_mogc)],['k--';'r- ']); legend off
xlabel(''); ylabel(''); set(gca,'xtick',[],'ytick',[])
title('MoG improved by Log-densities','fontsize',12)
drawnow
```
This example shows the use of the logdens() routine. It improves the classification in the tails of the distribution, which is especially important in high-dimensional spaces. To this end it is combined with normalization, generating posterior probabilities. Logdens() can only be applied to classifiers based on normal densities and Parzen estimates.
9. PRTools 4.0 release notes

This section supplies some information about changes in PRTools4.0 with respect to the PRTools3.1 versions. Changes are major and sometimes incompatible. A number of changes only involve the fundamental definitions, but are not is not yet implemented on the user level.

9.1 Datasets

The dataset construct has been entirely redefined and rewritten. See datasets (section 5.2) for an online description. Many fields are added. There are separate commands for setting and getting each field separately like setlabels(A,labels).

The main change for the user is that there are three different types of labels supported: ‘crisp’ (as it was), ‘soft’ (on the [0,1] interval) and ‘targets’ (a multidimensional vector for each object).

In the present state all higher level commands work for crisp labels and some for soft labels (e.g. for normal distributions) but nothing for targets. Also checking for appropriate labels is not done yet. As long as crisp labels are most routines work like before.

A new system has been created for keeping track of images stored as features or objects. In the size fields of datasets the image sizes are stored.

Datasets, classes and features may have names that are used to annotate plots.

During creation of a dataset objects are given a unique identifier, that is not changed anymore by PRTools. This enables the user to retrieve the original object from, for instance, the classification dataset, also after random selection of a test set. See setident, getident, findident and seldat.

Objects may be unlabeled. Such objects are not used for training classifiers.

For features domains may be defined for their values. Checking is done when dataset values or domain definitions change. See setfeatdom.

Programmers have to take care that all needed information is passed from one dataset to the other. The best thing to do is to ‘copy’ old datasets and create a new one by changing the data, e.g. B = setdata(A,data,featlab) creates B out of A with new data and new names for the features, assuming that we have the same objects, object labels, prior probabilities, etcetera.

9.2 Mappings

The mapping construct has been redefined and rewritten as well. See mappings for online information. Now a clear distinction is made between four types of mappings: untrained, trained, fixed and combiner. In the mapping definition the programmer has to specify the type explicitly. PRTools has to know about these types as they are treated differently:

- untrained mapping cannot map data, but define the choice of the mapping and contain some parameter choices, e.g. \( W = \text{ldc}([],1e^{-6}) \) defines a regularization value. Untrained mappings are useful for routines like cleval and featself that evaluate or use arbitrary untrained classifiers. \( V = A*W \) produces a ‘trained’ mapping. How training (and also execution) of mappings
is done is not hidden anymore for the user. Each mapping definition contains a `mapping_file` field that points to the file by which this further processing is performed.

- **trained mappings** map a dataset from one space to another, so \( D = B*V \) maps the dataset \( B \) by a trained classifier \( V \) from the feature space to a ‘classification’ space: each object has values for each class, e.g. a distance, a density, a posterior probability, a membership, etcetera. Routines for trained mappings typically have three ways they are called by PRTools and thereby have three program sections: the untrained call or definition, the training and the execution. See `kernelm` for a typical example. Sometimes execution is shared by some routines, e.g. `normal_map` handles all execution of normal densities based mappings.

- **fixed mappings** are like trained mappings but don’t find their parameter values by training. Instead, they are set by other routines or by the user. As a result they don’t have a part for training. So if \( W = \text{sigm}([], p) \), defining the sigmoid mapping, then \( W \) is called fixed (and not untrained) as \( A*W \) results in a dataset \( B = \text{sigm}(A, p) \) and not in a trained mapping.

- **combiners** are mappings that know how to handle other mappings. If \( V \) is a mapping and \( W \) is a combiner (e.g. \( W = \text{maxc} \)) then \( V*W \) results in a call like \( U = \text{maxc}(V) \), in which \( U \) is an untrained or a trained mapping, dependent on \( V \). If \( W \) is not a combiner, then \( V*W \) is stored as such in \( U \) (called a sequential mapping, which again can be trained or untrained) and execution is postponed until a dataset has to be processed by \( A*U = A*(V*W) \). How this is done depends on the mapping types of \( V \) and \( W \).

All the above is not really of importance for the users of PRTools, but just for programmers that like to write new mappings. For some users it may be of interest that the overload of the ‘\*’ operator can always be avoided by `map()`, e.g. \( V = A*W \) is identical to \( V = \text{map}(A, W) \).

The use of prior probabilities is now restricted to density based classifiers and the computation of means and covariance matrices over classes. If this has to be avoided, use \( A = \text{setprior}(A, []) \), by which class priors are made identical to class frequencies.

### 9.3 The user level

The old set of user routines has been corrected for the new definitions of datasets and mappings. During this revision some old constructs have been upgraded or removed. Some routines have been simplified (like `testc`, the new version of `testc`). Also `plotd` has been renamed to `plotc` for more consistency: `plotc` plots classifiers, `plotm` mappings (densities). Plotting routines have been extended and another default font size is introduced. On the whole, PRTools should behave about the same as before on the user level. Existing macros, however, have to be checked for sure.

Important for users is that mappings like \( B*\text{fisherc}(A) \) now output unnormalized posterior probability estimates (class memberships) or for density based classifiers (\( B*\text{qdc}(A) \)) the true density. So this output is always positive. The routine `classc` takes care of normalization, converting outputs into proper posterior estimates: \( B*\text{lmc}(A)*\text{classc}, \) or \( B*\text{qdc}(A)*\text{classc} \). This new implementation may result in accuracy problems as densities may suffer from underflows in large areas of the feature space. For the normal density based classifiers like `ldc`, `qdc` and `udc` this can be circumvented by the use of `logdens` in the classifier definition (e.g. \( B*(\text{qdc}(A)*\text{logdens}) \)). In that case log-densities are stored instead of densities.
10. PRTools 4.1 release notes

This section supplies some information about changes in PRTools 4.1 with respect to PRTools 4.0. A number of new possibilities have been created important for the handling of large datasets, multiple labels of objects, the optimisation of complexity and regularisation parameters and the handling of regression problems.

10.1 Compatibility

Changes are generally upwards compatible. With a few exceptions old routines should still work. The main exception is that the undocumented feature of PRTools 4.0 to obtain fields from dataset and mapping variables using the dot-construct (e.g. classnames = a.lablist) has been changed. From now on the official and guaranteed way to address fields is by using the get-commands (e.g. classnames = getlablist(a)). The reason is that for a number of fields subfields have been defined using structures, structure arrays and cell arrays. So users are urged to use the get-and set-commands as also in future releases the constructions may change. PRTools still recognizes datasets contructed in the old way and automatically converts them.

10.2 Datafiles

A new object class, datafile, has been created. The datafile class inherits most of the fields and methods of the dataset class, but extends them by allowing data that is not in core but stored in files on disk. As these may be large, handling of datafiles is restricted to administration, like desired sampling of objects and features and preprocessing (e.g. filtering and resizing of images). At some moment a datafile has to be converted into a dataset and it should fit then in the available memory. Datafiles are important to the extend PRTools possibilities with preprocessing and feature measurements within the same framework. Thereby classifiers may be designed and trained that can directly operate on raw images or other signals without the need to convert them first to datasets. For more information read datafiles help file.

10.3 Image processing routines

In relation with the above a large set of image processing routines operating on datafiles and datasets has been included. They are helpful to convert (sets of) images to features and datasets. A number of them assume that the dip_image toolbox is available.

10.4 Multiple labels

For some applications it is useful to have multiple labelings of the objects. For instance, pixels may be labeled according to the image region (grass, water, rock) as well as to the image category (mountains, seaside, city) as well as to some origin (France, England, Norway). A provision has been created to enable this. The various labelings and corresponding priors (and targets in case of soft labeling) are stored in the dataset, but just one of them is active and is accessed by getlablist, getlabels, getprior and getnlab. For more information read the multi_labeling help file.
10.5 Optimisation of complexity parameters and regularisation

Many trainable classifiers and mappings depend on some parameter controlling its complexity or regularisation. A general routine has been created to optimise such parameters by cross-validation. This is always done in a standard way: 20 steps of 5-fold cross-validation. This increases the training time roughly by a factor 100. The automatic optimisation is activated (for the routines for which it is implemented) by using a NaN in the function call. So \( w=\text{ldc}(a) \) uses no regularisation, \( w=\text{ldc}(a,1e-6) \) uses a user defined value of 1e-6 and \( w=\text{ldc}(a,\text{NaN}) \) activates the automatic optimisation. The actually used parameter value may be retrieved afterwards by the routine \text{getopt_pars}. 

10.6 Regression

PRTools has already for a long time the possibility of datasets consisting of feature based vectors with one or more desired target values (the have the label type ‘targets’ instead of ‘crisp’ or ‘soft’). Now a set of routines has been added to make use of this option, e.g. \text{linarr} for linear regression, \text{svmr} for support vector machine regression and \text{testr} of evaluation.

10.7 Object and dataset annotation

Datasets and objects inside datasets have fields to annotate them (‘user’ and ‘ident’). They are now structures and the routines for setting (\text{setuser} and \text{setident}) and reading (\text{getuser} and \text{getident}) can handle them.

10.8 Kernels

A general routine has been added for defining kernels: \text{kernelm}. This routine may be used in the support vector classifiers \text{svec} and \text{nusvc} as well as in the general kernel based classifier \text{kernelec}.

10.9 Support vector classifiers

The call of the general support vector classifier \text{svec} has been upgraded such that it included a recognizable kernel definition. In addition two other support vector classifiers are added, \text{nusvc} for using a regularisation parameter based on the expected error and \text{rbsvc}, which is parameter free as it estimates automatically the optimal radial basis kernel and the regularisation parameter.

10.10 Rejects

Routines \text{rejectc} and \text{rejectm} have been added to facilitate the construction of rejecting classifiers. They add class labels on the output that are NaN or ‘’ (empty string), the PRTools standard label for unlabeled objects.