TMVA 4
Toolkit for Multivariate Data Analysis with ROOT

Users Guide

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Abstract — In high-energy physics, with the search for ever smaller signals in ever larger data sets, it has become essential to extract a maximum of the available information from the data. Multivariate classification methods based on machine learning techniques have become a fundamental ingredient to most analyses. Also the multivariate classifiers themselves have significantly evolved in recent years. Statisticians have found new ways to tune and to combine classifiers to further gain in performance. Integrated into the analysis framework ROOT, TMVA is a toolkit which hosts a large variety of multivariate classification algorithms. Training, testing, performance evaluation and application of all available classifiers is carried out simultaneously via user-friendly interfaces. With version 4, TMVA has been extended to multivariate regression of a real-valued target vector. Regression is invoked through the same user interfaces as classification. TMVA 4 also features more flexible data handling allowing one to arbitrarily form combined MVA methods. A generalised boosting method is the first realisation benefiting from the new framework.

TMVA 4.3.0 for ROOT >= 6.12/00 — Toolkit for Multivariate Data Analysis with ROOT
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1 Introduction

The Toolkit for Multivariate Analysis (TMVA) provides a ROOT-integrated [1] environment for the processing, parallel evaluation and application of multivariate classification and – since TMVA version 4 – multivariate regression techniques. All multivariate techniques in TMVA belong to the family of “supervised learning” algorithms. They make use of training events, for which the desired output is known, to determine the mapping function that either describes a decision boundary (classification) or an approximation of the underlying functional behaviour defining the target value (regression). The mapping function can contain various degrees of approximations and may be a single global function, or a set of local models. TMVA is specifically designed for the needs of high-energy physics (HEP) applications, but should not be restricted to these. The package includes:

- Rectangular cut optimisation (binary splits, Sec. 8.1).
- Projective likelihood estimation (Sec. 8.2).
- Multi-dimensional likelihood estimation (PDE range-search – Sec. 8.3, PDE-Foam – Sec. 8.4, and k-NN – Sec. 8.5).
- Linear and nonlinear discriminant analysis (H-Matrix – Sec. 8.6, Fisher – Sec. 8.7, LD – Sec. 8.8, FDA – Sec. 8.9).
- Artificial neural networks (three different multilayer perceptron implementations – Sec. 8.10).
- Support vector machine (Sec. 8.12).
- Boosted/bagged decision trees (Sec. 8.13).
- Predictive learning via rule ensembles (RuleFit, Sec. 8.14).
- A generic boost classifier allowing one to boost any of the above classifiers (Sec. 10).
- A generic category classifier allowing one to split the training data into disjoint categories with independent MVAs.

The software package consists of abstract, object-oriented implementations in C++/ROOT for each of these multivariate analysis (MVA) techniques, as well as auxiliary tools such as parameter fitting and transformations. It provides training, testing and performance evaluation algorithms

---

1 A classification problem corresponds in more general terms to a discretised regression problem. A regression is the process that estimates the parameter values of a function, which predicts the value of a response variable (or vector) in terms of the values of other variables (the input variables). A typical regression problem in High-Energy Physics is for example the estimation of the energy of a (hadronic) calorimeter cluster from the cluster’s electromagnetic cell energies. The user provides a single dataset that contains the input variables and one or more target variables. The interface to defining the input and target variables, the booking of the multivariate methods, their training and testing is very similar to the syntax in classification problems. Communication between the user and TMVA proceeds conveniently via the Factory and Reader classes. Due to their similarity, classification and regression are introduced together in this Users Guide. Where necessary, differences are pointed out.
and visualisation scripts. Detailed descriptions of all the TMVA methods and their options for classification and (where available) regression tasks are given in Sec. 8. Their training and testing is performed with the use of user-supplied data sets in form of ROOT trees or text files, where each event can have an individual weight. The true sample composition (for event classification) or target value (for regression) in these data sets must be supplied for each event. Preselection requirements and transformations can be applied to input data. TMVA supports the use of variable combinations and formulas with a functionality similar to the one available for the \texttt{Draw} command of a ROOT tree.

TMVA works in transparent factory mode to provide an objective performance comparison between the MVA methods: all methods see the same training and test data, and are evaluated following the same prescriptions within the same execution job. A \textit{Factory} class organises the interaction between the user and the TMVA analysis steps. It performs preanalysis and preprocessing of the training data to assess basic properties of the discriminating variables used as inputs to the classifiers. The linear correlation coefficients of the input variables are calculated and displayed. For regression, also nonlinear correlation measures are given, such as the correlation ratio and mutual information between input variables and output target. A preliminary ranking is derived, which is later superseded by algorithm-specific variable rankings. For classification problems, the variables can be linearly transformed (individually for each MVA method) into a non-correlated variable space, projected upon their principle components, or transformed into a normalised Gaussian shape. Transformations can also be arbitrarily concatenated.

To compare the signal-efficiency and background-rejection performance of the classifiers, or the average variance between regression target and estimation, the analysis job prints – among other criteria – tabulated results for some benchmark values (see Sec. 3.1.9). Moreover, a variety of graphical evaluation information acquired during the training, testing and evaluation phases is stored in a ROOT output file. These results can be displayed using macros, which are conveniently executed via graphical user interfaces (each one for classification and regression) that come with the TMVA distribution (see Sec. 3.3).

The TMVA training job runs alternatively as a ROOT script, as a standalone executable, or as a python script via the PyROOT interface. Each MVA method trained in one of these applications writes its configuration and training results in a result ("weight") file, which in the default configuration has human readable XML format.

A light-weight \textit{Reader} class is provided, which reads and interprets the weight files (interfaced by the corresponding method), and which can be included in any C++ executable, ROOT macro, or python analysis job (see Sec. 3.4).

For standalone use of the trained MVA method, TMVA also generates lightweight C++ response classes (not available for all methods), which contain the encoded information from the weight files so that these are not required anymore. These classes do not depend on TMVA or ROOT, neither on any other external library (see Sec. 3.5).

We have put emphasis on the clarity and functionality of the Factory and Reader interfaces to the user applications, which will hardly exceed a few lines of code. All MVA methods run with reasonable default configurations and should have satisfying performance for average applications. \textit{We stress}
however that, to solve a concrete problem, all methods require at least some specific tuning to deploy their maximum classification or regression capabilities. Individual optimisation and customisation of the classifiers is achieved via configuration strings when booking a method.

This manual introduces the TMVA Factory and Reader interfaces, and describes design and implementation of the MVA methods. It is not the aim here to provide a general introduction to MVA techniques. Other excellent reviews exist on this subject (see, e.g., Refs. [2–4]). The document begins with a quick TMVA start reference in Sec. 2, and provides a more complete introduction to the TMVA design and its functionality for both, classification and regression analyses in Sec. 3. Data preprocessing such as the transformation of input variables and event sorting are discussed in Sec. 4. In Sec. 5, we describe the techniques used to estimate probability density functions from the training data. Section 6 introduces optimisation and fitting tools commonly used by the methods. All the TMVA methods including their configurations and tuning options are described in Secs. 8.1–8.14. Guidance on which MVA method to use for varying problems and input conditions is given in Sec. 11. An overall summary of the implementation status of all TMVA methods is provided in Sec. 12.

Copyrights and credits

TMVA is an open source product. Redistribution and use of TMVA in source and binary forms, with or without modification, are permitted according to the terms listed in the BSD license.\(^2\) Several similar combined multivariate analysis (“machine learning”) packages exist with rising importance in most fields of science and industry. In the HEP community the package StatPatternRecognition [5, 6] is in use (for classification problems only). The idea of parallel training and evaluation of MVA-based classification in HEP has been pioneered by the Cornelius package, developed by the Tagging Group of the BABAR Collaboration [7]. See further credits and acknowledgments on page 161.

\(^2\)For the BSD license, see http://tmva.sourceforge.net/LICENSE.
2 TMVA Quick Start

To run TMVA it is not necessary to know much about its concepts or to understand the detailed functionality of the multivariate methods. Better, just begin with the quick start tutorial given below.

Classification and regression analyses in TMVA have similar training, testing and evaluation phases, and will be treated in parallel in the following.

2.1 How to download and build TMVA

TMVA is built upon ROOT (https://root.cern), so that for TMVA to run ROOT must be installed. Since ROOT version 5.11/06, TMVA is distributed as an integral part of ROOT and can be used from the ROOT prompt without further preparation.

The latest version of ROOT can be found at https://root.cern/downloading-root. The advanced user, should she require it, can build ROOT and TMVA from the source distribution. Instructions are found at https://root.cern/building-root.

2.2 Version compatibility

TMVA can be run with any ROOT version equal or above v5.08. The few occurring conflicts due to ROOT source code evolution after v5.08 are intercepted in TMVA via C++ preprocessor conditions.

2.3 The TMVA namespace

All TMVA classes are embedded in the namespace `TMVA`. For interactive access, or use in macros, the classes must thus be preceded by `TMVA::`.

One may also use the command `using namespace TMVA;` instead. However, this approach is not preferred since it can lead to ambiguities.

2.4 Example jobs

TMVA comes with example jobs for the training phase (this phase actually includes training, testing and evaluation) using the TMVA `Factory`, as well as the application of the training results in a classification or regression analysis using the TMVA `Reader`. The training examples are `TMVAClassification.C`, `TMVAMulticlass.C` and `TMVARegression.C`, and the application examples are `TMVAClassificationApplication.C`, `TMVAMulticlassApplication.C` and `TMVARegressionApplication.C`.
2.5 Running the example

Additionally TMVACrossValidation.C shows how cross validation is performed in TMVA, and examples in $ROOTSYS/tutorials/tmva/test/keras shows how to use TMVA with keras.

The above macros (extension .C) are located in the directory $ROOTSYS/tutorials/tmva/test.

Some examples are also provided in form of C++ executables (replace .C by .cxx). To build the executables, go to $ROOTSYS/tutorials/tmva/test, type make and then simply execute them by typing ./TMVAClassification, ./TMVAMulticlass or ./TMVARegression (and similarly for the applications).

2.5 Running the example

The most straightforward way to get started with TMVA is to simply run the TMVAClassification.C or TMVARegression.C example macros. Both use academic toy datasets for training and testing, which, for classification, consists of four linearly correlated, Gaussian distributed discriminating input variables, with different sample means for signal and background, and, for regression, has two input variables with fuzzy parabolic dependence on the target (fvalue), and no correlations among themselves. All classifiers are trained, tested and evaluated using the toy datasets in the same way the user is expected to proceed for his or her own data. It is a valuable exercise to look at the example file in more detail. Most of the command lines therein should be self explaining, and one will easily find how they need to be customized to apply TMVA to a real use case. A detailed description is given in Sec. 3.

The macros automatically fetch the data file from the web using the corresponding TFile constructor, e.g., TFile::Open("http://root.cern/files/tmva_class_example.root") (tmva_reg_example.root for regression). The example ROOT macros can be run directly in the $ROOTSYS/tutorials/tmva/test directory, or from anywhere after adding the macro directory to ROOT’s macro search path:

```
> root -l $ROOTSYS/tutorials/tmva/TMVAClassification.C
> echo "Unix.*.Root.MacroPath: $ROOTSYS/tutorials/tmva" >> ~/.rootrc
> root -l TMVAClassification.C
```

**Code Example 1:** Running the example TMVAClassification.C.

It is also possible to explicitly select the MVA methods to be processed:

```
> root -l '$ROOTSYS/tutorials/tmva/TMVAClassification.C("Fisher","BDT")'
```

**Code Example 2:** Running the example TMVAClassification.C and processing only the Fisher and BDT-classifier. Multiple classifiers are separated by commas. The others macros can be called accordingly.
where the names of the MVA methods are predefined in the macro.

The training job provides formatted output logging containing analysis information such as: linear correlation matrices for the input variables, correlation ratios and mutual information (see below) between input variables and regression targets, variable ranking, summaries of the MVA configurations, goodness-of-fit evaluation for PDFs (if requested), signal and background (or regression target) correlations between the various MVA methods, decision overlaps, signal efficiencies at benchmark background rejection rates (classification) or deviations from target (regression), as well as other performance estimators. Comparison between the results for training and independent test samples provides overtraining validation.

### 2.6 Displaying the results

After a successful training, TMVA provides so called “weight”-files that contain all information necessary to recreate the method without retraining. For further information, see the `TMVA::Reader` in Section 3.4. TMVA also provides a variety of control and performance plots that can be displayed via a dedicated GUI (see Fig. 1). The GUI can be launched with

```
> root -l -e 'TMVA::TMVAGui("/path/to/tmva/output/file.root")'
> root -l -e 'TMVA::TMVAMultiClassGui("/path/to/tmva/output/file.root")'
> root -l -e 'TMVA::TMVARegGui("/path/to/tmva/output/file.root")'
```

**Code Example 3:** Running the TMVA GUI's from the command line.

or through the adding line

```
if (not gROOT->IsBatch()) {
    TMVA::TMVAGui("/path/to/tmva/output/file.root");
}
```

**Code Example 4:** Automatically running a GUI when doing training/testing/evaluation.

to your training file. See the last lines of `TMVAClassification.C` for an example.

Examples for plots produced by these macros are given in Figs. 4–6 for a classification problem. The distributions of the input variables for signal and background according to our example job are shown in Fig. 3. It is useful to quantify the correlations between the input variables. These are drawn in form of a scatter plot with the superimposed profile for two of the input variables in Fig. 4 (upper left). As will be discussed in Sec. 4, TMVA allows to perform a linear decorrelation transformation of the input variables prior to the MVA training (for classification only). The result of such decorrelation is shown at the upper right hand plot of Fig. 4. The lower plots display the
Figure 1: Graphical user interfaces (GUI) to execute macros displaying training, test and evaluation results (cf. Tables 2 and 4 on page 42) for classification (left) and regression problems (right). The multiclass classification GUI is similar to the classification GUI, where some functionality is not available. The classification GUI can be launched manually by executing the script 
\$ROOTSYS/tmva/test/TMVAGui.C\n
in a ROOT session. To launch the multiclass or regression GUIs use the macros TMVARegGui.C or TMVAMultiClassGui.C, respectively.

Classification (left). The buttons behave as follows: (1a) plots the signal and background distributions of input variables (training sample), (1b–d) the same after applying the corresponding preprocessing transformation of the input variables, (2a–f) scatter plots with superimposed profiles for all pairs of input variables for signal and background and the applied transformations (training sample), (3) correlation coefficients between the input variables for signal and background (training sample), (4a/b) signal and background distributions for the trained classifiers (test sample/test and training samples superimposed to probe overtraining), (4c,d) the corresponding probability and Rarity distributions of the classifiers (where requested, cf. see Sec. 3.1.13), (5a) signal and background efficiencies and purities versus the cut on the classifier output for the expected numbers of signal and background events (before applying the cut) given by the user (an input dialog box pops up, where the numbers are inserted), (5b) background rejection versus signal efficiency obtained when cutting on the classifier outputs (ROC curve, from the test sample), (6) plot of so-called Parallel Coordinates visualising the correlations among the input variables, and among the classifier and the input variables, (7–13) show classifier specific diagnostic plots, and (14) quits the GUI. Titles greyed out indicate actions that are not available because the corresponding classifier has not been trained or because the transformation was not requested.

Regression (right). The buttons behave as follows: (1–3) same as for classification GUI, (4a–d) show the linear deviations between regression targets and estimates versus the targets or input variables for the test and training samples, respectively, (5) compares the average deviations between target and MVA output for the trained methods, and (6–8) are as for the classification GUI.
Figure 2: Graphical user interfaces (GUI) to execute macros displaying training, test and evaluation results (cf. Tables 2 and 4 on page 42) for multiclass classification (left). The multiclass classification GUI can be started by executing `root-e'^TMVA::TMVAMultiClassGui()’ at the prompt. To the right a sample output from the 1 vs. Rest ROC macro can be seen.

Multiclass classification (left). The main difference between the multiclass GUI and the binary one is the splitting of button (5b) of `TMVAGui.C into button (5) and (6) in `TMVAMultiClassGui.C. Button (5) draws 1 vs. Rest ROC curves where each class, in turn, is considered signal while the remaining are considered background. Button (6) instead considers all pairs of classes in turn, ignoring events belonging to other classes. These two visualisation provide a simplified view of of the high-dimensional ROC surface of multiclass problems.

1 vs. 1 ROC curve (right). Example of one of the plots generated for the 1 vs. Rest ROC curve button (5).

linear correlation coefficients between all input variables, for the signal and background training samples of the classification example.

Figure 5 shows several classifier output distributions for signal and background events based on the test sample. By TMVA convention, signal (background) events accumulate at large (small) classifier output values. Hence, cutting on the output and retaining the events with $y$ larger than the cut requirement selects signal samples with efficiencies and purities that respectively decrease and increase with the cut value. The resulting relations between background rejection versus signal efficiency are shown in Fig. 6 for all classifiers that were used in the example macro. This plot belongs to the class of Receiver Operating Characteristic (ROC) diagrams, which in its standard form shows the true positive rate versus the false positive rate for the different possible cutpoints of a hypothesis test.

For multiclass classification the corresponding GUI is show in Figure 2 (left). Some differences to the binary classification GUI exists. The ROC space is $C(C - 1)$-dimensional, i.e. proportional to the number of classes squared. Visualising this potentially high dimensional space quickly becomes intractable as the number of dimensions increases. To still gain insight in the performance of
the classifier, TMVA offers two projections of the ROC surface into 2D. In 1 vs. Rest, $C$ plots are generated where each class, in turn, is considered signal while the remaining are considered background. In 1 vs. 1, $C(C-1)$ plots are generated where each pair of classes are considered. Events which do not belong to either of the two classes are ignored for the calculation of that ROC curve. Example output for one classes in 1 vs. Rest is shown in Figure 2 (right).

As an example for multivariate regression, Fig. 7 displays the deviation between the regression output and target values for linear and nonlinear regression algorithms.

More macros are available to validate training and response of specific MVA methods. For example, the macro `likelihoodref.C` compares the probability density functions used by the likelihood classifier to the normalised variable distributions of the training sample. It is also possible to

Figure 3: Example plots for input variable distributions. The histogram limits are chosen to zoom into the bulk of the distributions, which may lead to truncated tails. The vertical text on the right-hand side of the plots indicates the under- and overflows. The limits in terms of multiples of the distribution’s RMS can be adjusted in the user script by modifying the variable `TMVA::gConfig().GetVariablePlotting()`. TimesRMS (cf. Code Example 29).
2.7 Getting help

Several help sources exist for TMVA (all web address given below are also linked from the TMVA home page http://tmva.sourceforge.net).

- TMVA tutorial: https://twiki.cern.ch/twiki/bin/view/TMVA.

- On request, the TMVA methods provide a help message with a brief description of the method, and hints for improving the performance by tuning the available configuration options. The message is printed when the option "H" is added to the configuration string while booking the method (switch off by setting "!H").
Figure 5: Example plots for classifier output distributions for signal and background events from the academic test sample. Shown are likelihood (upper left), PDER range search (upper right), Multilayer perceptron (MLP – lower left) and boosted decision trees.

- Please ask questions and/or report problems in the ROOT forum https://root-forum.cern.ch.

3 Using TMVA

A typical TMVA classification or regression analysis consists of two independent phases: the training phase, where the multivariate methods are trained, tested and evaluated, and an application phase, where the chosen methods are applied to the concrete classification or regression problem they have been trained for. An overview of the code flow for these two phases as implemented in the examples TMVAClassification.C and TMVAClassificationApplication.C (for classification – see Sec. 2.4),
and TMVAREgression.C and TMVAREgressionApplication.C (for regression) are sketched in Fig. 8. Multiclass classification does not differ much from two class classification from a technical point of view and differences will only be highlighted where necessary.

In the training phase, the communication of the user with the data sets and the MVA methods is performed via a Factory object, created at the beginning of the program. The TMVA Factory provides member functions to specify the training and test data sets, to register the discriminating input and – in case of regression – target variables, and to book the multivariate methods. Subsequently the Factory calls for training, testing and the evaluation of the booked MVA methods. Specific result (“weight”) files are created after the training phase by each booked MVA method.

The application of training results to a data set with unknown sample composition (classification) / target value (regression) is governed by the Reader object. During initialisation, the user registers the input variables together with their local memory addresses, and books the MVA methods that were found to be the most appropriate after evaluating the training results. As booking argument, the name of the weight file is given. The weight file provides for each of the methods full and consistent configuration according to the training setup and results. Within the event loop, the input variables are updated for each event and the MVA response values are computed. Some methods also provide the computation of errors.

For standalone use of the trained MVA methods, TMVA also generates lightweight C++ response classes, which contain the encoded information from the weight files so that these are not required anymore (cf. Sec. 3.5).

---

3This somewhat redundant operation is required to verify the correspondence between the Reader analysis and the weight files used.
3.1 The TMVA Factory

The TMVA training phase begins by instantiating a Factory object with configuration options listed in Table 1.

```
TMVA::Factory* factory = new TMVA::Factory( "<JobName>", outputFile, "<options>" );
```

Code Example 5: Instantiating a Factory class object. The first argument is the user-defined job name that will reappear in the names of the weight files containing the training results. The second argument is the pointer to a writable TFile output file created by the user, where control and performance histograms are stored.

### 3.1.1 Specifying training and test data

The input data sets used for training and testing of the multivariate methods need to be handed to the Factory. TMVA supports ROOT TTree and derived TChain objects as well as text files. If ROOT trees are used for classification problems, the signal and background events can be located in the same or in different trees. Data trees can be provided specifically for the purpose of either training or testing or for both purposes. In the latter case the factory then splits the tree into one part for training, the other for testing (see also section 3.1.4).

Overall weights can be specified for the signal and background training data (the treatment of event-by-event weights is discussed below).

Specifying **classification training and test data** in ROOT tree format with signal and background events being located in different trees:
Figure 8: Left: Flow (top to bottom) of a typical TMVA training application. The user script can be a ROOT macro, C++ executable, python script or similar. The user creates a ROOT TFile, which is used by the TMVA Factory to store output histograms and trees. After creation by the user, the Factory organises the user’s interaction with the TMVA modules. It is the only TMVA object directly created and owned by the user. First the discriminating variables that must be TFormula-compliant functions of branches in the training trees are registered. For regression also the target variable must be specified. Then, selected MVA methods are booked through a type identifier and a user-defined unique name, and configuration options are specified via an option string. The TMVA analysis proceeds by consecutively calling the training, testing and performance evaluation methods of the Factory. The training results for all booked methods are written to custom weight files in XML format and the evaluation histograms are stored in the output file. They can be analysed with specific macros that come with TMVA (cf. Tables 2 and 4).

Right: Flow (top to bottom) of a typical TMVA analysis application. The MVA methods qualified by the preceding training and evaluation step are now used to classify data of unknown signal and background composition or to predict a regression target. First, a Reader class object is created, which serves as interface to the method’s response, just as was the Factory for the training and performance evaluation. The discriminating variables and references to locally declared memory placeholders are registered with the Reader. The variable names and types must be equal to those used for the training. The selected MVA methods are booked with their weight files in the argument, which fully configures them. The user then runs the event loop, where for each event the values of the input variables are copied to the reserved memory addresses, and the MVA response values (and in some cases errors) are computed.
### Option Table 1: Configuration options reference for class: Factory.

Coloured output is switched on by default, except when running ROOT in batch mode (i.e., when the `-b` option of the CINT interpreter is invoked). The list of transformations contains a default set of data preprocessing steps for test and visualisation purposes only. The usage of preprocessing transformations in conjunction with MVA methods must be configured when booking the methods.

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>V</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Verbose flag</td>
</tr>
<tr>
<td>Color</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Flag for coloured screen output (default: True, if in batch mode: False)</td>
</tr>
<tr>
<td>Transformations</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>List of transformations to test; formatting example: Transformations=I;D;P;U;G,D, for identity, decorrelation, PCA, Uniform and Gaussianisation followed by decorrelation transformations</td>
</tr>
<tr>
<td>Silent</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Batch mode: boolean silent flag inhibiting any output from TMVA after the creation of the factory class object (default: False)</td>
</tr>
<tr>
<td>DrawProgressBar</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Draw progress bar to display training, testing and evaluation schedule (default: True)</td>
</tr>
<tr>
<td>AnalysisType</td>
<td>–</td>
<td>Auto</td>
<td>Classification, Regression, Multiclass, Auto</td>
<td>Set the analysis type (Classification, Regression, Multiclass, Auto) (default: Auto)</td>
</tr>
</tbody>
</table>

// Get the signal and background trees from TFile source(s);
// multiple trees can be registered with the Factory
TTree* sigTree = (TTree*)sigSrc->Get( "<YourSignalTreeName>" );
TTree* bkgTreeA = (TTree*)bkgSrc->Get( "<YourBackgrTreeName_A>" );
TTree* bkgTreeB = (TTree*)bkgSrc->Get( "<YourBackgrTreeName_B>" );
TTree* bkgTreeC = (TTree*)bkgSrc->Get( "<YourBackgrTreeName_C>" );

// Set the event weights per tree (these weights are applied in addition to individual event weights that can be specified)
Double_t sigWeight = 1.0;
Double_t bkgWeightA = 1.0, bkgWeightB = 0.5, bkgWeightC = 2.0;

// Register the trees
factory->AddSignalTree( sigTree, sigWeight );
factory->AddBackgroundTree( bkgTreeA, bkgWeightA );
factory->AddBackgroundTree( bkgTreeB, bkgWeightB );
factory->AddBackgroundTree( bkgTreeC, bkgWeightC );

**Code Example 6**: Registration of signal and background ROOT trees read from TFile sources. Overall signal and background weights per tree can also be specified. The TTree object may be replaced by a TChain. The trees will be later split by the factory into subsamples used for testing and training.
Specifying **classification training and test data** in ROOT tree format with signal and background events being located in the same tree:

```c++
TTree* inputTree = (TTree*)source->Get("<YourTreeName>" );
TCut signalCut = ...; // how to identify signal events
TCut backgrCut = ...; // how to identify background events
factory->SetInputTrees( inputTree, signalCut, backgrCut );
```

**Code Example 7**: Registration of a single ROOT tree containing the input data for signal and background, read from a TFile source. The TTree object may be replaced by a TChain. The cuts identify the event species.

Specifying **classification data** in ROOT tree format with signal and background training/test events being located in separate trees:

```c++
#include "TMVA/Types.h"
// Get the signal and background training and test trees from TFile source(s);
TTree* sigTreeTrain = (TTree*)sigSrc->Get("<YourSignalTrainTreeName>" );
TTree* bkgTreeTrain = (TTree*)bkgSrc->Get("<YourBackgrTrainTreeName>" );
TTree* sigTreeTest = (TTree*)sigSrc->Get("<YourSignalTestTreeName>" );
TTree* bkgTreeTest = (TTree*)bkgSrc->Get("<YourBackgrTestTreeName>" );

// Set the event weights (these weights are applied in
// addition to individual event weights that can be specified)
Double_t sigWeight = 1.0;
Double_t bkgWeight = 1.0;

// Register the trees
factory->AddSignalTree( sigTreeTrain, sigWeight, TMVA::Types::kTraining);
factory->AddBackgroundTree( bkgTreeTrain, bkgWeight, TMVA::Types::kTraining);
factory->AddSignalTree( sigTreeTest, sigWeight, TMVA::Types::kTesting);
factory->AddBackgroundTree( bkgTreeTest, bkgWeight, TMVA::Types::kTesting);
```

**Code Example 8**: Registration of signal and background ROOT trees read from TFile sources. The first two tree are specified to be used only for training the other two only for testing. Please note that the tree type testing/training requires the inclusion of the header file TMVA/Types.h.

Specifying **classification training and test data** in text format:
Code Example 9: Registration of signal and background text files used for training and testing. Names and types of the input variables are given in the first line, followed by the values.
Specifying **regression training and test data** in ROOT tree format:

```c
factory->AddRegressionTree( regTree, weight );
```

**Code Example 10**: Registration of a ROOT tree containing the input and target variables. An overall weight per tree can also be specified. The `TTree` object may be replaced by a `TChain`.

Rather than having only global weighting factors for individual input trees which allow to scale them to the same “luminosity”, individual event weights can be applied as well. These weights should be available event-by-event, i.e. as a column or a function of columns of the input data sets. To specify the weights to be used for the training use the command:

```c
factory->SetWeightExpression( "<YourWeightExpression>" );
```
or if you have different expressions (variables) used as weights in the signal and background trees:

```c
factory->SetSignalWeightExpression( "<YourSignalWeightExpression>" );
factory->SetBackgroundWeightExpression( "<YourBackgroundWeightExpression>" );
```

**Code Example 11**: Specification of individual weights for the training events. The expression must be a function of variables present in the input data set.

### 3.1.2 Negative event weights

In next-to-leading order Monte Carlo generators, events with (unphysical) negative weights may occur in some phase space regions. Such events are often troublesome to deal with, and it depends on the concrete implementation of the MVA method, whether or not they are treated properly. Among those methods that correctly incorporate events with negative weights are likelihood and multi-dimensional probability density estimators, but also decision trees. A summary of this feature for all TMVA methods is given in Table 7. In cases where a method does not properly treat events with negative weights, it is advisable to ignore such events for the training - but to include them in the performance evaluation to not bias the results. This can be explicitly requested for each MVA method via the boolean configuration option `IgnoreNegWeightsInTraining` (cf. Option Table 9 on page 71).

### 3.1.3 Defining input variables, spectators and targets

The variables in the input trees used to train the MVA methods are registered with the Factory using the `AddVariable` method. It takes the variable name (string), which must have a correspondence in the input ROOT tree or input text file, and optionally a number type (‘F’ (default) and ‘I’). The type is used to inform the method whether a variable takes continuous floating point or discrete
values.\footnote{For example for the projective likelihood method, a histogram out of discrete values would not (and should not) be interpolated between bins.} Note that ’F’ indicates any floating point type, i.e., \emph{float} and \emph{double}. Correspondingly, ’I’ stands for integer, \emph{including int, short, char}, and the corresponding \emph{unsigned} types. Hence, if a variable in the input tree is \emph{double}, it should be declared ’F’ in the \texttt{AddVariable} call.

It is possible to specify variable expressions, just as for the \texttt{TTree::Draw} command (the expression is interpreted as a \texttt{TTreeFormula}, including the use of arrays). Expressions may be abbreviated for more concise screen output (and plotting) purposes by defining shorthand-notation \emph{labels} via the assignment operator :=.

In addition, two more arguments may be inserted into the \texttt{AddVariable} call, allowing the user to specify \emph{titles} and \emph{units} for the input variables for displaying purposes.

The following code example revises all possible options to declare an input variable:

\begin{verbatim}
factory->AddVariable( "<YourDescreteVar>", 'I' );
factory->AddVariable( "log(<YourFloatingVar>)", 'F' );
factory->AddVariable( "SumLabel := <YourVar1>+<YourVar2>", 'F' );
factory->AddVariable( "<YourVar3>", "Pretty Title", "Unit", 'F' );
\end{verbatim}

\textbf{Code Example 12:} Declaration of variables used to train the MVA methods. Each variable is specified by its name in the training tree (or text file), and optionally a type (’F’ for floating point and ’I’ for integer, ’F’ is default if nothing is given). Note that ’F’ indicates any floating point type, i.e., \emph{float} and \emph{double}. Correspondingly, ’I’ stands for integer, \emph{including int, short, char}, and the corresponding \emph{unsigned} types. Hence, even if a variable in the input tree is \emph{double}, it should be declared ’F’ here. Here, \emph{YourVar1} has discrete values and is thus declared as an integer. Just as in the \texttt{TTree::Draw} command, it is also possible to specify expressions of variables. The := operator defines labels (third row), used for shorthand notation in screen outputs and plots. It is also possible to define titles and units for the variables (fourth row), which are used for plotting. If labels \emph{and} titles are defined, labels are used for abbreviated screen outputs, and titles for plotting.

It is possible to define \emph{spectator variables}, which are part of the input data set, but which are not used in the MVA training, test nor during the evaluation. They are copied into the \texttt{TestTree}, together with the used input variables and the MVA response values for each event, where the spectator variables can be used for correlation tests or others. Spectator variables are declared as follows:

\begin{verbatim}
factory->AddSpectator( "<YourSpectatorVariable>" );
factory->AddSpectator( "log(<YourSpectatorVariable>)" );
factory->AddSpectator( "<YourSpectatorVariable>", "Pretty Title", "Unit" );
\end{verbatim}

\textbf{Code Example 13:} Various ways to declare a spectator variable, not participating in the MVA analysis, but written into the final \texttt{TestTree}.
For a regression problem, the target variable is defined similarly, without however specifying a number type:

```cpp
factory->AddTarget( "<YourRegressionTarget1>" );
factory->AddTarget( "log(<YourRegressionTarget2>)" );
factory->AddTarget( "<YourRegressionTarget3>", "Pretty Title", "Unit" );
```

**Code Example 14**: Various ways to declare the target variables used to train a multivariate regression method. If the MVA method supports multi-target (multidimensional) regression, more than one regression target can be defined.

### 3.1.4 Preparing the training and test data

The input events that are handed to the Factory are internally copied and split into one training and one test ROOT tree. This guarantees a statistically independent evaluation of the MVA algorithms based on the test sample. The numbers of events used in both samples are specified by the user. They must not exceed the entries of the input data sets. In case the user has provided a ROOT tree, the event copy can (and should) be accelerated by disabling all branches not used by the input variables.

It is possible to apply selection requirements (cuts) upon the input events. These requirements can depend on any variable present in the input data sets, i.e., they are not restricted to the variables used by the methods. The full command is as follows:

```cpp
TCut preselectionCut = "<YourSelectionString>";
factory->PrepareTrainingAndTestTree( preselectionCut, "<options>" );
```

**Code Example 15**: Preparation of the internal TMVA training and test trees. The sizes (number of events) of these trees are specified in the configuration option string. For classification problems, they can be set individually for signal and background. Note that the preselection cuts are applied before the training and test samples are created, i.e., the tree sizes apply to numbers of selected events. It is also possible to choose among different methods to select the events entering the training and test trees from the source trees. All options are described in Option-Table 2. See also the text for further information.

For classification, the numbers of signal and background events used for training and testing are specified in the configuration string by the variables `nTrain_Signal`, `nTrain_Background`, `nTest_Signal` and `nTest_Background` (for example, "nTrain_Signal=5000:nTrain_Background=5000:nTest_Signal=4000:nTest_Background=5000"). The default value (zero) signifies that all available events are taken, e.g., if `nTrain_Signal=5000` and `nTest_Signal=0`, and if the total signal sample has 15000 events, then 5000 signal events are used for training and the remaining 10000 events are

---

5 A fully unbiased training and evaluation requires at least three statistically independent data sets. See comments in Footnote 9 on page 29.
used for testing. If \(n_{\text{Train\_Signal}}=0\) and \(n_{\text{Test\_Signal}}=0\), the signal sample is split in half for training and testing. The same rules apply to background. Since zero is default, not specifying anything corresponds to splitting the samples in two halves.

For regression, only the sizes of the train and test samples are given, e.g., "\(n_{\text{Train\_Regression}}=0: n_{\text{Test\_Regression}}=0\)", so that one half of the input sample is used for training and the other half for testing. If a tree is given to the factory as a training tree. The events of that tree can only be used for training. The same is true for test trees.

The option `SplitMode` defines how the training and test samples are selected from the source trees. With `SplitMode=Random`, events are selected randomly. With `SplitMode=Alternate`, events are chosen in alternating turns for the training and test samples as they occur in the source trees until the desired numbers of training and test events are selected. The training and test samples should contain the same number of events for each event class. In the `SplitMode=Block` mode the first \(n_{\text{Train\_Signal}}\) and \(n_{\text{Train\_Background}}\) (classification), or \(n_{\text{Train\_Regression}}\) events (regression) of the input data set are selected for the training sample, and the next \(n_{\text{Test\_Signal}}\) and \(n_{\text{Test\_Background}}\) or \(n_{\text{Test\_Regression}}\) events comprise the test data. This is usually not desired for data that contains varying conditions over the range of the data set. For the Random selection mode, the seed of the random generator can be set. With `SplitSeed=0` the generator returns a different random number series every time. The default seed of 100 ensures that the same training and test samples are used each time TMVA is run (as does any other seed apart from 0).

The option `MixMode` defines the order of how the training and test events of the different classes are combined into a training sample. It also defines the order in which they appear in the test sample. The available options for `MixMode` are the same as for `SplitMode`. By default, the same option is chosen for the `MixMode` as given in `SplitMode`. Again, with `MixMode=Random`, the order of the events in the samples is random. With `MixMode=Alternate` subsequent events are always of the next class (e.g. 0, 1, 2, 3, 0, 1, 2, 3, \ldots). With `MixMode=Block` all events of one class are inserted in a block into the training/test samples (e.g. 0, 0, \ldots, 0, 1, 1, \ldots, 1, 2, 2, \ldots, 2, \ldots).

In some cases event weights are given by Monte Carlo generators, and may turn out to be overall very small or large numbers. To avoid artifacts due to this, TMVA can internally renormalise the signal and background training(!) weights such that their respective sums of effective (weighted) events is equal. This is the default renormalisation and it can be modified with the configuration option `NormMode` (cf. Table 2). Possible settings are: None: no renormalisation is applied (the weights are used as given), NumEvents: renormalisation of the training events such that the sum of event weights of the Signal and Background events, respectively are equal to the number of events \(Ns,Nb\) requested in the call `Factory::PrepareTrainingAndTestTree("", "n_{\text{Train\_Signal}}=Ns, n_{\text{Train\_Background}}=Nb...", EqualNumEvents` (default): the event weights are renormalised such that both, the sum of all weighted signal training events equals the sum of all weights of the background training events. Note: All this renormalisation only affects the training events as the training of some classifiers is sensitive to the relative amount of signal and background in the training data. On the other hand, the background or signal efficiency of the trained classifier as determined from the test sample is independent of the relative abundance of signal and background events.
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SplitMode</td>
<td>–</td>
<td>Random</td>
<td>Random, Alternate, Block</td>
<td>Method of picking training and testing events (default: random)</td>
</tr>
<tr>
<td>MixMode</td>
<td>–</td>
<td>SameAsSplitMode</td>
<td>SameAsSplitMode, Random, Alternate, Block</td>
<td>Method of mixing events of different classes into one dataset (default: SameAsSplitMode)</td>
</tr>
<tr>
<td>SplitSeed</td>
<td>–</td>
<td>100</td>
<td>–</td>
<td>Seed for random event shuffling</td>
</tr>
<tr>
<td>NormMode</td>
<td>–</td>
<td>EqualNumEvents</td>
<td>None, NumEvents, EqualNumEvents</td>
<td>Overall renormalisation of event-by-event weights used in the training (NumEvents: average weight of 1 per event, independently for signal and background; EqualNumEvents: average weight of 1 per event for signal, and sum of weights for background equal to sum of weights for signal)</td>
</tr>
<tr>
<td>nTrain_Signal</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>Number of training events of class Signal (default: 0 = all)</td>
</tr>
<tr>
<td>nTest_Signal</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>Number of test events of class Signal (default: 0 = all)</td>
</tr>
<tr>
<td>nTrain_Background</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>Number of training events of class Background (default: 0 = all)</td>
</tr>
<tr>
<td>nTest_Background</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>Number of test events of class Background (default: 0 = all)</td>
</tr>
<tr>
<td>V</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Verbosity (default: true)</td>
</tr>
<tr>
<td>VerboseLevel</td>
<td>–</td>
<td>Info</td>
<td>Debug, Verbose, Info</td>
<td>VerboseLevel (Debug/Verbose/Info)</td>
</tr>
<tr>
<td>Correlations</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Whether to show variable correlations output (default: true)</td>
</tr>
</tbody>
</table>

**Option Table 2:** Configuration options reference in call `Factory::PrepareTrainingAndTestTree(..)`. For regression, `nTrain_Signal` and `nTest_Signal` are replaced by `nTrain_Regression` and `nTest_Regression`, respectively, and `nTrain_Background` and `nTest_Background` are removed. See also Code-Example 15 and comments in the text.
3.1 The TMVA Factory

3.1.5 Booking MVA methods

All MVA methods are booked via the Factory by specifying the method’s type, plus a unique name chosen by the user, and a set of specific configuration options encoded in a string qualifier. If the same method type is booked several times with different options (which is useful to compare different sets of configurations for optimisation purposes), the specified names must be different to distinguish the instances and their weight files. A booking example for the likelihood method is given in Code Example 16 below. Detailed descriptions of the configuration options are given in the corresponding tools and MVA sections of this Users Guide, and booking examples for most of the methods are given in Appendix A. With the MVA booking the initialisation of the Factory is complete and no MVA-specific actions are left to do. The Factory takes care of the subsequent training, testing and evaluation of the MVA methods.

```
factory->BookMethod( TMVA::Types::kLikelihood, "LikelihoodD", 
    "!H:!V:!TransformOutput:PDFInterpol=Spline2:\n    NSmoothSig[0]=20:NSmoothBkg[0]=20:NSmooth=5:\n    NAvevtPerBin=50:VarTransform=Decorrelate" );
```

Code Example 16: Example booking of the likelihood method. The first argument is a unique type enumerator (the available types can be looked up in src/Types.h), the second is a user-defined name which must be unique among all booked MVA methods, and the third is a configuration option string that is specific to the method. For options that are not explicitly set in the string default values are used, which are printed to standard output. The syntax of the options should be explicit from the above example. Individual options are separated by a ‘:’. Boolean variables can be set either explicitly as MyBoolVar=True/False, or just via MyBoolVar/!MyBoolVar. All specific options are explained in the tools and MVA sections of this Users Guide. There is no difference in the booking of methods for classification or regression applications. See Appendix A on page 163 for a complete booking list of all MVA methods in TMVA.

3.1.6 Help option for MVA booking

Upon request via the configuration option "H" (see code example above) the TMVA methods print concise help messages. These include a brief description of the algorithm, a performance assessment, and hints for setting the most important configuration options. The messages can also be evoked by the command factory->PrintHelpMessage("<MethodName>").

3.1.7 Training the MVA methods

The training of the booked methods is invoked by the command:

---

6In the TMVA package all MVA methods are derived from the abstract interface IMethod and the base class MethodBase.
factory->TrainAllMethods();

Code Example 17: Executing the MVA training via the Factory.

The training results are stored in the weight files which are saved in the directory `weights` (which, if not existing is created).\textsuperscript{7} The weight files are named `Jobname_MethodName.weights.(extension)`, where the job name has been specified at the instantiation of the Factory, and `MethodName` is the unique method name specified in the booking command. Each method writes a custom weight file in XML format (extension is `xml`), where the configuration options, controls and training results for the method are stored.

### 3.1.8 Testing the MVA methods

The trained MVA methods are applied to the test data set and provide scalar outputs according to which an event can be classified as either signal or background, or which estimate the regression target.\textsuperscript{8} The MVA outputs are stored in the test tree (`TestTree`) to which a column is added for each booked method. The tree is eventually written to the output file and can be directly analysed in a ROOT session. The testing of all booked methods is invoked by the command:

factory->TestAllMethods();

Code Example 18: Executing the validation (testing) of the MVA methods via the Factory.

### 3.1.9 Evaluating the MVA methods

The Factory and data set classes of TMVA perform a preliminary property assessment of the input variables used by the MVA methods, such as computing correlation coefficients and ranking the variables according to their separation (for classification), or according to their correlations with the target variable(s) (for regression). The results are printed to standard output.

The performance evaluation in terms of signal efficiency, background rejection, faithful estimation of a regression target, etc., of the trained and tested MVA methods is invoked by the command:

\textsuperscript{7}The default weight file directory name can be modified from the user script through the global configuration variable (`TMVA::gConfig().GetIONames()).fWeightFileDir`.

\textsuperscript{8}In classification mode, TMVA discriminates signal from background in data sets with unknown composition of these two samples. In frequent use cases the background (sometimes also the signal) consists of a variety of different populations with characteristic properties, which could call for classifiers with more than two discrimination classes. However, in practice it is usually possible to serialise background fighting by training individual classifiers for each background source, and applying consecutive requirements to these. Since TMVA 4, the framework directly supports multi-class classification. However, some MVA methods have not yet been prepared for it.
3.1 The TMVA Factory

```cpp
factory->EvaluateAllMethods();
```

**Code Example 19:** Executing the performance evaluation via the Factory.

The performance measures differ between classification and regression problems. They are summarised below.

### 3.1.10 Classification performance evaluation

After training and testing, the linear correlation coefficients among the classifier outputs are printed. In addition, overlap matrices are derived (and printed) for signal and background that determine the fractions of signal and background events that are equally classified by each pair of classifiers. This is useful when two classifiers have similar performance, but a significant fraction of non-overlapping events. In such a case a combination of the classifiers (e.g., in a Committee classifier) could improve the performance (this can be extended to any combination of any number of classifiers).

The optimal method to be used for a specific analysis strongly depends on the problem at hand and no general recommendations can be given. To ease the choice TMVA computes a number of benchmark quantities that assess the performance of the methods on the independent test sample. For classification these are

- **The signal efficiency at three representative background efficiencies** (the efficiency is equal to 1 − rejection) obtained from a cut on the classifier output. Also given is the area of the background rejection versus signal efficiency function (the larger the area the better the performance).

- **The separation** $\langle S^2 \rangle$ of a classifier $y$, defined by the integral [7]

  $$
  \langle S^2 \rangle = \frac{1}{2} \int \frac{(\hat{y}_S(y) - \hat{y}_B(y))^2}{\hat{y}_S(y) + \hat{y}_B(y)} dy,
  $$

  where $\hat{y}_S$ and $\hat{y}_B$ are the signal and background PDFs of $y$, respectively (cf. Sec. 3.1.13). The separation is zero for identical signal and background shapes, and it is one for shapes with no overlap.

- **The discrimination significance** of a classifier, defined by the difference between the classifier means for signal and background divided by the quadratic sum of their root-mean-squares.

The results of the evaluation are printed to standard output. Smooth background rejection/efficiency versus signal efficiency curves are written to the output ROOT file, and can be plotted using custom macros (see Sec. 3.3).
3.1.11 Regression performance evaluation

Ranking for regression is based on the correlation strength between the input variables or MVA method response and the regression target. Several correlation measures are implemented in TMVA to capture and quantify nonlinear dependencies. Their results are printed to standard output.

- The Correlation between two random variables $X$ and $Y$ is usually measured with the correlation coefficient $\rho$, defined by

$$\rho(X, Y) = \frac{\text{cov}(X,Y)}{\sigma_X \sigma_Y}, \quad (2)$$

The correlation coefficient is symmetric in $X$ and $Y$, lies within the interval $[-1, 1]$, and quantifies by definition a linear relationship. Thus $\rho = 0$ holds for independent variables, but the converse is not true in general. In particular, higher order functional or non-functional relationships may not, or only marginally, be reflected in the value of $\rho$ (see Fig. 9).

- The correlation ratio is defined by

$$\eta^2(Y|X) = \frac{\sigma_{E(Y|X)}}{\sigma_Y}, \quad (3)$$

where

$$E(Y|X) = \int y \ P(y|x) \ dy, \quad (4)$$

is the conditional expectation of $Y$ given $X$ with the associated conditional probability density function $P(Y|X)$. The correlation ratio $\eta^2$ is in general not symmetric and its value lies within [0, 1], according to how well the data points can be fitted with a linear or nonlinear regression curve. Thus non-functional correlations cannot be accounted for by the correlation ratio. The following relations can be derived for $\eta^2$ and the squared correlation coefficient $\rho^2$ [9]:

- $\rho^2 = \eta^2 = 1$, if $X$ and $Y$ are in a strict linear functional relationship.
- $\rho^2 \leq \eta^2 = 1$, if $X$ and $Y$ are in a strict nonlinear functional relationship.
- $\rho^2 = \eta^2 < 1$, if there is no strict functional relationship but the regression of $X$ on $Y$ is exactly linear.
- $\rho^2 < \eta^2 < 1$, if there is no strict functional relationship but some nonlinear regression curve is a better fit than the best linear fit.

Some characteristic examples and their corresponding values for $\eta^2$ are shown in Fig. 9. In the special case, where all data points take the same value, $\eta$ is undefined.

- Mutual information allows to detect any predictable relationship between two random variables, be it of functional or non-functional form. It is defined by [10]

$$I(X,Y) = \sum_{X,Y} P(X,Y) \ln \frac{P(X,Y)}{P(X)P(Y)}, \quad (5)$$
Figure 9: Various types of correlations between two random variables and their corresponding values for the correlation coefficient $\rho$, the correlation ratio $\eta$, and mutual information $I$. Linear relationship (upper left), functional relationship (upper right), non-functional relationship (lower left), and independent variables (lower right).

where $P(X,Y)$ is the joint probability density function of the random variables $X$ and $Y$, and $P(X)$, $P(Y)$ are the corresponding marginal probabilities. Mutual information originates from information theory and is closely related to entropy which is a measure of the uncertainty associated with a random variable. It is defined by

$$H(X) = - \sum_X P(X) \ln P(X) ,$$

where $X$ is the discrete random variable and $P(X)$ the associated probability density function.
The connection between the two quantities is given by the following transformation

\[ I(X,Y) = \sum_{X,Y} P(X,Y) \ln \frac{P(X,Y)}{P(X)P(Y)} \] (7)

\[ = \sum_{X,Y} P(X,Y) \ln \frac{P(X|Y)}{P_X(X)} \] (8)

\[ = -\sum_{X,Y} P(X,Y) \ln P(X) + \sum_{X,Y} P(X,Y) \ln P(X|Y) \] (9)

\[ = -\sum_{X,Y} P(X) \ln P(X) - (\sum_{X,Y} P(X,Y) \ln P(X|Y)) \] (10)

\[ = H(X) - H(X|Y) \] (11)

where \( H(X|Y) \) is the conditional entropy of \( X \) given \( Y \). Thus mutual information is the reduction of the uncertainty in variable \( X \) due to the knowledge of \( Y \). Mutual information is symmetric and takes positive absolute values. In the case of two completely independent variables \( I(X,Y) \) is zero.

For experimental measurements the joint and marginal probability density functions are a priori unknown and must be approximated by choosing suitable binning procedures such as kernel estimation techniques (see, e.g., [11]). Consequently, the values of \( I(X,Y) \) for a given data set will strongly depend on the statistical power of the sample and the chosen binning parameters.

For the purpose of ranking variables from data sets of equal statistical power and identical binning, however, we assume that the evaluation from a simple two-dimensional histogram without further smoothing is sufficient.

A comparison of the correlation coefficient \( \rho \), the correlation ratio \( \eta \), and mutual information \( I \) for linearly correlated two-dimensional Gaussian toy MC simulations is shown in Table 1.

### Table 1: Comparison of the correlation coefficient \( \rho \), correlation ratio \( \eta \), and mutual information \( I \) for two-dimensional Gaussian toy Monte-Carlo distributions with linear correlations as indicated (20000 data points/100 × 100 bins).

<table>
<thead>
<tr>
<th>( \rho_{PDF} )</th>
<th>0.0</th>
<th>0.1</th>
<th>0.2</th>
<th>0.3</th>
<th>0.4</th>
<th>0.5</th>
<th>0.6</th>
<th>0.7</th>
<th>0.8</th>
<th>0.9</th>
<th>0.9999</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \rho )</td>
<td>0.006</td>
<td>0.092</td>
<td>0.191</td>
<td>0.291</td>
<td>0.391</td>
<td>0.492</td>
<td>0.592</td>
<td>0.694</td>
<td>0.795</td>
<td>0.898</td>
<td>1.0</td>
</tr>
<tr>
<td>( \eta^2 )</td>
<td>0.004</td>
<td>0.012</td>
<td>0.041</td>
<td>0.089</td>
<td>0.156</td>
<td>0.245</td>
<td>0.354</td>
<td>0.484</td>
<td>0.634</td>
<td>0.806</td>
<td>1.0</td>
</tr>
<tr>
<td>( I )</td>
<td>0.093</td>
<td>0.099</td>
<td>0.112</td>
<td>0.139</td>
<td>0.171</td>
<td>0.222</td>
<td>0.295</td>
<td>0.398</td>
<td>0.560</td>
<td>0.861</td>
<td>3.071</td>
</tr>
</tbody>
</table>

3.1.12 Overtraining

Overtraining occurs when a machine learning problem has too few degrees of freedom, because too many model parameters of an algorithm were adjusted to too few data points. The sensitivity to
overtraining therefore depends on the MVA method. For example, a Fisher (or linear) discriminant can hardly ever be overtrained, whereas, without the appropriate counter measures, boosted decision trees usually suffer from at least partial overtraining, owing to their large number of nodes. Overtraining leads to a seeming increase in the classification or regression performance over the objectively achievable one, if measured on the training sample, and to an effective performance decrease when measured with an independent test sample. A convenient way to detect overtraining and to measure its impact is therefore to compare the performance results between training and test samples. Such a test is performed by TMVA with the results printed to standard output.

Various method-specific solutions to counteract overtraining exist. For example, binned likelihood reference distributions are smoothed before interpolating their shapes, or unbinned kernel density estimators smear each training event before computing the PDF; neural networks steadily monitor the convergence of the error estimator between training and test samples\(^9\) suspending the training when the test sample has passed its minimum; the number of nodes in boosted decision trees can be reduced by removing insignificant ones (“tree pruning”), etc.

\subsection{3.1.13 Other representations of MVA outputs for classification: probabilities and probability integral transformation (\textit{Rarity})}

In addition to the MVA response value \(y\) of a classifier, which is typically used to place a cut for the classification of an event as either signal or background, or which could be used in a subsequent likelihood fit, TMVA also provides the classifier’s signal and background PDFs, \(\hat{y}_{S(B)}\). The PDFs can be used to derive classification probabilities for individual events, or to compute any kind of transformation of which the \textit{Probability integral transformation} (\textit{Rarity}) transformation is implemented in TMVA.

\begin{itemize}
\item \textbf{Classification probability}: The techniques used to estimate the shapes of the PDFs are those developed for the likelihood classifier (see Sec. 8.2.2 for details) and can be customised individually for each method (the control options are given in Sec. 8). The probability for event \(i\) to be of signal type is given by,
\end{itemize}

\[ P_S(i) = \frac{\hat{f}_S \cdot \hat{y}_S(i)}{\hat{f}_S \cdot \hat{y}_S(i) + (1 - \hat{f}_S) \cdot \hat{y}_B(i)}, \tag{12} \]

where \(\hat{f}_S = N_S / (N_S + N_B)\) is the expected signal fraction, and \(N_{S(B)}\) is the expected number of signal (background) events (default is \(\hat{f}_S = 0.5\)).\(^{10}\)

\footnote{\(\hat{f}_S\) distributions may exhibit a somewhat peculiar structure with frequent narrow peaks. They are generated by regions of classifier output values in which \(\hat{y}_S \propto \hat{y}_B\) for which \(P_S\) becomes a constant.}

\footnote{Proper training and validation requires three statistically independent data sets: one for the parameter optimisation, another one for the overtraining detection, and the last one for the performance validation. In TMVA, the last two samples have been merged to increase statistics. The (usually insignificant) bias introduced by this on the evaluation results does not affect the analysis as far as classification cut efficiencies or the regression resolution are independently validated with data.}
Figure 10: Example plots for classifier probability integral transformation distributions for signal and background events from the academic test sample. Shown are likelihood (left) and Fisher (right).

- **Probability Integral Transformation**: The Probability integral transformation $\mathcal{R}(y)$ of a classifier $y$ is given by the integral

$$\mathcal{R}(y) = \int_{-\infty}^{y} \hat{y}_B(y') \ dy' ,$$

which is defined such that $\mathcal{R}(y_B)$ for background events is uniformly distributed between 0 and 1, while signal events cluster towards 1. The signal distributions can thus be directly compared among the various classifiers. The stronger the peak towards 1, the better is the discrimination.

Another useful aspect of the probability integral transformation is the possibility to directly visualise deviations of a test background (which could be physics data) from the training sample, by exhibition of non-uniformity.

The probability integral transformation distributions of the Likelihood and Fisher classifiers for the example used in Sec. 2 are plotted in Fig. 10. Since Fisher performs better (cf. Fig. 6 on page 12), its signal distribution is stronger peaked towards 1. By construction, the background distributions are uniform within statistical fluctuations.

The probability and probability integral transformation distributions can be plotted with dedicated macros, invoked through corresponding GUI buttons.

### 3.2 Cross Validation

The goal of cross validation, and the larger framework of validation, is to estimate the performance of a machine learning model.

A model $\hat{f}(x|\theta)$ takes as input a data point $x$ and outputs a prediction for that data point given a set of tunable parameters $\theta$. The parameters are tuned through a training process using a training
set, \( \mathcal{T} \), for which the true output is known. This results in a model whose parameters are tuned to perform as good as possible on new, unseen, data.

The training error, \( \text{err}_T \), for a model is defined as

\[
\text{err}_T = \frac{1}{N_t} \sum_{n=1}^{N_t} L(y_n, \hat{f}(x_n)) \tag{14}
\]

where \( N_t \) is the number of events used for training, \( L \) is a chosen loss function, \( \hat{f} \) is our model, and \( x_n \) and \( y_n \) are points in our training set.

The training error, in general, is a poor estimator of the performance of the model on new, unseen data. It is generally a decreasing function of the number of training iterations and unless the method is simple, i.e. has few tunable parameters, it can start to adapt to the noise in the training data. When this happens the training error continues to go down but the general performance, the error on data outside of the training set, starts increasing. This effect is called overfitting.

The test error, or prediction error, is defined as the expected error when the model is applied to new, unseen data.

\[
\text{Err}_T = E \left[ L(Y, \hat{f}(X)) | \mathcal{T} \right] \tag{15}
\]

using the same notation as Eq 14 and where \((X,Y)\) are two random variables drawn from the joint probability distribution. Here the model, \( \hat{f} \), is trained using the training set, \( \mathcal{T} \), and the error is evaluated over all possible inputs in the input space.

A related measure, the expected prediction error, additionally averages over all possible training sets

\[
\text{Err} = E \left[ L(Y, \hat{f}(X)) \right] = E \left[ \text{Err}_T \right] \tag{16}
\]

This notation is inspired by [55]. To understand, and to more easily remember, what each quantity signifies one can consider whether it considers concrete data or random variables. The training error is defined for events in the training set; we can actually compute the value and thus uses a minuscule initial letter and an overbar. The prediction error is defined over the complete input space and uses random variables in the definition thus has a capital letter. The subscript signifies what data was used to train the model.

The simplest way to reliably estimate \( \text{Err}_T \) is to partition the initial data set into two parts and use one part for training and one part for testing. In the case where access to data is unlimited this method yields an optimal estimate.

However, often access to data is limited; For example in physics where the cost of generating Monte Carlo samples can be prohibitive, or in medical surveys where the number of respondents is limited. This means a choice has to be made, how much data should be used for training, and how much for evaluating the performance?

As a larger fraction of events are used for training, the performance of the final model increases due to better tuned parameters. However, our estimation of that performance becomes increasingly uncertain due to the limited size of the test set.
One way to reap the benefits of a large training set and large test set is to use cross validation. This discussion will focus on one technique in particular: K-folds.

In k-folds cross validation, initially introduced in [56], a data set is split into equal sized partitions, or folds. A model is then trained using one fold as test set and the concatenation of the remaining folds as training set. This procedure is repeated until each fold has been used as test set exactly once. In this way data efficiency is gained at the cost of an increased computational burden.

The expected prediction error of a model trained with the procedure can then be estimated as the average of the error of each individual fold.

$$\text{Err} = \frac{1}{K} \sum \text{Err}_k.$$ \hspace{1cm} (17)

Having many folds of adequate size gives the best estimation. Increasing the number of folds will yield more models to average over, increasing the confidence of how consistent the model achieves a given level of performance. However, it reduces the statistical strength of each fold. In practise, 10 folds may give a good trade-off between effects and can serve as a good initial guess.

**On validation and model selection** Ideally, the test set would only be used a single time to evaluate the performance of a model. If the data set is reused, an amount of bias is introduced. A common practise in the initial phase of an analysis is to try several ideas out to get a grip on what works and what works not. To select the best model from the set of proposed ideas is called model selection.

To solve this problem, the data can be partitioned into an additional set, often called the validation set. This set can be reused, introducing bias in the performance estimation as estimated by that set. A final test set is then used on the unique model selected in the selection process. Model selection encompasses both selecting between model types such as BDT, k-nearest neighbours, SVM etcetera; and choosing hyperparameters for a model.

One way to realise how this process can introduce bias is to consider it a training session where the model to use is a hyperparameter. The discussion about overfitting below then applies. This is primarily a concern when the models under consideration are prone to overfitting, i.e. when then number of tunable parameters of the model is much larger than the number of events in the training data set. This because a model not prone to overfitting will have very consistent generalisation performance. With a model that can overfit, you can get lucky and find a configuration of parameters that give a good performance on the limited data set.

TMVA facilitates either performance evaluation of a single model, or model selection from a set, but not both at the same time. Setting up an analysis which encompasses both is possible and requires manual set up of surrounding code.
Figure 11: Model 1, 2, and 3 are used only for performance estimation and are then thrown away. A final model (model 4) is trained using the complete dataset.

3.2.1 Using CV in practise

Using k-folds for estimating the performance of a training procedure for a model is an efficient use of data, but it does not result in a final model which to apply to data.

Two schemes for generating a final model, \( \hat{f}(x, |\theta) \), where the parameters \( \theta \) are fixed after training together with an estimate of its performance are presented here.

Retraining A simple approach to evaluating the performance of a model using k-folds cross validation is to approximate the error as described in Section 3.2.

We then retrain the model using all available training data and assumes the performance estimation from the previous step holds for this final model.

The estimate of the model performance when trained on \( N_t \) events will be similar to the performance of a model trained on \( N \) events.

The reasoning is that cross validation estimates the expected error averaged over all training sets of size \( N_t \), \( \text{Err}_{N_t} \). Then, if \( N \approx N_t \), we make the approximation \( \text{Err}_{N_t} \approx \text{Err}_N \).

That is, put in a different way: On average, the final model will have the given average performance. Note that there are two averages here since our estimation was not done for the trained model but for a similar one.

Cross validation in application A problem with the previous approach is that the estimation is not done for the final model, but rather of the average final model. This complicates statistical
3.2.2 Implementation

Cross validation in TMVA is implemented as a wrapper around the Factory class. This means the cross validated work flow uses a separate entry point, but effort is made to maintain similarity between the two interfaces. Do note however that there is a difference in how the dataloader is handled. For the Factory a separate dataloader can be associated with each booked method, in cross validation the same dataloader is applied to all booked methods. See Example 21 for an example of a simple set up.

Cross validation is supported for all analysis types that the Factory supports, i.e. Classification, Regression and Multiclass.

After training and evaluation, output files suitable for analysis with the different TMVA GUIs can be generated. All analysis types are supported with some caveats.

The current cross validation implementation supports k-folds splitting. This splitting mechanism is implemented separately from the dataset splitting mechanism and currently only uses events the dataloader puts in the test set. See Example 20 for an example of how this is done for a simple Classification set up.

The k-folds splitting mechanism supports two modes of operations. Random splitting and functional splitting. The random splitting is activated when the splitExpr option is left out or given as the empty string. In this case events are assigned to folds randomly (where the random generator is...
seeded with the seed provided in the option SplitSeed). Random splitting is only supported for training and evaluation, not in the application phase. This makes it suitable for quickly getting up and running, trying different methods out, but not for final analysis and application to data.

For data application, the second mode of operation, functional splitting, is preferred. Using the option splitExpr you can define an expression that is evaluated for each event. The result of evaluation is used for fold assignment. See Section 3.2.4 for more details.

```cpp
auto d = new TMVA::DataLoader("<datasetName>");
d->AddVariable("x");
d->AddSpectator("eventID");
d->AddSignalTree(sigTree);
d->AddBackgroundTree(bkgTree);
d->PrepareTrainingAndTestTree("", "nTest_Signal=0"
                           "nTest_Background=0");
```

**Code Example 20:** Setting up a typical dataloader for using with cross validation. It is assumed that you have available to TTrees, one with signal events, one with background events, both with suitable variables defined. Cross validation in TMVA uses a separate splitting mechanism which is applied after the ordinary splitting step. It uses only events in the training set, the test set is currently left unused. To make all events available to cross validation, they are placed in the training set by setting the size of the test set to zero.

```cpp
auto d = new TMVA::DataLoader("<datasetName>");
auto f = TFile::Open("path/to/file");
TString opt = "!V:!Silent:!Correlations"
             "AnalysisType=Classification"
             "NumFolds=2";
TMVA::CrossValidation cv {"<jobname>", d, f, opt};

cv.BookMethod(TMVA::Types::kBDT, "BDT", "<options>");
cv.Evaluate();
```

**Code Example 21:** Minimal example to get cross validation up and running using the dataloader defined in 20. The example sets up a cross validated BDT with 2 folds where the folds are assigned randomly. The results will be available for inspection with the TMVA GUI interface under "path/to/file".

### 3.2.3 Cross validation options

Constructing a `CrossValidation` object is very similar to how one would construct a TMVA Factory with the exception of how the data loader is handled. In TMVA cross validation, you are responsible for constructing the data loader. When passed to the `CrossValidation` constructor it takes ownership and makes sure that the memory is properly released after training and evaluation.
An schematic of how to construct a new `CrossValidation` object is detailed in Code Example 22.

```c
CrossValidation cv {"<jobname>" , dataloader , "<options>"};
```

**Code Example 22:** Constructing a CrossValidation instance: the first argument is a job name, which will get prepended to all files produced by the CrossValidation factory; The second is the data loader that will provide data for all methods booked through the CrossValidation class; The third is a list of options configuring this instance. Available options can be found in Table ?? . Individual options are separated by a ':'. See Sec. 3.1.5 for more information on the booking.

`CrossValidation` wraps a TMVA Factory internally and thus supports all options that the Factory supports. In addition it, can be configured with the following options.

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SplitExpr</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>Expression used to assign events to folds. If not given or given as &quot;&quot; events will be assigned to folds randomly.</td>
</tr>
<tr>
<td>SplitSeed</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>Only used when SplitExpr is &quot;&quot;. Determines the seed for the random assignment of events to folds.</td>
</tr>
<tr>
<td>NumFolds</td>
<td>–</td>
<td>2</td>
<td>–</td>
<td>Number of folds to generate.</td>
</tr>
<tr>
<td>FoldFileOutput</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>If given a TMVA output file will be generated for each fold. File name will be the same as specified for the combined output with a _foldX suffix.</td>
</tr>
<tr>
<td>OutputEnsembling</td>
<td>–</td>
<td>&quot;None&quot;</td>
<td>&quot;None&quot; , &quot;Avg&quot;</td>
<td>Combines output from contained methods (only in application phase). If None, SplitExpr is used to assign event to fold, no combination is performed. Valid values are &quot;None&quot;, and &quot;Avg&quot;.</td>
</tr>
</tbody>
</table>

### 3.2.4 K-folds splitting

TMVA currently supports k-folds cross validation. In this scheme, events are assigned to one of K folds. The training is then run K times, each time using one fold as the test data and the rest as training data. TMVA supports to modes of assigning events to folds: Random assignment, and assignment through an expression. The option `SplitExpr` selects what mode to use and what expression to evaluate.

The splitting mechanism used by the k-folds implementation is different from the one that the TMVA dataloader uses to partition the input into a training and test set. With k-folds, only events
in the training set after \texttt{PrepareTrainingAndTestTree} has been called are picked up by the k-folds splitting. To use all available events for cross validation one can explicitly set the number of test events to zero, as is shown in Code Example 23.

\begin{verbatim}
dataloader.PrepareTrainingAndTestTree("<sigcut>", "<bkgcut>", "<options>:nTest_Signal=0:nTest_Background=0");
\end{verbatim}

\textbf{Code Example 23:}

\textbf{Random splitting} \quad Random splitting is the default mode of operation if \texttt{SplitExpr} is not specified to the \texttt{CrossValidation} constructor, or if it is given as \texttt{"SplitExpr=\"}. Random splitting is usually used to estimate the performance of a training scheme. After the estimation a final model is created by retraining the method on all available training data. This final model is then applied to new, unlabelled data. More details can be found in 3.2.1.

Be careful if using the resulting models in the TMVA application phase and ensure that you use a separate dataset that has not been used for training. Since the training error can be significantly smaller than the generalisation error, there estimate thus derived will be optimistically biased. This applies equally to a model trained on the complete data set as well as the models resulting from the individual folds.

\textbf{Splitting with an expression} \quad If an expression is provided to the \texttt{CrossValidation} constructor through the option \texttt{SplitExpr}, events will be assigned to folds according to this expression. This is true both for training and application.

The expression can use spectators defined in the dataset and a special variable \texttt{numFolds} that will be replaced with the corresponding concrete number at split time.

The idea behind using an expression for splitting is to generate a final model for which the performance estimates acquired through cross validation applies. As such the result of evaluating the expression should in essence be a uniformly distributed random number, independent of the content of an event. It is therefore preferred to assign a globally unique number to event at creation time and use this as input.

An example can be seen in Code Example 24. The syntax is a consequence of the expression being backed by a TFormula. Spectators and context variables are made available through TFormula named parameters, which are accessed with brackets "[]". TFormulas only handle floating point operands which necessitates the integer casts. See the documentation of TFormula for more information on syntax.
auto * d = new TMVA::DataLoader("<name>");
d.AddVariable("x");
d.AddVariable("y");
d.AddSpectator("eventNumber");

TString splitExpr = "int([eventNumber])\%int([numFolds])";
TString options = TString::Format("SplitExpr=%s", splitExpr);

TMVA::CrossValidation cv {"<jobname>", d, options};

Code Example 24: The split expression uses spectators defined on the TMVA::DataSet to calculate the fold assignment for each fold. Note that code for adding data, methods, and for training has been abbreviated.

3.2.5 Output

Cross validation in TMVA provides several different outputs to facilitate analysis. Firstly, a file suitable for analysis with the GUIs presented in Section 2.6 is produced after a successful training. One can also request the generation of additional files for per-fold analysis with the GUIs.

Finally, a selection of statistics are collected during training and made available through a programmatic interface.

**Offline analysis through the TMVA GUI**  Output files suitable for analysis with the TMVA GUI is produced after a completed training if a TFile is provided the CrossValidation constructor. For each fold, the independent test set is evaluated and added to the output file.

Optionally files with per-fold training and test set output files can be generated.

All analysis types are supported, i.e. classification, multiclass, and regression.

The provided output file will be contain all events provided the test set, since k-folds splitting can ensure that each training event belongs only to one test set.

Do note that the training set produced when using the TMVA Factory is difficult to define in a meaningful way here since each event will be used as training, multiple times, and test event. However, for implementation considerations the training set will still be present in the output file.

In the current version it is a copy of the test set, however the contents of the training set cannot be depended on as it may change in future releases.

To generate output files for offline analysis of the separate folds one submits the option foldOutputFiles=True to the CrossValidation constructor. The output files will use the same name as the provided TFile, with an extra suffix, _foldK, where K indicates the corresponding fold.
There are no caveats when analysing the per-fold output files, as compared to above.

```
root -l -e 'TMVA::TMVAGui("<path/to/file>")'
```

**Code Example 25:** Command line command to run to inspect the resulting output file using the classification GUI. For the multiclass and regression GUIs, use `TMVA::TMVAMultiClassGui` and `TMVA::TMVARegressionGui` respectively. Note that "<path/to/file>" can be either the output file provided to the `CrossValidation` constructor, or one of the fold outputs. The per-fold output files have the same name, with an extra suffix, _foldK, where K indicates fold id.

**Summary statistics after training** Some statistics are collected during training and evaluation on a per-fold basis. These are made available through the `TMVA::CrossValidation::GetResults` method.

Tracked metrics include, ROC curves and integral, signal significance, separation and background efficiencies.

An example of how this is used in practise is seen in Code Example 26 and an example output of running that code can be seen in Code Example 27.

For full documentation of available statistics, see root.cern.ch/doc/master/classTMVA_1_1CrossValidationResult.html.

```
TMVA::CrossValidation cv {"<jobname>", <dataloader>, <outputFile>, "<options>");
cv.BookMethod(<methodType>, "<methodName>", "<options>");
cv.Evaluate();

size_t iMethod = 0;
for (auto && result : cv.GetResults()) {
    std::cout << "Summary for method "
              << cv.GetMethods()[iMethod++].GetValue<TString>("MethodName")
              << std::endl;
    for (UInt_t iFold = 0; iFold<cv.GetNumFolds(); ++iFold) {
        std::cout << "\tFold " << iFold << ": "
                  << "ROC int: " << result.GetROCValues()[iFold]
                  << ", "
                  << "BkgEff@SigEff=0.3: " << result.GetEff30Values()[iFold]
                  << std::endl;
    }
}
```

**Code Example 26:** Statistics is available through the `CrossValidationResults` object. For example, ROC integrals and efficiencies can be accessed per fold. See the ROOT reference for up to date interface documentation.
Summary for method BDT
Fold 0: ROC int: 0.961109, BkgEff@SigEff=0.3: 0.961
Fold 1: ROC int: 0.974712, BkgEff@SigEff=0.3: 0.986

Summary for method Fisher
Fold 0: ROC int: 0.964606, BkgEff@SigEff=0.3: 0.963
Fold 1: ROC int: 0.977215, BkgEff@SigEff=0.3: 0.992


### 3.2.6 Application

Application is the phase where the model is presented with new, unlabelled data. This requires a final model which is ready to accept events. The naive approach of cross validation does not produce a final model to be evaluated, but instead produces one model for each fold. Generating a such a model can be done in a multitude of ways including the approaches presented in Section ??.

When using the first approach, retraining your model on the complete data set, you would do this using the TMVA Factory as detailed in Section 3.1 and the process of applying data using the TMVA Reader as detailed in section 3.4.

When using the second approach, cross validation in application, TMVA simplifies the process for you. You need to specify a split expression so that TMVA can partition the input data. After training, a combined model is produced, containing the models for the folds and the split expression.

This model can be used with the regular TMVA Reader and ensures that applied events are evaluated by the corresponding model, as indicated by the split expression. You as the end-user need only ensure that the same variables and spectators are defined for the application data set as in training, and that the weight files for each fold model reside in the same directory as the combined model.

```cpp
TMVA::Reader reader {"!Color:!Silent:!V");

reader.AddVariable("x", &x);
reader.AddVariable("y", &y);
reader.AddSpectator("eventID", &eventID);

reader.BookMVA("<methodName>", "<path/to/weight/file.xml>");
```

Code Example 28: Using a model training with cross validation in application uses the same interface as the non cross validated case. One must make sure, however, that the variable and spectator declarations are the same as in training, and the per-fold model weight files reside in the same folder as the combined model weight file specified by "<path/to/weight/file.xml>".
3.3 ROOT macros to plot training, testing and evaluation results

TMVA provides simple GUIs (TMVAGui.C and TMVARegGui.C, see Fig. 1), which interface ROOT macros that visualise the various steps of the training analysis. The macros are respectively located in TMVA/macros/ (Sourceforge.net distribution) and $ROOTSYS/tmva/test/ (ROOT distribution), and can also be executed from the command line. They are described in Tables 2 and 4. All plots drawn are saved as png files (or optionally as eps, gif files) in the macro subdirectory plots which, if not existing, is created.

The binning and histogram boundaries for some of the histograms created during the training, testing and evaluation phases are controlled via the global singleton class TMVA::Config. They can be modified as follows:

```c++
// Modify settings for the variable plotting
(TMVA::gConfig().GetVariablePlotting()).fTimesRMS = 8.0;
(TMVA::gConfig().GetVariablePlotting()).fNbins1D = 60.0;
(TMVA::gConfig().GetVariablePlotting()).fNbins2D = 300.0;

// Modify the binning in the ROC curve (for classification only)
(TMVA::gConfig().GetVariablePlotting()).fNbinsXOfROCCurve = 100;

// For file name settings, modify the struct TMVA::Config::IONames
(TMVA::gConfig().GetIONames()).fWeightFileDir = "myWeightFileDir";
```

**Code Example 29:** Modifying global parameter settings for the plotting of the discriminating input variables. The values given are the TMVA defaults. Consult the class files Config.h and Config.cxx for all available global configuration variables and their default settings, respectively. Note that the additional parentheses are mandatory when used in CINT.

3.4 The TMVA Reader

After training and evaluation, the most performing MVA methods are chosen and used to classify events in data samples with unknown signal and background composition, or to predict values of a regression target. An example of how this *application phase* is carried out is given in TMVA/macros/TMVAClassificationApplication.C and TMVA/macros/TMVAREgressionApplication.C (Sourceforge.net), or $ROOTSYS/tmva/test/TMVAClassificationApplication.C and $ROOTSYS/tmva/test/TMVAREgressionApplication.C (ROOT).

Analogously to the Factory, the communication between the user application and the MVA methods is interfaced by the TMVA *Reader*, which is created by the user:
<table>
<thead>
<tr>
<th>Macro</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>variables.C</td>
<td>Plots the signal and background MVA input variables (training sample). The second argument sets the directory, which determines the preprocessing type (InputVariables_Id for default identity transformation, cf. Sec. 4.1). The third argument is a title, and the fourth argument is a flag whether or not the input variables served a regression analysis.</td>
</tr>
<tr>
<td>correlationsscatter.C</td>
<td>Plots superimposed scatters and profiles for all pairs of input variables used during the training phase (separate plots for signal and background in case of classification). The arguments are as above.</td>
</tr>
<tr>
<td>correlations.C</td>
<td>Plots the linear correlation matrices for the input variables in the training sample (distinguishing signal and background for classification).</td>
</tr>
<tr>
<td>mvas.C</td>
<td>Plots the classifier response distributions of the test sample for signal and background. The second argument (HistType=0,1,2,3) allows to also plot the probability (1) and probability integral transformation (2) distributions of the classifiers, as well as a comparison of the output distributions between test and training samples. Plotting of probability and probability integral transformation requires the CreateMVAPdfs option for the classifier to be set to true.</td>
</tr>
<tr>
<td>mvaeffs.C</td>
<td>Signal and background efficiencies, obtained from cutting on the classifier outputs, versus the cut value. Also shown are the signal purity and the signal efficiency times signal purity corresponding to the expected number of signal and background events before cutting (numbers given by user). The optimal cuts according to the best significance are printed on standard output.</td>
</tr>
<tr>
<td>efficiencies.C</td>
<td>Background rejection (second argument type=2, default), or background efficiency (type=1), versus signal efficiency for the classifiers (test sample). The efficiencies are obtained by cutting on the classifier outputs. This is traditionally the best plot to assess the overall discrimination performance (ROC curve).</td>
</tr>
<tr>
<td>paracoor.C</td>
<td>Draws diagrams of “Parallel coordinates” \cite{33} for signal and background, used to visualise the correlations among the input variables, but also between the MVA output and input variables (indicating the importance of the variables).</td>
</tr>
</tbody>
</table>

Table 2: ROOT macros for the representation of the TMVA input variables and classification results. All macros take as first argument the name of the ROOT file containing the histograms (default is TMVA.root). They are conveniently called via the TMVAGui.C GUI (the first three macros are also called from the regression GUI TMVARegGui.C). Macros for the representation of regression results are given in Table 3. Plotting macros for MVA method specific information are listed in Table 4.
### Macro Description

<table>
<thead>
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</tr>
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<tbody>
<tr>
<td>deviations.C</td>
<td>Plots the linear deviation between regression target value and MVA response or input variables for test and training samples.</td>
</tr>
<tr>
<td>regression_average_devs.C</td>
<td>Draws the average deviation between the MVA output and the regression target value for all trained methods.</td>
</tr>
</tbody>
</table>

**Table 3:** ROOT macros for the representation of the TMVA regression results. All macros take as first argument the name of the ROOT file containing the histograms (default is `TMVA.root`). They are conveniently called from the `TMVAREgGui.C` GUI.

```c++
TMVA::Reader* reader = new TMVA::Reader( "<options>" );
```

**Code Example 30:** Instantiating a Reader class object. The only options are the booleans: `V` for verbose, `Color` for coloured output, and `Silent` to suppress all output.

### 3.4.1 Specifying input variables

The user registers the names of the input variables with the Reader. They are required to be the same (and in the same order) as the names used for training (this requirement is not actually mandatory, but enforced to ensure the consistency between training and application). Together with the name is given the address of a local variable, which carries the updated input values during the event loop.

```c++
Int_t localDescreteVar;
Float_t localFloatingVar, locaSum, localVar3;
reader->AddVariable( "<YourDescreteVar>", &localDescreteVar );
reader->AddVariable( "log(<YourFloatingVar>)", &localFloatingVar );
reader->AddVariable( "SumLabel := <YourVar1>+<YourVar2>", &locaSum );
reader->AddVariable( "<YourVar3>", &localVar3 );
```

**Code Example 31:** Declaration of the variables and references used as input to the methods (cf. Code Example 12). The order and naming of the variables must be consistent with the ones used for the training. The local variables are updated during the event loop, and through the references their values are known to the MVA methods. The variable type must be either `float` or `int` (`double` is not supported).
Macro Description

**likelihoodrefs.C**  Plots the reference PDFs of all input variables for the projective likelihood method and compares it to original distributions obtained from the training sample.

**network.C**  Draws the TMVA-MLP architecture including weights after training (does not work for the other ANNs).

**annconvergencetest.C**  Plots the MLP error-function convergence versus the training epoch for training and test events (does not work for the other ANNs).

**BDT.C(i)**  Draws the i-th decision tree of the trained forest (default is i=1). The second argument is the weight file that contains the full architecture of the forest (default is `weights/TMVAClassification_BDT.weights.xml`).

**BDTControlPlots.C**  Plots distributions of boost weights throughout forest, boost weights versus decision tree, error fraction, number of nodes before and after pruning and the coefficient $\alpha$.

**mvarefs.C**  Plots the PDFs used to compute the probability response for a classifier, and compares it to the original distributions.

**PlotFoams.C**  Draws the signal and background foams created by the method PDE-Foam.

**rulevis.C**  Plots the relative importance of rules and linear terms. The 1D plots show the accumulated importance per input variable. The 2D scatter plots show the same but correlated between the input variables. These plots help to identify regions in the parameter space that are important for the model.

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</tr>
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**Table 4:** List of ROOT macros representing results for specific MVA methods. The macros require that these methods have been included in the training. All macros take as first argument the name of the ROOT file containing the histograms (default is TMVA.root).

### 3.4.2 Booking MVA methods

The selected MVA methods are booked with the Reader using the weight files from the preceding training job:

```cpp
reader->BookMVA( "<YourMethodName>", "<path/JobName_MethodName.weights.xml>" );
```

**Code Example 32:** Booking a multivariate method. The first argument is a user defined name to distinguish between methods (it does not need to be the same name as for training, although this could be a useful choice). The true type of the method and its full configuration are read from the weight file specified in the second argument. The default structure of the weight file names is: `path/(JobName)_(MethodName).weights.xml`. 
3.4.3 Requesting the MVA response

Within the event loop, the response value of a classifier, and – if available – its error, for a given set of input variables computed by the user, are obtained with the commands:

```c++
localDiscreteVar = treeDiscreteVar; // reference could be implicit
localFloatingVar = log(treeFloatingVar);
localSum = treeVar1 + treeVar2;
localVar3 = treeVar3; // reference could be implicit

// Classifier response
Double_t mvaValue = reader->EvaluateMVA( "<YourMethodName>" );

// Error on classifier response – must be called after "EvaluateMVA"
// (not available for all methods, returns -1 in that case)
Double_t mvaErr = reader->GetMVAError();
```

Code Example 33: Updating the local variables for an event, and obtaining the corresponding classifier output and error (if available – see text).

The output of a classifier may then be used for example to put a cut that increases the signal purity of the sample (the achievable purities can be read off the evaluation results obtained during the test phase), or it could enter a subsequent maximum-likelihood fit, or similar. The error reflects the uncertainty, which may be statistical, in the output value as obtained from the training information.

For regression, multi-target response is already supported in TMVA, so that the retrieval command reads:

```c++
// Regression response for one target
Double_t regValue = (reader->EvaluateRegression( "<YourMethodName>" ))[0];
```

Code Example 34: Obtaining the regression output (after updating the local variables for an event – see above). For multi-target regression, the corresponding vector entries are filled.

The output of a regression method could be directly used for example as energy estimate for a calorimeter cluster as a function of the cell energies.

The rectangular cut classifier is special since it returns a binary answer for a given set of input variables and cuts. The user must specify the desired signal efficiency to define the working point according to which the Reader will choose the cuts:
Boo_t passed = reader->EvaluateMVA( "Cuts", signalEfficiency );

**Code Example 35:** For the cut classifier, the second parameter gives the desired signal efficiency according to which the cuts are chosen. The return value is 1 for passed and 0 for retained. See Footnote 20 on page 71 for information on how to determine the optimal working point for known signal and background abundance.

Instead of the classifier response values, one may also retrieve the ratio (12) from the Reader, which, if properly normalised to the expected signal fraction in the sample, corresponds to a probability. The corresponding command reads:

```
Double_t pSig = reader->GetProba( "<YourClassifierName>", sigFrac );
```

**Code Example 36:** Requesting the event’s signal probability from a classifier. The signal fraction is the parameter $f_S$ in Eq. (12).

Similarly, the probability integral transformation ($Rarity$) (13) of a classifier is retrieved by the command

```
Double_t rarity = reader->GetRarity( "<YourClassifierName>");
```

**Code Example 37:** Requesting the event’s probability integral transformation distribution from a classifier.

### 3.5 An alternative to the Reader: standalone C++ response classes

To simplify the portability of the trained MVA response to any application the TMVA methods generate after the training lightweight standalone C++ response classes including in the source code the content of the weight files.\(^{11}\) These classes do not depend on ROOT, neither on any other non-standard library. The names of the classes are constructed out of `Read+"MethodName"`, and they inherit from the interface class `IClassifierReader`, which is written into the same C++ file. These standalone classes are presently only available for classification.

An example application (ROOT script here, not representative for a C++ standalone application) for a Fisher classifier is given in Code-Example 38. The example is also available in the macro `TMVA/macros/ClassApplication.C` (Sourceforge.net). These classes are C++ representations of the information stored in the weight files. Any change in the training parameters will generate a

\(^{11}\) At present, the class making functionality has been implemented for all MVA methods with the exception of cut optimisation, PDE-RS, PDE-Foam and k-NN. While for the former classifier the cuts can be easily implemented into the user application, and do not require an extra class, the implementation of a response class for PDE-RS or k-NN requires a copy of the entire analysis code, which we have not attempted so far. We also point out that the use of the standalone C++ class for BDT is not practical due to the colossal size of the generated code.
// load the generated response class into macro and compile it (ROOT)
// or include it into any C++ executable
gROOT->LoadMacro( "TMVAClassification_Fisher.class.C++" ); // usage in ROOT

// define the names of the input variables (same as for training)
std::vector<std::string> inputVars;
inputVars.push_back( "<YourVar1>" );
inputVars.push_back( "log(<YourVar2>)" );
inputVars.push_back( "<YourVar3>+<YourVar4>" );

// create a class object for the Fisher response
IClassifierReader* fisherResponse = new ReadFisher( inputVars );

// the user’s event loop ...
std::vector<double> inputVec( 3 );
for (...) {
    // compute the input variables for the event
    inputVec[0] = treeVar1;
    inputVec[1] = TMath::Log(treeVar2);
    inputVec[2] = treeVar3 + treeVar4;

    // get the Fisher response
    double fiOut = fisherResponse->GetMvaValue( inputVec );
    // ... use fiOut
}

Code Example 38: Using a standalone C++ class for the classifier response in an application (here of the Fisher discriminant). See also the example code in TMVA/macros/ClassApplication.C (Sourceforge.net).

new class, which must be updated in the corresponding application.\textsuperscript{12}

For a given test event, the MVA response returned by the standalone class is identical to the one returned by the Reader. Nevertheless, we emphasise that the recommended approach to apply the training results is via the Reader.

\textsuperscript{12}We are aware that requiring recompilation constitutes a significant shortcoming. we consider to upgrade these classes to reading the XML weight files, which entails significant complications if the independence of any external library shall be conserved.
4 Data Preprocessing

It is possible to preprocess the discriminating input variables or the training events prior to presenting them to a multivariate method. Preprocessing can be useful to reduce correlations among the variables, to transform their shapes into more appropriate forms, or to accelerate the response time of a method (event sorting).

The preprocessing is completely transparent to the MVA methods. Any preprocessing performed for the training is automatically performed in the application through the Reader class. All the required information is stored in the weight files of the MVA method. The preprocessing methods normalisation and principal component decomposition are available for input and target variables, gaussianization, uniformization and decorrelation discussed below can only be used for input variables.

Apart from five variable transformation methods mentioned above, an unsupervised variable selection method Variance Threshold is also implemented in TMVA. It follows a completely different processing pipeline. It is discussed in detail in section 4.2.

4.1 Transforming input variables

Currently five preprocessing transformations are implemented in TMVA:

- variable normalisation;
- decorrelation via the square-root of the covariance matrix;
- decorrelation via a principal component decomposition;
- transformation of the variables into Uniform distributions (“Uniformization”).
- transformation of the variables into Gaussian distributions (“Gaussianisation”).

Normalisation and principal component decomposition can be applied to input and target variables, the other transformations can only be used for the input variables. The transformations be performed for all input variables of for a selected subset of the input variables only. The latter is especially useful in situations where only some variables are correlated or the correlations are very different for different subsets.

All transformations can be performed separately for signal and background events because their correlation patterns are usually different.\footnote{Different transformations for signal and background events are only useful for methods that explicitly distinguish signal and background hypotheses. This is the case for the likelihood and PDE-RS classifiers. For all other methods the user must choose which transformation to use.}

Technically, any transformation of the input and target variables is performed “on the fly” when the event is requested from the central DataSet class. The preprocessing is hence fully transparent to
4.1 Transforming input variables

![Diagram of variable transformation interface in TMVA]

Figure 13: Schematic view of the variable transformation interface implemented in TMVA. Each concrete MVA method derives from `MethodBase` (interfaced by `IMethod`), which holds a protected member object of type `TransformationHandler`. In this object a list of objects derived from `VariableTransformBase` which are the implementations of the particular variable transformations available in TMVA is stored. The construction of the concrete variable transformation objects proceeds in `MethodBase` according to the transformation methods requested in the option string. The events used by the MVA methods for training, testing and final classification (or regression) analysis are read via an API of the `TransformationHandler` class, which itself reads the events from the `DataSet` and applies subsequently all initialised transformations. The `DataSet` fills the current values into the reserved event addresses (the event content may either stem from the training or testing trees, or is set by the user application via the `Reader` for the final classification/regression analysis). The `TransformationHandler` class ensures the proper transformation of all events seen by the MVA methods.

the MVA methods. Any preprocessing performed for the training is automatically also performed in the application through the Reader class. All the required information is stored in the weight files of the MVA method. Each MVA method carries a variable transformation type together with a pointer to the object of its transformation class which is owned by the `DataSet`. If no preprocessing is requested, an identity transform is applied. The `DataSet` registers the requested transformations and takes care not to recreate an identical transformation object (if requested) during the training phase. Hence if two MVA methods wish to apply the same transformation, a single object is shared between them. Each method writes its transformation into its weight file once the training has converged. For testing and application of an MVA method, the transformation is read from the weight file and a corresponding transformation object is created. Here each method owns its transformation so that no sharing of potentially different transformation objects occurs (they may have been obtained with different training data and/or under different conditions). A schematic view of the variable transformation interface used in TMVA is drawn in Fig. 13.

4.1.1 Variable normalisation

Minimum and maximum values for the variables to be transformed are determined from the training events and used to linearly scale these input variables to lie within $[-1, 1]$. Such a transformation is useful to allow direct comparisons between the MVA weights assigned to the variables, where large
absolute weights may indicate strong separation power. Normalisation may also render minimisation processes, such as the adjustment of neural network weights, more effective.

### 4.1.2 Variable decorrelation

A drawback of, for example, the projective likelihood classifier (see Sec. 8.2) is that it ignores correlations among the discriminating input variables. Because in most realistic use cases this is not an accurate conjecture it leads to performance loss. Also other classifiers, such as rectangular cuts or decision trees, and even multidimensional likelihood approaches underperform in presence of variable correlations.

Linear correlations, measured in the training sample, can be taken into account in a straightforward manner through computing the square-root of the covariance matrix. The square-root of a matrix $C$ is the matrix $C'$ that multiplied with itself yields $C$: $C = (C')^2$. TMVA computes the square-root matrix by means of diagonalising the (symmetric) covariance matrix

$$D = S^T C S \quad \Rightarrow \quad C' = S \sqrt{D} S^T,$$

where $D$ is a diagonal matrix, and where the matrix $S$ is symmetric. The linear decorrelation of the selected variables is then obtained by multiplying the initial variable tuple $x$ by the inverse of the square-root matrix

$$x \mapsto (C')^{-1} x .$$

The decorrelation is complete only for linearly correlated and Gaussian distributed variables. In situations where these requirements are not fulfilled only little additional information can be recovered by the decorrelation procedure. For highly nonlinear problems the performance may even become worse with linear decorrelation. Nonlinear methods without prior variable decorrelation should be used in such cases.

### 4.1.3 Principal component decomposition

Principal component decomposition or principal component analysis (PCA) as presently applied in TMVA is not very different from the above linear decorrelation. In common words, PCA is a linear transformation that rotates a sample of data points such that the maximum variability is visible. It thus identifies the most important gradients. In the PCA-transformed coordinate system, the largest variance by any projection of the data comes to lie on the first coordinate (denoted the first principal component), the second largest variance on the second coordinate, and so on. PCA can thus be used to reduce the dimensionality of a problem (initially given by the number of input variables) by removing dimensions with insignificant variance. This corresponds to keeping lower-order principal components and ignoring higher-order ones. This latter step however goes beyond straight variable transformation as performed in the preprocessing steps discussed here (it rather represents itself a full classification method). Hence all principal components are retained here.

The tuples $x_U^{PC}(i) = (x_{U,1}^{PC}(i), \ldots, x_{U,n_{var}}^{PC}(i))$ of principal components of a tuple of input variables $x(i) = (x_1(i), \ldots, x_{n_{var}}(i))$, measured for the event $i$ for signal ($U = S$) and background ($U = B$),
are obtained by the transformation

$$x_{PC}^{U,k}(i) = \sum_{\ell=1}^{n_{\text{var}}} (x_{U,\ell}(i) - \bar{x}_{U,\ell}) v_{U,\ell}^{(k)}, \quad \forall k = 1, n_{\text{var}}. \quad (20)$$

The tuples $\bar{x}_{U}$ and $v_{U}^{(k)}$ are the sample means and eigenvectors, respectively. They are computed by the ROOT class TPrincipal. The matrix of eigenvectors $V_{U} = (v_{U}^{(1)}, \ldots, v_{U}^{(n_{\text{var}})})$ obeys the relation

$$C_{U} \cdot V_{U} = D_{U} \cdot V_{U}, \quad (21)$$

where $C$ is the covariance matrix of the sample $U$, and $D_{U}$ is the tuple of eigenvalues. As for the preprocessing described in Sec. 4.1.2, the transformation (20) eliminates linear correlations for Gaussian variables.

### 4.1.4 Uniform and Gaussian transformation of variables (“Uniformisation” and “Gaussianisation”)

The decorrelation methods described above require linearly correlated and Gaussian distributed input variables. In real-life HEP applications this is however rarely the case. One may hence transform the variables prior to their decorrelation such that their distributions become Gaussian. The corresponding transformation function is conveniently separated into two steps: first, transform a variable into a uniform distribution using its cumulative distribution function obtained from the training data (this transformation is identical to the probability integral transformation (“Rarity”) introduced in Sec. 3.1.13 on page 29); second, use the inverse error function to transform the uniform distribution into a Gaussian shape with zero mean and unity width. As for the other transformations, one needs to choose which class of events (signal or background) is to be transformed and hence, for the input variables (Gaussianisation is not available for target values), it is only possible to transform signal or background into proper Gaussian distributions (except for classifiers testing explicitly both hypotheses such as likelihood methods). Hence a discriminant input variable $x$ with the probability density function $\hat{x}$ is transformed as follows

$$x \mapsto \sqrt{2} \cdot \text{erf}^{-1} \left( 2 \cdot \int_{-\infty}^{x} \hat{x}(x') \, dx' - 1 \right). \quad (22)$$

A subsequent decorrelation of the transformed variable tuple sees Gaussian distributions, but most likely non-linear correlations as a consequence of the transformation (22). The distributions obtained after the decorrelation may thus not be Gaussian anymore. It has been suggested that iterating Gaussianisation and decorrelation more than once may improve the performance of likelihood methods (see next section).

---

14 The cumulative distribution function $F(x)$ of the variable $x$ is given by the integral $F(x) = \int_{-\infty}^{x} \hat{x}(x') \, dx'$, where $\hat{x}$ is the probability density function of $x$. 
4.1.5 Booking and chaining transformations for some or all input variables

Variable transformations to be applied prior to the MVA training (and application) can be defined independently for each MVA method with the booking option \texttt{VarTransform=<type>}, where \texttt{<type>} denotes the desired transformation (or chain of transformations). The available transformation types are normalisation, decorrelation, principal component analysis and Gaussianisation, which are labelled by \texttt{Norm}, \texttt{Deco}, \texttt{PCA}, \texttt{Uniform}, \texttt{Gauss}, respectively, or, equivalently, by the short-hand notations \texttt{N}, \texttt{D}, \texttt{P}, \texttt{U}, \texttt{G}.

Transformations can be \emph{chained} allowing the consecutive application of all defined transformations to the variables for each event. For example, the above Gaussianisation and decorrelation sequence would be programmed by \texttt{VarTransform=G,D}, or even \texttt{VarTransform=G,D,G,D} in case of two iterations (instead of the comma ",", a "+" can be equivalently used as chain operator, i.e. \texttt{VarTransform=G+D+G+D}). The ordering of the transformations goes from left (first) to right (last).

\begin{verbatim}
factory->BookMethod( TMVA::Types::kLD, "LD_GD", "H:!V:VarTransform=G,D");
\end{verbatim}

\textbf{Code Example 39:} Booking of a linear discriminant (LD) classifier with Gaussianisation and decorrelation for all input variables.

By default, the transformations are computed with the use of all training events. It is possible to specify the use of a specific class only (e.g., \texttt{Signal}, \texttt{Background}, \texttt{Regression}) by attaching \texttt{<class name>} to the user option – where \texttt{<class name>} has to be replaced by the actual class name (e.g., \texttt{Signal}) – which defines the transformation (e.g., \texttt{VarTransform=G,Signal}). A complex transformation option might hence look like \texttt{VarTransform=D,G,Signal,N}. The attachment \texttt{AllClasses} is equivalent to the default, where events from all classes are used.

\begin{verbatim}
factory->BookMethod( TMVA::Types::kLD, "LD_GDSignal", "H:!V:VarTransform= G_Signal,D_Signal");
\end{verbatim}

\textbf{Code Example 40:} Booking of a linear discriminant (LD) classifier where the Gaussianisation and decorrelation transformations are computed from the signal events only.

By default, all input variables are transformed when booking a transformation. In many situations however, it is desirable to apply a transformation only to a subset of all variables\textsuperscript{15}. Such a selection can be achieved by specifying the variable names separated by commas in brackets after the transformation name, i.e. \texttt{VarTransform=N(var2, var1, myvar3)}. Alternatively the variables and targets can be denoted by their position \texttt{V(index)} (for input variables) and \texttt{T(index)} (for targets) where the index is the position of the variable (target) according to the order in which the variables where defined to the factory (e.g. \texttt{VarTransform=N(V0, V3, T3, ).} The numbering starts with

\textsuperscript{15}As decorrelation for example is only reasonable if the variables have strong “linear” correlations, it might be wise to apply the decorrelation only to those variables that have linear correlations.
0 and the ordering of the variables within the parentheses has no impact. To address all variables (targets) the keywords _V_ and _T_ can be used. The result of a transformation is written back into the same variables. For instance in the case of the principal component analysis transformation the first (and most important) eigenvalue is written into the variable with the smallest index, the second eigenvalue into the variable with the second lowest index and so forth. Input variables and targets can only be transformed at the same time if each variable is transformed independently. This is the case for the normalisation transformation.

Transformations of variable subsets can also be chained. For example, in `VarTransform=N+P(_V0_,var2, _V4_)+P(_T_) Background+G_SIGNAL`, first all variables and targets are normalized with respect to all events, then the variables with the indices 0 and 4 and the variable `var2` are PCA transformed, subsequently all targets are PCA transformed with respect to the background events and finally all variables are gaussianised with respect to the signal events.

```c
factory->BookMethod( TMVA::Types::kLD, "LD_DSubset", "H:!V:VarTransform= \nD(_V0_,var2)" );
```

**Code Example 41:** Booking of a linear discriminant (LD) classifier where only the first variable and the variable labeled `var2` are subject to a decorrelation transformation.

### 4.2 Variable selection based on variance

In high energy physics and machine learning problems, we often encounter datasets which have large number of input variables. However to extract maximum information from the data, we need to select the relevant input variables for the multivariate classification and regression methods implemented in TMVA. Variance Threshold is a simple unsupervised variable selection method which automates this process. It computes weighted variance $\sigma_j^2$ for each variable $j=1, \ldots, n_{\text{var}}$ and ignores the ones whose variance lie below a specific threshold. Weighted variance for each variable is defined as follows:

$$
\sigma_j^2 = \frac{\sum_{i=1}^{N} w_i (x_j(i) - \mu_j)^2}{\sum_{i=1}^{N} w_i} \tag{23}
$$

where $N$ is the total number of events in a dataset, $x(i) = (x_1(i), \ldots, x_{n_{\text{var}}}(i))$ denotes each event $i$ and $w_i$ is the weight of each event. Weighted mean $\mu_j$ is defined as:

$$
\mu_j = \frac{\sum_{i=1}^{N} w_i x_j(i)}{\sum_{i=1}^{N} w_i} \tag{24}
$$

Unlike above five variable transformation method, this Variance Threshold method is implemented in DataLoader class. After loading dataset in the DataLoader object, we can apply this method. It returns a new DataLoader with the selected variables which have variance strictly greater than the threshold value passed by user. Default value of threshold is zero i.e. remove the variables which have same value in all the events.
Code Example 42: VT stands for Variance Threshold. Parameter passed to VarTransform method is just a single string. String strictly follows either of the above two formats for Variance Threshold otherwise method would raise an error.

4.3 Binary search trees

When frequent iterations over the training sample need to be performed, it is helpful to sort the sample before using it. Event sorting in binary trees is employed by the MVA methods rectangular cut optimisation, PDE-RS and k-NN. While the former two classifiers rely on the simplest possible binary tree implementation, k-NN uses on a better performing kd-tree (cf. Ref. [12]).

Efficiently searching for and counting events that lie inside a multidimensional volume spanned by the discriminating input variables is accomplished with the use of a binary tree search algorithm [13]. It is realised in the class BinarySearchTree, which inherits from BinaryTree, and which is also employed to grow decision trees (cf. Sec. 8.13). The amount of computing time needed to sort \(N\) events into the tree is \(\sum_{i=1}^{N} \ln_2(i) = \ln_2(N!) \approx N \ln_2 N\). Finding the events within the tree which lie in a given volume is done by comparing the bounds of the volume with the coordinates of the events in the tree. Searching the tree once requires a CPU time that is \(\propto \ln_2 N\), compared to \(\propto N^{n_{var}}\) without prior event sorting.

5 Probability Density Functions – the PDF Class

Several methods and functionalities in TMVA require the estimation of probability densities (PDE) of one or more correlated variables. One may distinguish three conceptually different approaches to PDEs: (i) parametric approximation, where the training data are fitted with a user-defined parametric function, (ii) nonparametric approximation, where the data are fitted piecewise using simple standard functions, such as a polynomial or a Gaussian, and (iii) nearest-neighbour estimation, where the average of the training data in the vicinity of a test event determines the PDF.

---

\(^{16}\)The following is extracted from Ref. [14] for a two-dimensional range search example. Consider a random sequence of signal events \(e_i(x_1, x_2), i = 1, 2, \ldots\), which are to be stored in a binary tree. The first event in the sequence becomes by definition the topmost node of the tree. The second event \(e_2(x_1, x_2)\) shall have a larger \(x_1\)-coordinate than the first event, therefore a new node is created for it and the node is attached to the first node as the right child (if the \(x_1\)-coordinate had been smaller, the node would have become the left child). Event \(e_3\) shall have a larger \(x_1\)-coordinate than event \(e_1\), it therefore should be attached to the right branch below \(e_1\). Since \(e_2\) is already placed at that position, now the \(x_2\)-coordinates of \(e_2\) and \(e_3\) are compared, and, since \(e_3\) has a larger \(x_2\), \(e_3\) becomes the right child of the node with event \(e_2\). The tree is sequentially filled by taking every event and, while descending the tree, comparing its \(x_1\) and \(x_2\) coordinates with the events already in place. Whether \(x_1\) or \(x_2\) are used for the comparison depends on the level within the tree. On the first level, \(x_1\) is used, on the second level \(x_2\), on the third again \(x_1\) and so on.
All multidimensional PDEs used in TMVA are based on nearest-neighbour estimation with however quite varying implementations. They are described in Secs. 8.3, 8.4 and 8.5.

One-dimensional PDFs in TMVA are estimated by means of nonparametric approximation, because parametric functions cannot be generalised to a-priori unknown problems. The training data can be in form of binned histograms, or unbinned data points (or “quasi-unbinned” data, i.e., histograms with very narrow bins). In the first case, the bin centres are interpolated with polynomial spline curves, while in the latter case one attributes a kernel function to each event such that the PDF is represented by the sum over all kernels. Beside a faithful representation of the training data, it is important that statistical fluctuations are smoothed out as much as possible without destroying significant information. In practise, where the true PDFs are unknown, a compromise determines which information is regarded significant and which is not. Likelihood methods crucially depend on a good-quality PDF representation. Since the PDFs are strongly problem dependent, the default configuration settings in TMVA will almost never be optimal. The user is therefore advised to scrutinise the agreement between training data and PDFs via the available plotting macros, and to optimise the settings.

In TMVA, all PDFs are derived from the PDF class, which is instantiated via the command (usually hidden in the MVA methods for normal TMVA usage):

```cpp
PDF* pdf = new PDF( "<options>", "Suffix", defaultPDF );
pdf->BuildPDF( SourceHistogram );
double p = pdf->GetVal( x );
```

Code Example 43: Creating and using a PDF class object. The first argument is the configuration options string. Individual options are separated by a ‘:’. The second optional argument is the suffix appended to the options used in the option string. The suffix is added to the option names given in the Option Table 3 in order to distinguish variables and types. The third (optional) object is a PDF from which default option settings are read. The histogram specified in the second line is a TH1* object from which the PDF is built. The third line shows how to retrieve the PDF value at a given test value 'x'.

Its configuration options are given in Option Table 3.

5.1 Nonparametric PDF fitting using spline functions

Polynomial splines of various orders are fitted to one-dimensional (1D) binned histograms according to the following procedure.

1. The number of bins of the TH1 object representing the distribution of the input variable is driven by the options NAvEvtPerBin or Nbins (cf. Option Table 3). Setting Nbins enforces a fixed number of bins, while NAvEvtPerBin defines an average number of entries required per bin. The upper and lower bounds of the histogram coincide with the limits found in the data (or they are [−1, 1] if the input variables are normalised).
**Option Table 3:** Configuration options for class: PDF. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. In case a suffix is defined for the PDF, it is added in the end of the option name. For example for PDF with suffix MVAPdf the number of smoothing iterations is given by the option NSmoothMVAPdf (see Option Table 11 on page 75 for a concrete use of the PDF options in a MVA method).

<table>
<thead>
<tr>
<th>Option</th>
<th>Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDFInterpol</td>
<td>KDE,</td>
<td>The method of interpolating the reference histograms:</td>
</tr>
<tr>
<td></td>
<td>Spline0, Spline1, Spline2⋆,</td>
<td>either by using the unbinned kernel density estimator (KDE), or by various</td>
</tr>
<tr>
<td></td>
<td>Spline3, Spline5</td>
<td>degrees of spline functions (note that currently the KDE characteristics</td>
</tr>
<tr>
<td></td>
<td></td>
<td>cannot be changed individually but apply to all variables that select KDE)</td>
</tr>
<tr>
<td>NSmooth</td>
<td>0</td>
<td>Number of smoothing iterations for the input histograms; if set, MinNSmooth</td>
</tr>
<tr>
<td></td>
<td></td>
<td>and MaxNSmooth are ignored</td>
</tr>
<tr>
<td>MinNSmooth</td>
<td>-1</td>
<td>Minimum number of smoothing iteration for the input histograms; for bins</td>
</tr>
<tr>
<td></td>
<td></td>
<td>with least relative error (see text)</td>
</tr>
<tr>
<td>MaxNSmooth</td>
<td>-1</td>
<td>Maximum number of smoothing iteration for the input histograms; for bins</td>
</tr>
<tr>
<td></td>
<td></td>
<td>with most relative error (see text)</td>
</tr>
<tr>
<td>Nbins</td>
<td>0</td>
<td>Number of bins used to build the reference histogram; if set to value &gt; 0,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NAvEvtPerBin is ignored</td>
</tr>
<tr>
<td>NAvEvtPerBin</td>
<td>50</td>
<td>Average number of events per bin in the reference histogram (see text)</td>
</tr>
<tr>
<td>KDEtype</td>
<td>Gauss⋆</td>
<td>KDE kernel type (currently only Gauss)</td>
</tr>
<tr>
<td>KDEIter</td>
<td>Nonadaptive⋆, Adaptive</td>
<td>Non-adaptive or adaptive number of iterations (see text)</td>
</tr>
<tr>
<td>KDEFineFactor</td>
<td>1</td>
<td>Fine-tuning factor for the adaptive KDE</td>
</tr>
<tr>
<td>KDEborder</td>
<td>None⋆, Renorm, Mirror</td>
<td>Method for correcting histogram boundary/border effects</td>
</tr>
<tr>
<td>CheckHist</td>
<td>False</td>
<td>Sanity comparison of derived high-binned interpolated PDF histogram versus</td>
</tr>
<tr>
<td></td>
<td></td>
<td>the original PDF function</td>
</tr>
</tbody>
</table>
2. The histogram is smoothed adaptively between MinNSmooth and MaxNSmooth times, using TH1::Smooth( ) – an implementation of the 353QH-twice algorithm [20]. The appropriate number of smoothing iterations is derived with the aim to preserve statistically significant structures, while smoothing out fluctuations. Bins with the largest (smallest) relative statistical error are maximally (minimally) smoothed. All other bins are smoothed between MaxNSmooth and MinNSmooth times according to a linear function of their relative errors. During the smoothing process a histogram with the suffix NSmooth is created for each variable, where the number of smoothing iterations applied to each bin is stored.

3. The smoothed TH1 object is internally cloned and the requested polynomial splines are used to interpolate adjacent bins. All spline classes are derived from ROOT’s TSpline class. Available are: polynomials of degree 0 (the original smoothed histogram is kept), which is used for discrete variables; degree 1 (linear), 2 (quadratic), 3 (cubic) and degree 5. Splines of degree two or more render the PDF continuous and differentiable in all points excluding the interval borders. In case of a likelihood analysis, this ensures the same property for the likelihood ratio (40). Since cubic (and higher) splines equalise the first and second derivatives at the spline transitions, the resulting curves, although mathematically smooth, can wiggle in quite unexpected ways. Furthermore, there is no local control of the spline: moving one control point (bin) causes the entire curve to change, not just the part near the control point. To ensure a safe interpolation, quadratic splines are used by default.

4. To speed up the numerical access to the probability densities, the spline functions are stored into a finely binned (10^4 bins) histogram, where adjacent bins are interpolated by a linear function. Only after this step, the PDF is normalised according to Eq. (42).

5.2 Nonparametric PDF parameterisation using kernel density estimators

Another type of nonparametric approximation of a 1D PDF are kernel density estimators (KDE). As opposed to splines, KDEs are obtained from unbinned data. The idea of the approach is to estimate the shape of a PDF by the sum over smeared training events. One then finds for a PDF \( p(x) \) of a variable \( x \) [21]

\[
p(x) = \frac{1}{Nh} \sum_{i=1}^{N} K\left( \frac{x-x_i}{h} \right) = \frac{1}{N} \sum_{i=1}^{N} K_h(x-x_i),
\]

where \( N \) is the number of training events, \( K_h(t) = K(t/h)/h \) is the kernel function, and \( h \) is the bandwidth of the kernel (also termed the smoothing parameter). Currently, only a Gaussian form of \( K \) is implemented in TMVA, where the exact form of the kernel function is of minor relevance for the quality of the shape estimation. More important is the choice of the bandwidth.

The KDE smoothing can be applied in either non-adaptive (NA) or adaptive form (A), the choice of which is controlled by the option KDEiter. In the non-adaptive case the bandwidth \( h_{NA} \) is kept constant for the entire training sample. As optimal bandwidth can be taken the one that minimises the asymptotic mean integrated square error (AMISE). For the case of a Gaussian kernel function
this leads to [21]

\[ h_{NA} = \left( \frac{4}{3} \right)^{1/5} \sigma_x N^{-1/5}, \] (26)

where \( \sigma_x \) is the RMS of the variable \( x \).

The so-called sample point adaptive method uses as input the result of the non-adaptive KDE, but also takes into account the local event density. The adaptive bandwidth \( h_A \) then becomes a function of \( p(x) \) [21]

\[ h_A(x) = \frac{h_{NA}}{\sqrt{p(x)}}. \] (27)

The adaptive approach improves the shape estimation in regions with low event density. However, in regions with high event density it can give rise to “over-smoothing” of fine structures such as narrow peaks. The degree of smoothing can be tuned by multiplying the bandwidth \( h_A(x) \) with the user-specified factor \( \text{KDEFineFactor} \).

For practical reasons, the KDE implementation in TMVA differs somewhat from the procedure described above. Instead of unbinned training data a finely-binned histogram is used as input, which allows to significantly speed up the algorithm. The calculation of the optimal bandwidth \( h_{NA} \) is performed in the dedicated class \( \text{KDEKernel} \). If the algorithm is run in the adaptive mode, the non-adaptive step is also performed because its output feeds the computation of \( h_A(x) \) for the adaptive part. Subsequently, a smoothed high-binned histogram estimating the PDF shape is created by looping over the bins of the input histogram and summing up the corresponding kernel functions, using \( h_{NA} \) (\( h_A(x) \)) in case of the non-adaptive (adaptive) mode. This output histogram is returned to the PDF class.

Both the non-adaptive and the adaptive methods can suffer from the so-called boundary problem at the histogram boundaries. It occurs for instance if the original distribution is non-zero below a physical boundary value and zero above. This property cannot be reproduced by the KDE procedure. In general, the stronger the discontinuity the more acute is the boundary problem. TMVA provides three options under the term \( \text{KDEborder} \) that allow to treat boundary problems.

- **KDEborder=None**
  No boundary treatment is performed. The consequence is that close to the boundary the KDE result will be inaccurate: below the boundary it will underestimate the PDF while it will not drop to zero above. In TMVA the PDF resulting from KDE is a (finely-binned) histogram, with bounds equal to the minimum and the maximum values of the input distribution. Hence, the boundary value will be at the edge of the PDF (histogram), and a drop of the PDF due to the proximity of the boundary can be observed (while the artificial enhancement beyond the boundary will fall outside of the histogram). In other words, for training events that are close to the boundary some fraction of the probability “flows” outside the histogram (probability leakage). As a consequence, the integral of the kernel function inside the histogram borders is smaller than one.

- **KDEborder=Renorm**
  The probability leakage is compensated by renormalising the kernel function such that the integral inside the histogram borders is equal to one.
6 Optimisation and Fitting

Several MVA methods (notably cut optimisation and FDA) require general purpose parameter fitting to optimise the value of an estimator. For example, an estimator could be the sum of the deviations of classifier outputs from 1 for signal events and 0 for background events, and the parameters are adjusted so that this sum is as small as possible. Since the various fitting problems call for dedicated solutions, TMVA has a fitter base class, used by the MVA methods, from which all concrete fitters inherit. The consequence of this is that the user can choose whatever fitter is deemed suitable and can configure it through the option string of the MVA method. At present, four fitters are implemented and described below: Monte Carlo sampling, Minuit minimisation, a Genetic Algorithm, Simulated Annealing. They are selected via the configuration option of the corresponding MVA method for which the fitter is invoked (see Option Table 4). Combinations of MC and GA with Minuit are available for the FDA method by setting the Converger option, as described in Option Table 16.

6.1 Monte Carlo sampling

The simplest and most straightforward, albeit inefficient fitting method is to randomly sample the fit parameters and to choose the set of parameters that optimises the estimator. The priors used for the sampling are uniform or Gaussian within the parameter limits. The specific configuration options for the MC sampling are given in Option Table 5.
### Option Table 5: Configuration options reference for fitting method: Monte Carlo sampling (MC).

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SampleSize</td>
<td>–</td>
<td>100000</td>
<td>–</td>
<td>Number of Monte Carlo events in toy sample</td>
</tr>
<tr>
<td>Sigma</td>
<td>–</td>
<td>-1</td>
<td>–</td>
<td>If &gt; 0: new points are generated according to Gauss around best value and with Sigma in units of interval length</td>
</tr>
<tr>
<td>Seed</td>
<td>–</td>
<td>100</td>
<td>–</td>
<td>Seed for the random generator (0 takes random seeds)</td>
</tr>
</tbody>
</table>

For fitting problems with few local minima out of which one is a global minimum the performance can be enhanced by setting the parameter *Sigma* to a positive value. The newly generated parameters are then not any more independent of the parameters of the previous samples. The random generator will throw random values according to a Gaussian probability density with the mean given by the currently known best value for that particular parameter and the width in units of the interval size given by the option *Sigma*. Points which are created out of the parameter’s interval are mapped back into the interval.

### 6.2 Minuit minimisation

Minuit is the standard multivariate minimisation package used in HEP [17]. Its purpose is to find the minimum of a multi-parameter estimator function and to analyse the shape of the function around the minimum (error analysis). The principal application of the TMVA fitters is simple minimisation, while the shape of the minimum is irrelevant in most cases. The use of Minuit is therefore not necessarily the most efficient solution, but because it is a very robust tool we have included it here. Minuit searches the solution along the direction of the gradient until a minimum or an boundary is reached (MIGRAD algorithm). It does not attempt to find the global minimum but is satisfied with local minima. If during the error analysis with MINOS, the minimum smaller values than the local minimum might be obtained. In particular, the use of MINOS may as a side effect of an improved error analysis uncover a convergence in a local minimum, in which case MIGRAD minimisation is invoked once again. If multiple local and/or global solutions exist, it might be preferable to use any of the other fitters which are specifically designed for this type of problem.

The configuration options for Minuit are given in Option Table 6.

### 6.3 Genetic Algorithm

A Genetic Algorithm is a technique to find approximate solutions to optimisation or search problems. The problem is modelled by a group (*population*) of abstract representations (*genomes*) of possible
### 6.3 Genetic Algorithm

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ErrorLevel</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>TMinuit: error level: 0.5=logL fit, 1=chi-squared fit</td>
</tr>
<tr>
<td>PrintLevel</td>
<td>–</td>
<td>-1</td>
<td>–</td>
<td>TMinuit: output level: -1=least, 0, +1=all garbage</td>
</tr>
<tr>
<td>FitStrategy</td>
<td>–</td>
<td>2</td>
<td>–</td>
<td>TMinuit: fit strategy: 2=best</td>
</tr>
<tr>
<td>PrintWarnings</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>TMinuit: suppress warnings</td>
</tr>
<tr>
<td>UseImprove</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>TMinuit: use IMPROVE</td>
</tr>
<tr>
<td>UseMinos</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>TMinuit: use MINOS</td>
</tr>
<tr>
<td>SetBatch</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>TMinuit: use batch mode</td>
</tr>
<tr>
<td>MaxCalls</td>
<td>–</td>
<td>1000</td>
<td>–</td>
<td>TMinuit: approximate maximum number of function calls</td>
</tr>
<tr>
<td>Tolerance</td>
<td>–</td>
<td>0.1</td>
<td>–</td>
<td>TMinuit: tolerance to the function value at the minimum</td>
</tr>
</tbody>
</table>


solutions (*individuals*). By applying means similar to processes found in biological evolution the individuals of the population should evolve towards an optimal solution of the problem. Processes which are usually modelled in evolutionary algorithms — of which Genetic Algorithms are a subtype — are inheritance, mutation and “sexual recombination” (also termed crossover).

Apart from the abstract representation of the solution domain, a *fitness* function must be defined. Its purpose is the evaluation of the goodness of an individual. The fitness function is problem dependent. It either returns a value representing the individual’s goodness or it compares two individuals and indicates which of them performs better.

The Genetic Algorithm proceeds as follows:

- **Initialisation**: A starting population is created. Its size depends on the problem to be solved. Each individual belonging to the population is created by randomly setting the parameters of the abstract representation (variables), thus producing a point in the solution domain of the initial problem.

- **Evaluation**: Each individual is evaluated using the fitness function.

- **Selection**: Individuals are kept or discarded as a function of their fitness. Several selection procedures are possible. The simplest one is to separate out the worst performing fraction of the population. Another possibility is to decide on the individual’s survival by assigning probabilities that depend on the individual’s performance compared to the others.

- **Reproduction**: The surviving individuals are copied, mutated and crossed-over until the initial population size is reached again.
The TMVA Genetic Algorithm provides controls that are set through configuration options (cf. Table 7). The parameter PopSize determines the number of individuals created at each generation of the Genetic Algorithm. At the initialisation, all parameters of all individuals are chosen randomly. The individuals are evaluated in terms of their fitness, and each individual giving an improvement is immediately stored.

Individuals with a good fitness are selected to engender the next generation. The new individuals are created by crossover and mutated afterwards. Mutation changes some values of some parameters of some individuals randomly following a Gaussian distribution function. The width of the Gaussian can be altered by the parameter SC\_factor. The current width is multiplied by this factor when within the last SC\_steps generations more than SC\_rate improvements have been obtained. If there were SC\_rate improvements the width remains unchanged. Were there, on the other hand, less than SC\_rate improvements, the width is divided by SC\_factor. This allows to influence the speed of searching through the solution domain.

The cycle of evaluating the fitness of the individuals of a generation and producing a new generation is repeated until the improvement of the fitness within the last Steps has been less than ConvCrit.
The minimisation is then considered to have converged. The whole cycle from initialisation over fitness evaluation, selection, reproduction and determining the improvement is repeated \texttt{Cycles} times, before the Genetic Algorithm has finished.

\section*{Guidelines for optimising the GA}

\texttt{PopSize} is the most important value for enhancing the quality of the results. This value is by default set to 300, but can be increased to 1000 or more only limited by the resources available. The calculation time of the GA should increase with \texttt{O(PopSize)}.

\texttt{Steps} is set by default to 40. This value can be increased moderately to about 60. Time consumption increases non linearly but at least with \texttt{O(Steps)}.

\texttt{Cycles} is set by default to 3. In this case, the GA is called three times independently from each other. With \texttt{SaveBestCycle} and \texttt{SaveBestGen} it is possible to set the number of best results which shall be stored each cycle of even each generation. These stored results are reused in the last cycle. That way the last cycle is not independent from the others, but incorporates their best results. The number of cycles can be increased moderately to about 10. The time consumption of GA rises with about \texttt{O(Cycles)}.

\section*{6.4 Simulated Annealing}

Simulated Annealing also aims at solving a minimisation problem with several discrete or continuous, local or global minima. The algorithm is inspired by the process of annealing which occur in condensed matter physics. When first heating and then slowly cooling down a metal ("annealing") its atoms move towards a state of lowest energy, while for sudden cooling the atoms tend to freeze in intermediate states higher energy. For infinitesimal annealing activity the system will always converge in its global energy minimum (see, e.g., Ref. [18]). This physical principle can be converted into an algorithm to achieve slow, but correct convergence of an optimisation problem with multiple solutions. Recovery out of local minima is achieved by assigning the probability [19]

\[ p(\Delta E) \propto \exp\left( -\frac{\Delta E}{T} \right) , \]

(28)

to a perturbation of the parameters leading to a shift $\Delta E$ in the energy of the system. The probability of such perturbations to occur decreases with the size of a positive energy coefficient of the perturbation, and increases with the ambient temperature ($T$).

\section*{Guidelines for optimising SA}

The TMVA implementation of Simulated Annealing includes various different adaptive adjustments of the perturbation and temperature gradients. The adjustment procedure is chosen by setting \texttt{KernelTemp} to one of the following values.
• **Increasing Adaptive Approach** (IncAdaptive). The algorithm seeks local minima and explores their neighbourhood, while changing the ambient temperature depending on the number of failures in the previous steps. The performance can be improved by increasing the number of iteration steps (MaxCalls), or by adjusting the minimal temperature (MinTemp). Manual adjustments of the speed of the temperature increase (TempScale and AdaptiveSpeed) for individual data sets might also improve the performance.

• **Decreasing Adaptive Approach** (DecAdaptive). The algorithm calculates the initial temperature (based on the effectiveness of large steps) and defines a multiplier to ensure that the minimal temperature is reached in the requested number of iteration steps. The performance can be improved by adjusting the minimal temperature (MinTemp) and by increasing number of steps (MaxCalls).

• **Other Kernels.** Several other procedures to calculate the temperature change are also implemented. Each of them starts with the maximum temperature (MaxTemp) and decreases while changing the temperature according to:

\[
\text{Temperature}^{(k)} = \begin{cases} 
\text{Sqrt} : & \frac{\text{InitialTemp}}{\sqrt{k+2}} \cdot \text{TempScale} \\
\text{Log} : & \frac{\text{InitialTemp}}{\ln(k+2)} \cdot \text{TempScale} \\
\text{Homo} : & \frac{\text{InitialTemp}}{k+2} \cdot \text{TempScale} \\
\text{Sin} : & \frac{\sin(k/\text{TempScale})+1}{k+1} \cdot \text{InitialTemp} + \varepsilon \\
\text{Geo} : & \text{CurrentTemp} \cdot \text{TempScale}
\end{cases}
\]

Their performances can be improved by adjusting the initial temperature InitialTemp (\(=\) Temperature\(^{(k=0)}\)), the number of iteration steps (MaxCalls), and the multiplier that scales the temperature decrease (TempScale).

The configuration options for the Simulated Annealing fitter are given in Option Table 8.

### 6.5 Combined fitters

For MVA methods such as FDA, where parameters of a discrimination function are adjusted to achieve optimal classification performance (cf. Sec. 8.9), the user can choose to combine Minuit parameter fitting with Monte Carlo sampling or a Genetic Algorithm. While the strength of Minuit is the speedy detection of a nearby local minimum, it might not find a better global minimum. If several local minima exist Minuit will find different solutions depending on the start values for the fit parameters. When combining Minuit with Monte Carlo sampling or a Genetic Algorithm, Minuit uses starting values generated by these methods. The subsequent fits then converge in local minima. Such a combination is usually more efficient than the uncontrolled sampling used in Monte Carlo techniques. When combined with a Genetic Algorithm the user can benefit from the advantages of both methods: the Genetic Algorithm to roughly locate the global minimum, and Minuit to find an accurate solution for it (for an example see the FDA method).
### Option Table 8: Configuration options reference for fitting method: Simulated Annealing (SA).

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MaxCalls</td>
<td>–</td>
<td>100000</td>
<td>–</td>
<td>Maximum number of minimisation calls</td>
</tr>
<tr>
<td>InitialTemp</td>
<td>–</td>
<td>1e+06</td>
<td>–</td>
<td>Initial temperature</td>
</tr>
<tr>
<td>MinTemp</td>
<td>–</td>
<td>1e-06</td>
<td>–</td>
<td>Minimum temperature</td>
</tr>
<tr>
<td>Eps</td>
<td>–</td>
<td>1e-10</td>
<td>–</td>
<td>Epsilon</td>
</tr>
<tr>
<td>TempScale</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Temperature scale</td>
</tr>
<tr>
<td>AdaptiveSpeed</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Adaptive speed</td>
</tr>
<tr>
<td>TempAdaptiveStep</td>
<td>–</td>
<td>0.009875</td>
<td>–</td>
<td>Step made in each generation temperature adaptive</td>
</tr>
<tr>
<td>UseDefaultScale</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Use default temperature scale for temperature minimisation algorithm</td>
</tr>
<tr>
<td>UseDefaultTemp</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Use default initial temperature</td>
</tr>
</tbody>
</table>

The configuration options for the combined fit methods are the inclusive sum of all the individual fitter options. It is recommended to use Minuit in batch mode (option `SetBatch`) and without MINOS (option `!UseMinos`) to prevent TMVA from flooding the output with Minuit messages which cannot be turned off, and to speed up the individual fits. It is however important to note that the combination of MIGRAD and MINOS together is less susceptible to getting caught in local minima.

### 7 Boosting and Bagging

Boosting is a way of enhancing the classification and regression performance (and increasing the stability with respect to statistical fluctuations in the training sample) of typically weak MVA methods by sequentially applying an MVA algorithm to reweighted (boosted) versions of the training data and then taking a weighted majority vote of the sequence of MVA algorithms thus produced. It has been introduced to classification techniques in the early '90s [25] and in many cases this simple strategy results in dramatic performance increases.

Although one may argue that bagging (cf. Sec. 7.3) is not a genuine boosting algorithm, we include it in the same context and typically when discussing boosting we also refer to bagging. The most commonly boosted methods are decision trees. However, as described in Sec. 10.1 on page 147, any MVA method may be boosted with TMVA. Hence, although the following discussion refers to decision trees, it also applies to other methods. (Note however that “Gradient Boost” is only
available for decision trees).

### 7.1 Adaptive Boost (AdaBoost)

The most popular boosting algorithm is the so-called AdaBoost (adaptive boost) [26]. In a classification problem, events that were misclassified during the training of a decision tree are given a higher event weight in the training of the following tree. Starting with the original event weights when training the first decision tree, the subsequent tree is trained using a modified event sample where the weights of previously misclassified events are multiplied by a common boost weight $\alpha$.

The boost weight is derived from the misclassification rate, $\text{err}$, of the previous tree:

$$\alpha = \frac{1 - \text{err}}{\text{err}}.$$  \hspace{1cm} (29)

The weights of the entire event sample are then renormalised such that the sum of weights remains constant.

We define the result of an individual classifier as $h(x)$, with $(x$ being the tuple of input variables) encoded for signal and background as $h(x) = +1$ and $-1$, respectively. The boosted event classification $y_{\text{Boost}}(x)$ is then given by

$$y_{\text{Boost}}(x) = \frac{1}{N_{\text{collection}}} \cdot \sum_i N_{\text{collection}} \cdot \ln(\alpha_i) \cdot h_i(x),$$  \hspace{1cm} (30)

where the sum is over all classifiers in the collection. Small (large) values for $y_{\text{Boost}}(x)$ indicate a background-like (signal-like) event. Equation (30) represents the standard boosting algorithm.

AdaBoost performs best on weak classifiers, meaning small individual decision trees with a tree depth of often as small 2 or 3, that have very little discrimination power by themselves. Given such small trees, they are much less prone to overtraining compared to simple decision trees and as an ensemble outperform them typically by far. The performance is often further enhanced by forcing a “slow learning” and allowing a larger number of boost steps instead. The learning rate of the AdaBoost algorithm is controled by a parameter $\beta$ giving as an exponent to the boost weight $\alpha \rightarrow \alpha^\beta$, which can be modified using the configuration option string of the MVA method to be boosted (see Option Tables 25 and 27 on pages 131 and 131 for boosted decision trees, and Option Table 30 for general classifier boosting 10.1).

For regression trees, the AdaBoost algorithm needs to be modified. TMVA uses here the so-called AdaBoost.R2 algorithm [28]. The idea is similar to AdaBoost albeit with a redefined loss per event to account for the deviation of the estimated target value from the true one. Moreover, as there are no longer correctly and wrongly classified events, all events need to be reweighted depending on

---

17By construction, the error rate is $\text{err} \leq 0.5$ as the same training events used to classify the output nodes of the previous tree are used for the calculation of the error rate.
their individual loss, which – for event $k$ – is given by

- **Linear**: $L(k) = \frac{|y(k) - \hat{y}(k)|}{\max_{k'}(|y(k') - \hat{y}(k')|)}$, \hspace{1cm} (31)
- **Square**: $L(k) = \left[\frac{|y(k) - \hat{y}(k)|}{\max_{k'}(|y(k') - \hat{y}(k')|)}\right]^2$, \hspace{1cm} (32)
- **Exponential**: $L(k) = 1 - \exp\left[-\frac{|y(k) - \hat{y}(k)|}{\max_{k'}(|y(k') - \hat{y}(k')|)}\right]$. \hspace{1cm} (33)

The average loss of the classifier $y^{(i)}$ over the whole training sample, $\langle L \rangle^{(i)} = \sum_{k} w(k') L^{(i)}(k')$, can be considered to be the analogon to the error fraction in classification. Given $\langle L \rangle$, one computes the quantity $\beta^{(i)} = \langle L \rangle^{(i)}/(1 - \langle L \rangle^{(i)})$, which is used in the boosting of the events, and for the combination of the regression methods belonging to the boosted collection. The boosting weight, $w^{(i+1)}(k)$, for event $k$ and boost step $i + 1$ thus reads

$$w^{(i+1)}(k) = w^{(i)}(k) \cdot \beta^{1-L^{(i)}(k)}.$$ \hspace{1cm} (34)

The sum of the event weights is again renormalised to reproduce the original overall number of events. The final regressor, $y_{\text{Boost}}$, uses the weighted median, $\tilde{y}(i)$, where $(i)$ is chosen so that it is the minimal $(i)$ that satisfies the inequality

$$\sum_{t \in \text{sorted collection}} \ln \frac{1}{\beta(t)} \geq \frac{1}{2} \sum_{t} \ln \frac{1}{\beta(t)}.$$ \hspace{1cm} (35)

### 7.2 Gradient Boost

The idea of function estimation through boosting can be understood by considering a simple additive expansion approach. The function $F(x)$ under consideration is assumed to be a weighted sum of parametrised base functions $f(x; a_m)$, so-called “weak learners”. From a technical point of view any TMVA classifier could act as a weak learner in this approach, but decision trees benefit most from boosting and are currently the only classifier that implements GradientBoost (a generalisation may be included in future releases). Thus each base function in this expansion corresponds to a decision tree

$$F(x; P) = \sum_{m=0}^{M} \beta_m f(x; a_m); \hspace{1cm} P \in \{\beta_m; a_m\}_0^M.$$ \hspace{1cm} (36)

The boosting procedure is now employed to adjust the parameters $P$ such that the deviation between the model response $F(x)$ and the true value $y$ obtained from the training sample is minimised. The deviation is measured by the so-called loss-function $L(F, y)$, a popular choice being squared error loss $L(F, y) = (F(x) - y)^2$. It can be shown that the loss function fully determines the boosting procedure.

The most popular boosting method, AdaBoost, is based on exponential loss, $L(F, y) = e^{-F(x)y}$, which leads to the well known reweighting algorithm described in Sec. 7.1. Exponential loss has
the shortcoming that it lacks robustness in presence of outliers or mislabelled data points. The performance of AdaBoost therefore is expected to degrade in noisy settings.

The \textit{GradientBoost} algorithm attempts to cure this weakness by allowing for other, potentially more robust, loss functions without giving up on the good out-of-the-box performance of AdaBoost. The current TMVA implementation of GradientBoost uses the binomial log-likelihood loss

\begin{equation}
L(F,y) = \ln \left( 1 + e^{-2F(x)y} \right),
\end{equation}

for classification. As the boosting algorithm corresponding to this loss function cannot be obtained in a straightforward manner, one has to resort to a steepest-descent approach to do the minimisation. This is done by calculating the current gradient of the loss function and then growing a regression tree whose leaf values are adjusted to match the mean value of the gradient in each region defined by the tree structure. Iterating this procedure yields the desired set of decision trees which minimises the loss function. Note that GradientBoost can be adapted to any loss function as long as the calculation of the gradient is feasible.

Just like AdaBoost, GradientBoost works best on weak classifiers, meaning small individual decision trees with a depth of often just 2 to 4. Given such small trees, they are much less prone to overtraining compared to simple decision trees. Its robustness can be enhanced by reducing the learning rate of the algorithm through the \textit{Shrinkage} parameter (cf. Option Table 25 on page 131), which controls the weight of the individual trees. A small shrinkage (0.1–0.3) demands more trees to be grown but can significantly improve the accuracy of the prediction in difficult settings.

In certain settings GradientBoost may also benefit from the introduction of a bagging-like resampling procedure using random subsamples of the training events for growing the trees. This is called \textit{stochastic gradient boosting} and can be enabled by selecting the \textit{UseBaggedGrad} option. The sample fraction used in each iteration can be controlled through the parameter \textit{BaggingSampleFraction}, where typically the best results are obtained for values between 0.5 and 0.8.

For regression tasks, GradientBoost employs the Huber loss function \cite{47}, which features the robustness of least-absolute-deviation loss for error distributions with broad tails, while maintaining the high efficiency of least-squares loss for normally distributed errors. For moderately broad tails, it should surpass both least-squares and least-absolute-deviation loss.

\begin{equation}
L(F,y) = \begin{cases} 
\frac{1}{2} (y - F(x))^2 & |y - F| \leq \delta, \\
\delta(|y - F| - \delta/2) & |y - F| > \delta.
\end{cases}
\end{equation}

All GradientBoost options for classification described above can be also applied for regression tasks, but tests have shown that stochastic gradient boosting may not perform very well for regression problems.

### 7.3 Bagging

The term \textit{Bagging} denotes a resampling technique where a classifier is repeatedly trained using resampled training events such that the combined classifier represents an average of the individual
Bagging 69 classifiers. A priori, bagging does not aim at enhancing a weak classifier in the way adaptive or gradient boosting does, and is thus not a “boosting” algorithm in a strict sense. Instead it effectively smears over statistical representations of the training data and is hence suited to stabilise the response of a classifier. In this context it is often accompanied also by a significant performance increase compared to the individual classifier.

Resampling includes the possibility of replacement, which means that the same event is allowed to be (randomly) picked several times from the parent sample. This is equivalent to regarding the training sample as being a representation of the probability density distribution of the parent sample: indeed, if one draws an event out of the parent sample, it is more likely to draw an event from a region of phase-space that has a high probability density, as the original data sample will have more events in that region. If a selected event is kept in the original sample (that is when the same event can be selected several times), the parent sample remains unchanged so that the randomly extracted samples will have the same parent distribution, albeit statistically fluctuated. Training several classifiers with different resampled training data, and combining them into a collection, results in an averaged classifier that, just as for boosting, is more stable with respect to statistical fluctuations in the training sample.

Technically, resampling is implemented by applying random Poisson weights to each event of the parent sample.
8 The TMVA Methods

All TMVA classification and regression methods (in most cases, a method serves both analysis goals) inherit from `MethodBase`, which implements basic functionality like the interpretation of common configuration options, the interaction with the training and test data sets, I/O operations and common performance evaluation calculus. The functionality each MVA method is required to implement is defined in the abstract interface `IMethod`. Each MVA method provides a function that creates a rank object (of type `Ranking`), which is an ordered list of the input variables prioritised according to criteria specific to that method. Also provided are brief method-specific help notes (option `Help`, switched off by default) with information on the adequate usage of the method and performance optimisation in case of unsatisfying results.

If the option `CreateMVAPdfs` is set TMVA creates signal and background PDFs from the corresponding MVA response distributions using the training sample (cf. Sec. 3.1.7). The binning and smoothing properties of the underlying histograms can be customised via controls implemented in the `PDF` class (cf. Sec. 5 and Option Table 3 on page 56). The options specific to `MethodBase` are listed in Option Table 9. They can be accessed by all MVA methods.

The following sections describe the methods implemented in TMVA. For each method we proceed according to the following scheme: 

(i) a brief introduction, 
(ii) the description of the booking options required to configure the method, 
(iii) a description of the the method and TMVA implementation specifications for classification and – where available – for regression, 
(iv) the properties of the variable ranking, and 
(v) a few comments on performance, favourable (and disfavoured) use cases, and comparisons with other methods.

8.1 Rectangular cut optimisation

The simplest and most common classifier for selecting signal events from a mixed sample of signal and background events is the application of an ensemble of rectangular cuts on discriminating variables. Unlike all other classifiers in TMVA, the cut classifier only returns a binary response (signal or background). The optimisation of cuts performed by TMVA maximises the background rejection at given signal efficiency, and scans over the full range of the latter quantity. Dedicated analysis optimisation for which, e.g., the signal significance is maximised requires the expected signal and background yields to be known before applying the cuts. This is not the case for a

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18 Two constructors are implemented for each method: one that creates the method for a first time for training with a configuration (“option”) string among the arguments, and another that recreates a method from an existing weight file. The use of the first constructor is demonstrated in the example macros `TMVAClassification.C` and `TMVARegression.C`, while the second one is employed by the Reader in `TMVAClassificationApplication.C` and `TMVARegressionApplication.C`. Other functions implemented by each methods are: `Train` (called for training), `Write/ReadWeightsToStream` (I/O of specific training results), `WriteMonitoringHistosToFile` (additional specific information for monitoring purposes) and `CreateRanking` (variable ranking).

19 Note that cut optimisation is not a multivariate analyser method but a sequence of univariate ones, because no combination of the variables is achieved. Neither does a cut on one variable depend on the value of another variable (like it is the case for Decision Trees), nor can a, say, background-like value of one variable in a signal event be counterweighed by signal-like values of the other variables (like it is the case for the likelihood method).
8.1 Rectangular cut optimisation

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>V</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Verbose output (short form of VerbosityLevel below - overrides the latter one)</td>
</tr>
<tr>
<td>VerbosityLevel</td>
<td>–</td>
<td>Default</td>
<td>Default, Debug, Verbose, Info, Warning, Error, Fatal</td>
<td>Verbosity level</td>
</tr>
<tr>
<td>VarTransform</td>
<td>–</td>
<td>None</td>
<td>–</td>
<td>List of variable transformations performed before training, e.g., D_Background, P_Signal, G_N_AllClasses for: Decorrelation, PCA-transformation, Gaussianisation, Normalisation, each for the given class of events ('AllClasses' denotes all events of all classes, if no class indication is given, 'All' is assumed)</td>
</tr>
<tr>
<td>H</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Print method-specific help message</td>
</tr>
<tr>
<td>CreateMVAPdfs</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Create PDFs for classifier outputs (signal and background)</td>
</tr>
<tr>
<td>IgnoreNegWeightsInTraining</td>
<td>False</td>
<td>–</td>
<td>–</td>
<td>Events with negative weights are ignored in the training (but are included for testing and performance evaluation)</td>
</tr>
</tbody>
</table>

Option Table 9: Configuration options that are common for all classifiers (but which can be controlled individually for each classifier). Values given are defaults. If predefined categories exist, the default category is marked by a *. The lower options in the table control the PDF fitting of the classifiers (required, e.g., for the Rarity calculation).

multi-purpose discrimination and hence not used by TMVA. However, the cut ensemble leading to maximum significance corresponds to a particular working point on the efficiency curve, and can hence be easily derived after the cut optimisation scan has converged.\(^{20}\)

TMVA cut optimisation is performed with the use of multivariate parameter fitters interfaced by the class FitterBase (cf. Sec. 6). Any fitter implementation can be used, where however because of the peculiar, non-unique solution space only Monte Carlo sampling, Genetic Algorithm, and

\(^{20}\) Assuming a large enough number of events so that Gaussian statistics is applicable, the significance for a signal is given by

\[ S = \varepsilon_S N_S / \sqrt{\varepsilon_S N_S + \varepsilon_B(\varepsilon_S) N_B}, \]

where \(\varepsilon_S(B)\) and \(N_S(B)\) are the signal and background efficiencies for a cut ensemble and the event yields before applying the cuts, respectively. The background efficiency \(\varepsilon_B\) is expressed as a function of \(\varepsilon_S\) using the TMVA evaluation curve obtained from the test data sample. The maximum significance is then found at the root of the derivative

\[ \frac{dS}{d\varepsilon_S} = N_S \frac{2\varepsilon_B(\varepsilon_S) N_B + \varepsilon_S \left( N_S - \frac{dN_S}{d\varepsilon_S} \right) N_B}{2 \left( \varepsilon_S N_S + \varepsilon_B(\varepsilon_S) N_B \right)^{3/2}} = 0, \tag{39} \]

which depends on the problem.
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>FitMethod</td>
<td>-</td>
<td>GA</td>
<td>GA, SA, MC, MCEvents, MINUIT, EventScan</td>
<td>Minimisation Method (GA, SA, and MC are the primary methods to be used; the others have been introduced for testing purposes and are depreciated)</td>
</tr>
<tr>
<td>EffMethod</td>
<td>-</td>
<td>EffSel</td>
<td>EffSel, EffPDF</td>
<td>Selection Method</td>
</tr>
<tr>
<td>CutRangeMin</td>
<td>Yes</td>
<td>-1</td>
<td>-</td>
<td>Minimum of allowed cut range (set per variable)</td>
</tr>
<tr>
<td>CutRangeMax</td>
<td>Yes</td>
<td>-1</td>
<td>-</td>
<td>Maximum of allowed cut range (set per variable)</td>
</tr>
<tr>
<td>VarProp</td>
<td>Yes</td>
<td>NotEnforced</td>
<td>NotEnforced, FMax, FMin, FSmart</td>
<td>Categorisation of cuts</td>
</tr>
</tbody>
</table>

Option Table 10: Configuration options reference for MVA method: Cuts. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured.

Simulated Annealing show satisfying results. Attempts to use Minuit (SIMPLEX or MIGRAD) have not shown satisfactory results, with frequently failing fits.

The training events are sorted in binary trees prior to the optimisation, which significantly reduces the computing time required to determine the number of events passing a given cut ensemble (cf. Sec. 4.3).

### 8.1.1 Booking options

The rectangular cut optimisation is booked through the Factory via the command:

```plaintext
factory->BookMethod( Types::kCuts, "Cuts", "<options>" );
```

**Code Example 44**: Booking of the cut optimisation classifier: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ‘:’. See Sec. 3.1.5 for more information on the booking.

The configuration options for the various cut optimisation techniques are given in Option Table 10.
8.1.2 Description and implementation

The cut optimisation analysis proceeds by first building binary search trees for signal and background. For each variable, statistical properties like mean, root-mean-squared (RMS), variable ranges are computed to guide the search for optimal cuts. Cut optimisation requires an estimator that quantifies the goodness of a given cut ensemble. Maximising this estimator minimises (maximises) the background efficiency, \( \varepsilon_B \) (background rejection, \( r_B = 1 - \varepsilon_B \)) for each signal efficiency \( \varepsilon_S \).

All optimisation methods (fitters) act on the assumption that one minimum and one maximum requirement on each variable is sufficient to optimally discriminate signal from background (i.e., the signal is clustered). If this is not the case, the variables must be transformed prior to the cut optimisation to make them compliant with this assumption.

For a given cut ensemble the signal and background efficiencies are derived by counting the training events that pass the cuts and dividing the numbers found by the original sample sizes. The resulting efficiencies are therefore rational numbers that may exhibit visible discontinuities when the number of training events is small and an efficiency is either very small or very large. Another way to compute efficiencies is to parameterise the probability density functions of all input variables and thus to achieve continuous efficiencies for any cut value. Note however that this method expects the input variables to be uncorrelated! Non-vanishing correlations would lead to incorrect efficiency estimates and hence to underperforming cuts. The two methods are chosen with the option \texttt{EffMethod} set to \texttt{EffSel} and \texttt{EffPDF}, respectively.

Monte Carlo sampling

Each generated cut sample (cf. Sec. 6.1) corresponds to a point in the \((\varepsilon_S, r_B)\) plane. The \( \varepsilon_S \) dimension is (finely) binned and a cut sample is retained if its \( r_B \) value is larger than the value already contained in that bin. This way a reasonably smooth efficiency curve can be obtained if the number of input variables is not too large (the required number of MC samples grows with powers of \( 2^n_{\text{var}} \)).

Prior information on the variable distributions can be used to reduce the number of cuts that need to be sampled. For example, if a discriminating variable follows Gaussian distributions for signal and background, with equal width but a larger mean value for the background distribution, there is no useful minimum requirement (other than \(-\infty\)) so that a single maximum requirement is sufficient for this variable. To instruct TMVA to remove obsolete requirements, the option \texttt{VarProp[i]} must be used, where \([i]\) indicates the counter of the variable (following the order in which they have been registered with the Factory, beginning with 0) must be set to either \texttt{FMax} or \texttt{FMin}. TMVA is capable of automatically detecting which of the requirements should be removed. Use the option \texttt{VarProp[i]=FSmart} (where again \([i]\) must be replaced by the appropriate variable counter, beginning with 0). Note that in many realistic use cases the mean values between signal and background of a variable are indeed distinct, but the background can have large tails. In such a case, the removal of a requirement is inappropriate, and would lead to underperforming cuts.
Genetic Algorithm

Genetic Algorithm (cf. Sec. 6.3) is a technique to find approximate solutions to optimisation or search problems. Apart from the abstract representation of the solution domain, a fitness function must be defined. In cut optimisation, the fitness of a rectangular cut is given by good background rejection combined with high signal efficiency.

At the initialization step, all parameters of all individuals (cut ensembles) are chosen randomly. The individuals are evaluated in terms of their background rejection and signal efficiency. Each cut ensemble giving an improvement in the background rejection for a specific signal efficiency bin is immediately stored. Each individual’s fitness is assessed, where the fitness is largely determined by the difference of the best found background rejection for a particular bin of signal efficiency and the value produced by the current individual. The same individual that has at one generation a very good fitness will have only average fitness at the following generation. This forces the algorithm to focus on the region where the potential of improvement is the highest. Individuals with a good fitness are selected to produce the next generation. The new individuals are created by crossover and mutated afterwards. Mutation changes some values of some parameters of some individuals randomly following a Gaussian distribution function, etc. This process can be controlled with the parameters listed in Option Table 7, page 62.

Simulated Annealing

Cut optimisation using Simulated Annealing works similarly as for the Genetic Algorithm and achieves comparable performance. The same fitness function is used to estimator the goodness of a given cut ensemble. Details on the algorithm and the configuration options can be found in Sec. 6.4 on page 63.

8.1.3 Variable ranking

The present implementation of Cuts does not provide a ranking of the input variables.

8.1.4 Performance

The Genetic Algorithm currently provides the best cut optimisation convergence. However, it is found that with rising number of discriminating input variables the goodness of the solution found (and hence the smoothness of the background-rejections versus signal efficiency plot) deteriorates quickly. Rectangular cut optimisation should therefore be reduced to the variables that have the largest discriminating power.

If variables with excellent signal from background separation exist, applying cuts can be quite competitive with more involved classifiers. Cuts are known to underperform in presence of strong nonlinear correlations and/or if several weakly discriminating variables are used. In the latter case,
8.2 Projective likelihood estimator (PDE approach)

The method of maximum likelihood consists of building a model out of probability density functions (PDF) that reproduces the input variables for signal and background. For a given event, the likelihood for being of signal type is obtained by multiplying the signal probability densities of all input variables, which are assumed to be independent, and normalising this by the sum of the signal and background likelihoods. Because correlations among the variables are ignored, this PDE approach is also called “naive Bayes estimator”, unlike the full multidimensional PDE approaches such as PDE-range search, PDE-foam and k-nearest-neighbour discussed in the subsequent sections, which approximate the true Bayes limit.

8.2.1 Booking options

The likelihood classifier is booked via the command:

```cpp
factory->BookMethod( Types::kLikelihood, "Likelihood", "<options>" );
```

Code Example 45: Booking of the (projective) likelihood classifier: the first argument is the predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ‘:’. See Sec. 3.1.5 for more information on the booking.

The likelihood configuration options are given in Option Table 11.

8.2.2 Description and implementation

The likelihood ratio $y_L(i)$ for event $i$ is defined by

$$y_L(i) = \frac{L_S(i)}{L_S(i) + L_B(i)},$$  \hspace{1cm} (40)
where
\[ L_{SB}(i) = \prod_{k=1}^{n_{\text{var}}} p_{SB,k}(x_k(i)), \] (41)

and where \( p_{SB,k} \) is the signal (background) PDF for the \( k \)th input variable \( x_k \). The PDFs are normalised
\[ \int_{-\infty}^{+\infty} p_{SB,k}(x_k) dx_k = 1, \quad \forall k. \] (42)

It can be shown that in absence of model inaccuracies (such as correlations between input variables not removed by the de-correlation procedure, or an inaccurate probability density model), the ratio (40) provides optimal signal from background separation for the given set of input variables.

Since the parametric form of the PDFs is generally unknown, the PDF shapes are empirically approximated from the training data by nonparametric functions, which can be chosen individually for each variable and are either polynomial splines of various degrees fitted to histograms or unbinned kernel density estimators (KDE), as discussed in Sec. (5).

A certain number of primary validations are performed during the PDF creation, the results of which are printed to standard output. Among these are the computation of a \( \chi^2 \) estimator between all nonzero bins of the original histogram and its PDF, and a comparison of the number of outliers (in sigmas) found in the original histogram with respect to the (smoothed) PDF shape, with the statistically expected one. The fidelity of the PDF estimate can be also inspected visually by executing the macro likelihoodrefs.C (cf. Table 4).

**Transforming the likelihood output**

If a data-mining problem offers a large number of input variables, or variables with excellent separation power, the likelihood response \( y_L \) is often strongly peaked at 0 (background) and 1 (signal). Such a response is inconvenient for the use in subsequent analysis steps. TMVA therefore allows to transform the likelihood output by an inverse sigmoid function that zooms into the peaks
\[ y_L(i) \rightarrow y_L'(i) = -\tau^{-1} \ln(y_L^{-1} - 1), \] (43)

where \( \tau = 15 \) is used. Note that \( y_L'(i) \) is no longer contained within \([0,1]\) (see Fig. 14). The transformation (43) is enabled (disabled) with the booking option TransformOutput=True(False).

**8.2.3 Variable ranking**

The present likelihood implementation does not provide a ranking of the input variables.
8.2.4 Performance

Both the training and the application of the likelihood classifier are very fast operations that are suitable for large data sets.

The performance of the classifier relies on the accuracy of the likelihood model. Because high fidelity PDF estimates are mandatory, sufficient training statistics is required to populate the tails of the distributions. The neglect of correlations between input variables in the model (41), often leads to a diminution of the discrimination performance. While linear Gaussian correlations can be rotated away (see Sec. 4.1), such an ideal situation is rarely given. Positive correlations lead to peaks at both $y_L \to 0, 1$. Correlations can be reduced by categorising the data samples and building an independent likelihood classifier for each event category. Such categories could be geometrical regions in the detector, kinematic properties, etc. In spite of this, realistic applications with a large number of input variables are often plagued by irreducible correlations, so that projective likelihood approaches like the one discussed here are under-performing. This finding led to the development of the many alternative classifiers that exist in statistical theory today.

8.3 Multidimensional likelihood estimator (PDE range-search approach)

This is a generalization of the projective likelihood classifier described in Sec. 8.2 to $n_{\text{var}}$ dimensions, where $n_{\text{var}}$ is the number of input variables used. If the multidimensional PDF for signal and background (or regression data) were known, this classifier would exploit the full information contained in the input variables, and would hence be optimal. In practice however, huge training samples are necessary to sufficiently populate the multidimensional phase space.\footnote{Due to correlations between the input variables, only a sub-space of the full phase space may be populated.} Kernel estimation methods may be used to approximate the shape of the PDF for finite training statistics.

A simple probability density estimator denoted $PDE$ range search, or $PDE-RS$, has been suggested
The PDE for a given test event (discriminant) is obtained by counting the (normalised) number of training events that occur in the "vicinity" of the test event. The classification of the test event may then be conducted on the basis of the majority of the nearest training events. The $n_{\text{var}}$-dimensional volume that encloses the "vicinity" is user-defined and can be adaptive. A search method based on sorted binary trees is used to reduce the computing time for the range search. To enhance the sensitivity within the volume, kernel functions are used to weight the reference events according to their distance from the test event. PDE-RS is a variant of the k-nearest neighbour classifier described in Sec. 8.5.

### 8.3.1 Booking options

The PDE-RS classifier is booked via the command:

```cpp
factory->BookMethod( Types::kPDERS, "PDERS", "<options>" );
```

**Code Example 46:** Booking of PDE-RS: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ":". See Sec. 3.1.5 for more information on the booking.

The configuration options for the PDE-RS classifier are given in Option Table 12.

### 8.3.2 Description and implementation

#### Classification

To classify an event as being either of signal or of background type, a local estimate of the probability density of it belonging to either class is computed. The method of PDE-RS provides such an estimate by defining a volume ($V$) around the test event ($i$), and by counting the number of signal ($n_S(i, V)$) and background events ($n_B(i, V)$) obtained from the training sample in that volume. The ratio

$$y_{\text{PDE-RS}}(i, V) = \frac{1}{1 + r(i, V)},$$

is taken as the estimate, where $r(i, V) = (n_B(i, V)/N_B) \cdot (N_S/n_S(i, V))$, and $N_{S(B)}$ is the total number of signal (background) events in the training sample. The estimator $y_{\text{PDE-RS}}(i, V)$ peaks at 1 (0) for signal (background) events. The counting method averages over the PDF within $V$, and hence ignores the available shape information inside (and outside) that volume.

#### Binary tree search

Efficiently searching for and counting the events that lie inside the volume is accomplished with the use of a $n_{\text{var}}$-variable binary tree search algorithm [13] (cf. Sec. 4.3).
8.3 Multidimensional likelihood estimator (PDE range-search approach) 79

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>VolumeRangeMode</td>
<td>–</td>
<td>Adaptive</td>
<td>Unscaled, MinMax, RMS, Adaptive, kNN</td>
<td>Method to determine volume size</td>
</tr>
<tr>
<td>KernelEstimator</td>
<td>–</td>
<td>Box</td>
<td>Box, Sphere, Teepee, Gauss, Sinc3, Sinc5, Sinc7, Sinc9, Sinc11, Lanczos2, Lanczos3, Lanczos5, Lanczos8, Trim</td>
<td>Kernel estimation function</td>
</tr>
<tr>
<td>DeltaFrac</td>
<td>–</td>
<td>3</td>
<td>–</td>
<td>nEventsMin/Max for minmax and rms volume range</td>
</tr>
<tr>
<td>NEventsMin</td>
<td>–</td>
<td>100</td>
<td>–</td>
<td>nEventsMin for adaptive volume range</td>
</tr>
<tr>
<td>NEventsMax</td>
<td>–</td>
<td>200</td>
<td>–</td>
<td>nEventsMax for adaptive volume range</td>
</tr>
<tr>
<td>MaxVIterations</td>
<td>–</td>
<td>150</td>
<td>–</td>
<td>MaxVIterations for adaptive volume range</td>
</tr>
<tr>
<td>InitialScale</td>
<td>–</td>
<td>0.99</td>
<td>–</td>
<td>InitialScale for adaptive volume range</td>
</tr>
<tr>
<td>GaussSigma</td>
<td>–</td>
<td>0.1</td>
<td>–</td>
<td>Width (wrt volume size) of Gaussian kernel estimator</td>
</tr>
<tr>
<td>NormTree</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Normalize binary search tree</td>
</tr>
</tbody>
</table>

Option Table 12: Configuration options reference for MVA method: PDE-RS. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured.

Choosing a volume

The TMVA implementation of PDE-RS optionally provides four different volume definitions selected via the configuration option VolumeRangeMode.

- **Unscaled**
  The simplest volume definition consisting of a rigid box of size DeltaFrac, in units of the variables. This method was the one originally used by the developers of PDE-RS [14].

- **MinMax**
  The volume is defined in each dimension (i.e., input variable) with respect to the full range of values found for that dimension in the training sample. The fraction of this volume used for the range search is defined by the option DeltaFrac.

- **RMS**
  The volume is defined in each dimension with respect to the RMS of that dimension (input
variable), estimated from the training sample. The fraction of this volume used for the range search is defined by the option DeltaFrac.

- **Adaptive**
  A volume is defined in each dimension with respect to the RMS of that dimension, estimated from the training sample. The overall scale of the volume is adjusted individually for each test event such that the total number of events confined in the volume lies within a user-defined range (options NEventsMin/Max). The adjustment is performed by the class RootFinder, which is a C++ implementation of Brent's algorithm (translated from the CERNLIB function RZERO). The maximum initial volume (fraction of the RMS) and the maximum number of iterations for the root finding is set by the options InitialScale and MaxVIterations, respectively. The requirement to collect a certain number of events in the volume automatically leads to small volume sizes in strongly populated phase space regions, and enlarged volumes in areas where the population is scarce.

Although the adaptive volume adjustment is more flexible and should perform better, it significantly increases the computing time of the PDE-RS discriminant. If found too slow, one can reduce the number of necessary iterations by choosing a larger NEventsMin/Max interval.

**Event weighting with kernel functions**

One of the shortcomings of the original PDE-RS implementation is its sensitivity to the exact location of the sampling volume boundaries: an infinitesimal change in the boundary placement can include or exclude a training event, thus changing \( r(i,V) \) by a finite amount.\(^{22} \) In addition, the shape information within the volume is ignored.

Kernel functions mitigate these problems by weighting each event within the volume as a function of its distance to the test event. The farer it is away, the smaller is its weight. The following kernel functions are implemented in TMVA, and can be selected with the option KernelEstimator.

- **Box**
  Corresponds to the original rectangular volume element without application of event weights.

- **Sphere**
  A hyper-elliptic volume element is used without application of event weights. The hyper-ellipsoid corresponds to a sphere of constant fraction in the MinMax or RMS metrics. The size of the sphere can be chosen adaptive, just as for the rectangular volume.

- **Teepee**
  The simplest linear interpolation that eliminates the discontinuity problem of the box. The training events are given a weight that decreases linearly with their distance from the centre of the volume (the position of the test event). In other words: these events are convolved with the triangle or tent function, becoming a sort of teepee in multi-dimensions.

\(^{22}\)Such an introduction of artefacts by having sharp boundaries in the sampled space is an example of Gibbs's phenomenon, and is commonly referred to as ringing or aliasing.
8.3 Multidimensional likelihood estimator (PDE range-search approach)

- **Trim**
  Modified Teepee given by the function \((1 - \hat{x}^3)^3\), where \(\hat{x}\) is the normalised distance between test event and reference.

- **Gauss**
  The simplest well behaved convolution kernel. The width of the Gaussian (fraction of the volume size) can be set by the option `GaussSigma`.

Other kernels implemented for test purposes are “Sinc” and ”Lanczos” functions \(\propto \sin x/x\) of different (symmetric) orders. They exhibit strong peaks at zero and oscillating tails. The Gaussian and Teepee kernel functions are shown in Fig. 15.

**Regression**

Regression with PDE-RS proceeds similar to classification. The difference lies in the replacement of Eq. (44) by the average target value of all events belonging to the volume \(V\) defined by event \(i\) (the test event)

\[
y_{\text{PDE-RS}}(i, V) = \langle t(i, V) \rangle = \frac{\sum_{j \in V} w_j t_j f(\text{dis}(i, j))}{\sum_{j \in V} w_j f(\text{dis}(i, j))},
\]

where the sum is over all training events in \(V\), \(w_j\) and \(t_j\) are the weight and target value of event \(j\) in \(V\), \(\text{dis}(i, j)\) is a measure of the distance between events \(i\) and \(j\), and \(f(\ldots)\) is a kernel function.

**8.3.3 Variable ranking**

The present implementation of PDE-RS does not provide a ranking of the input variables.
8.3.4 Performance

As opposed to many of the more sophisticated data-mining approaches, which tend to present the user with a "black box", PDE-RS is simple enough that the algorithm can be easily traced and tuned by hand. PDE-RS can yield competitive performance if the number of input variables is not too large and the statistics of the training sample is ample. In particular, it naturally deals with complex nonlinear variable correlations, the reproduction of which may, for example, require involved neural network architectures.

PDE-RS is a slowly responding classifier. Only the training, i.e., the fabrication of the binary tree is fast, which is usually not the critical part. The necessity to store the entire binary tree in memory to avoid accessing virtual memory limits the number of training events that can effectively be used to model the multidimensional PDF. This is not the case for the other classifiers implemented in TMVA (with some exception for Boosted Decision Trees).

8.4 Likelihood estimator using self-adapting phase-space binning (PDE-Foam)

The PDE-Foam method [16] is an extension of PDE-RS, which divides the multi-dimensional phase space in a finite number of hyper-rectangles (cells) of constant event density. This “foam” of cells is filled with averaged probability density information sampled from the training data. For a given number of cells, the binning algorithm adjusts the size and position of the cells inside the multi-dimensional phase space based on a binary split algorithm that minimises the variance of the event density in the cell. The binned event density information of the final foam is stored in cells, organised in a binary tree, to allow a fast and memory-efficient storage and retrieval of the event density information necessary for classification or regression. The implementation of PDE-Foam is based on the Monte-Carlo integration package T Foam [15] included in ROOT.

In classification mode PDE-Foam forms bins of similar density of signal and background events or the ratio of signal to background. In regression mode the algorithm determines cells with small varying regression targets. In the following, we use the term density ($\rho$) for the event density in case of classification or for the target variable density in case of regression.

8.4.1 Booking options

The PDE-Foam classifier is booked via the command:

```c++
factory->BookMethod( Types::kPDEFoam, "PDEFoam", "<options>" );
```

Code Example 47: Booking of PDE-Foam: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ‘:’. See Sec. 3.1.5 for more information on the booking.
The configuration options for the PDE-Foam method are summarised in Option Table 13. Table 5 gives an overview of supported combinations of configuration options.

<table>
<thead>
<tr>
<th>Option</th>
<th>Classification</th>
<th>Regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>SigBgSeparate</td>
<td>True</td>
<td>False</td>
</tr>
<tr>
<td>MultiTargetRegression</td>
<td>–</td>
<td>False</td>
</tr>
<tr>
<td>Kernel</td>
<td>★</td>
<td>★</td>
</tr>
<tr>
<td>TargetSelection</td>
<td>◦</td>
<td>◦</td>
</tr>
<tr>
<td>TailCut</td>
<td>★</td>
<td>★</td>
</tr>
<tr>
<td>UseYesNoCell</td>
<td>★</td>
<td>★</td>
</tr>
<tr>
<td>MaxDepth</td>
<td>★</td>
<td>★</td>
</tr>
</tbody>
</table>

**Table 5:** Availability of options for the two classification and two regression modes of PDE-Foam. Supported options are marked by a ‘★’, while disregarded ones are marked by a ◦.

### 8.4.2 Description and implementation of the foam algorithm

Foams for an arbitrary event sample are formed as follows.

1. **Setup of binary search trees.** A binary search tree is created and filled with the \(d\)-dimensional event tuples form the training sample as for the PDE-RS method (cf. Sec. 8.3).

2. **Initialisation phase.** A foam is created, which at first consists of one \(d\)-dimensional hyper-rectangle (base cell). The coordinate system of the foam is normalised such that the base cell extends from 0 to 1 in each dimension. The coordinates of the events in the corresponding training tree are linearly transformed into the coordinate system of the foam.

3. **Growing phase.** A binary splitting algorithm iteratively splits cells of the foam along axis-parallel hyperplanes until the maximum number of active cells (set by \(n\text{ActiveCells}\)) is reached. The splitting algorithm minimises the relative variance of the density \(\sigma_\rho/\langle \rho \rangle\) across each cell (cf. Ref. [15]). For each cell \(n\text{Samp1}\) random points uniformly distributed over the cell volume are generated. For each of these points a small box centred around this point is defined. The box has a size of \(\text{VolFrac}\) times the size of the base cell in each dimension. The density is estimated as the number of events contained in this box divided by the volume of the box.\(^{23}\) The densities obtained for all sampled points in the cell are projected onto the \(d\) axes of the cell and the projected values are filled in histograms with \(n\text{Bin}\) bins. The next cell to be split and the corresponding division edge (bin) for the split are selected as the ones that have the largest relative variance. The two new daughter cells are marked as ‘active’ cells and the old mother cell is marked as ‘inactive’. A detailed description of the splitting algorithm can be found in Ref. [15]. The geometry of the final foam reflects the distribution of the training samples: phase-space regions where the density is constant are combined in large

\(^{23}\)In case of regression this is the average target value computed according to Eq. (45), page 81.
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SigBgSeparate</td>
<td>-</td>
<td>False</td>
<td>-</td>
<td>Separate foams for signal and background</td>
</tr>
<tr>
<td>TailCut</td>
<td>-</td>
<td>0.001</td>
<td>-</td>
<td>Fraction of outlier events that are excluded from the foam in each dimension</td>
</tr>
<tr>
<td>VolFrac</td>
<td>-</td>
<td>0.0666667</td>
<td>-</td>
<td>Size of sampling box, used for density calculation during foam build-up</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(maximum value: 1.0 is equivalent to volume of entire foam)</td>
</tr>
<tr>
<td>nActiveCells</td>
<td>-</td>
<td>500</td>
<td>-</td>
<td>Maximum number of active cells to be created by the foam</td>
</tr>
<tr>
<td>nSamp1</td>
<td>-</td>
<td>2000</td>
<td>-</td>
<td>Number of generated MC events per cell</td>
</tr>
<tr>
<td>nBin</td>
<td>-</td>
<td>5</td>
<td>-</td>
<td>Number of bins in edge histograms</td>
</tr>
<tr>
<td>Compress</td>
<td>-</td>
<td>True</td>
<td>-</td>
<td>Compress foam output file</td>
</tr>
<tr>
<td>MultiTargetRegression</td>
<td>-</td>
<td>False</td>
<td>-</td>
<td>Do regression with multiple targets</td>
</tr>
<tr>
<td>Nmin</td>
<td>-</td>
<td>100</td>
<td>-</td>
<td>Number of events in cell required to split cell</td>
</tr>
<tr>
<td>MaxDepth</td>
<td>-</td>
<td>0</td>
<td>-</td>
<td>Maximum depth of cell tree (0=unlimited)</td>
</tr>
<tr>
<td>FillFoamWithOrigWeights</td>
<td>-</td>
<td>False</td>
<td>-</td>
<td>Fill foam with original or boost weights</td>
</tr>
<tr>
<td>UseYesNoCell</td>
<td>-</td>
<td>False</td>
<td>-</td>
<td>Return -1 or 1 for bkg or signal like events</td>
</tr>
<tr>
<td>DTLogic</td>
<td>-</td>
<td>None</td>
<td>None, GiniIndex,</td>
<td>Use decision tree algorithm to split cell</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>MisClassificationError, CrossEntropy, GiniIndexWithLaplace, SdivSqrtS-plusB</td>
<td></td>
</tr>
<tr>
<td>Kernel</td>
<td>-</td>
<td>None</td>
<td>None, Gauss, LinNeighbors</td>
<td>Kernel type used</td>
</tr>
<tr>
<td>TargetSelection</td>
<td>-</td>
<td>Mean</td>
<td>Mean, Mpv</td>
<td>Target selection method</td>
</tr>
</tbody>
</table>

Option Table 13: Configuration options reference for MVA method: *PDE-Foam*. The options in Option Table 9 on page 71 can also be configured.
Figure 16: Projections of a two-dimensional foam with 500 cells for a Gaussian distribution on a two-dimensional histogram. The foam was created with 5000 events from the input tree. (a) shows the reconstructed distribution without kernel weighting and (b) shows the distribution weighted with a Gaussian kernel. The grey shades indicate the event density of the drawn cell. For more details about the projection function see the description on page 92.

4. **Filling phase.** Each active cell is filled with values that classify the event distribution within this cell and which permit the calculation of the classification or regression discriminator.

5. **Evaluation phase.** The estimator for a given event is evaluated based on the information stored in the foam cells. The corresponding foam cell, in which the event variables (\(d\)-dimensional vector) of a given event is contained, is determined with a binary search algorithm.\(^{24}\)

The initial trees which contain the training events and which are needed to evaluate the densities for the foam build-up, are discarded after the training phase. The memory consumption for the foam is 160 bytes per foam cell plus an overhead of 1.4 kbytes for the PDE-Foam object on a 64 bit architecture. Note that in the foam all cells created during the growing phase are stored within a binary tree structure. Cells which have been split are marked as inactive and remain empty. To reduce memory consumption, the cell geometry is not stored with the cell, but rather obtained recursively from the information about the division edge of the corresponding mother cell. This way only two short integer numbers per cell represent the information of the entire foam geometry: the division coordinate and the bin number of the division edge.

**PDE-Foam options**

- **TailCut** – *boundaries for foam geometry*

\(^{24}\)For an event that falls outside the foam boundaries, the cell with the smallest Cartesian distance to the event is chosen.
A parameter TailCut is used to define the geometry of the base cell(s) such that outliers of
the distributions in the training ensemble are not included in the volume of the base cell(s).
In a first step, the upper and lower limits for each input variable are determined from the
training sample. Upper and a lower bounds are then determined for each input variable,
such that on both sides a fraction TailCut of all events is excluded. The default value of
TailCut=0.001 ensures a sufficient suppression of outliers for most input distributions. For
cases where the input distributions have a fixed range or where they are discontinuous and/or
have peaks towards the edges, it can be appropriate to set TailCut to 0. Note that for event
classification it is guaranteed that the foam has an infinite coverage: events outside the foam
volume are assigned to the cell with the smallest Cartesian distance to the event.

- **nActiveCells** – maximum number of active foam cells
  In most cases larger nActiveCells values result in more accurate foams, but also lead to
  longer computation time during the foam formation, and require more storage memory. The
default value of 500 was found to be a good compromise for most practical applications if the
size of the training samples is of the order of $10^4$ events. Note that the total number of cells,
nCells, is given as $nCells = nActiveCells \cdot 2 - 1$, since the binary-tree structure of the foam
requires all inactive cells to remain part of the foam (see Growing phase).

- **VolFrac** – size of the probe volume for the density sampling of the training data
  The volume is defined as a $d$-dimensional box with edge length VolFrac times the extension
  of the base cell in each dimension. The default of $1/15$ results in a box with volume $1/15^d$
times the volume of the base cell. Smaller values of VolFrac increase the sampling speed,
but also make the algorithm more vulnerable to statistical fluctuations in the training sample
(overtraining). In cases with many observables ($> 5$) and small training samples ($< 10^4$),
VolFrac should be increased for better classification results.

- **nSampl** – number of samplings per cell and per cell-division step
  The computation time for the foam formation scales linearly with the number of samplings.
The default of 2000 was found to give sufficiently accurate estimates of the density distributions
with an acceptable computation time.\footnote{Contrary to the original application where an analytical function is used, increasing the number of samplings does not automatically lead to better accuracy. Since the density is defined by a limited event sample, it may occur that always the same event is found for all sample points.}

- **nBin** – number of bins for edge histograms
  The number of bins for the edge histograms used to determine the variance of the sampled
density distributions within one cell are set with the parameter nBin. The performance in
typical applications was found to be rather insensitive to the number of bins. The default value
of 5 gives the foam algorithm sufficient flexibility to find the division points, while maintaining
sufficient statistical accuracy also in case of small event samples.

- **Nmin** – Minimal number of events for cell split
  If Nmin is set to a value greater than zero, the foam will only consider cells with a number of
events greater than Nmin to be split. If no more cells are found during the growing phase for
which this condition is fulfilled, the foam will stop splitting cells, even if the target number of
cells is not yet reached. This option prevents the foam from adapting to statistical fluctuations
in the training samples (overtraining). Note that event weights are not taken into account for evaluating the number of events in the cell.

In particular for training samples with small event numbers of less than $10^4$ events this cut improves the performance. The default value of ($\text{Nmin}=100$) was found to give good results in most cases. It should be reduced in applications with very small training samples (less than 200 training events) and with many cells.

- **Kernel – cell weighting with kernel functions:**
  A Gaussian Kernel smoothing is applied during the evaluation phase, if the option Kernel is set to “Gauss”. In this case all cells contribute to the calculation of the discriminant for a given event, weighted with their Gaussian distance to the event. The average cell value $v$ (event density in case of separated foams, and the ratio $n_S/(n_S + n_B)$ in case of a single foam) for a given event $x = (x_1, \ldots, x_{n_{\text{var}}})$ is calculated by

  $$v = \frac{\sum_{\text{all cells } i} G(D(i, x), 0, \text{VolFrac}) \cdot v_i}{\sum_{\text{all cells } i} G(D(i, x), 0, \text{VolFrac}),}$$  (46)

  where $v_i$ is the output value in cell $i$, $G(x, x_0, \sigma) = \exp(-(x - x_0)^2/2\sigma^2)$, and $D(i, x)$ is the minimal Euclidean distance between $x$ and any point $y$ in cell $i$

  $$D(i, x) = \min_{y \in \text{cell } i} |x - y|.$$  (47)

  The Gaussian kernel avoids discontinuities of the discriminant values at the cell boundaries. In most cases it results in an improved separation power between signal and background. However, the time needed for classification increases due to the larger number of computations performed. A comparison between foams with and without Gaussian kernel can be seen in Fig. 16.

  A linear interpolation with adjacent cells in each dimension is applied during the classification phase, if the option Kernel is set to “LinNeighbors”. This results in faster classification than the Gauss weighting of all cells in the foam.

- **UseYesNoCell – Discrete classification output**
  If UseYesNoCell is set to False (default), the discriminant (52) is returned as classification output. If UseYesNoCell is set to True, $-1$ is returned for events with discriminant $< 0.5$ (background-like) and $+1$ is returned for events with $\geq 0.5$ (signal-like).

- **MaxDepth – Maximum cell tree depth**
  The cell tree depth can be limited by using this option. When MaxDepth is set to an integer value greater than zero, the created cell tree will not be deeper than MaxDepth. By convention the root node has depth 1, which implies that a foam with 2 active cells (3 cells in total) has depth 2. If $n_{\text{ActiveCells}} \geq 2^{\text{MaxDepth} - 1}$ the resulting cell tree will be completely balanced. When MaxDepth is set to 0, the cell tree depth is not limited.

- **DTLogic – Cell split algorithm**
  In order to emulate a decision tree-like cell splitting algorithm, the option DTLogic was introduced. When set to GiniIndex, MisClassificationError or CrossEntropy, the algorithm
projects all events in a cell onto the cell edges and probes nBin-1 division points such that the separation gain

\[ \text{gain(parent cell)} - \text{gain(daughter cell 1)} - \text{gain(daughter cell 2)} \] (48)

is maximal (see Sec. 8.13.3). For a given separation type and a given cell the gain is defined as

\begin{align*}
\text{GiniIndex} : & \quad \text{gain(cell)} = p(1 - p), \\
\text{MisClassificationError} : & \quad \text{gain(cell)} = 1 - \max(p, 1 - p), \\
\text{CrossEntropy} : & \quad \text{gain(cell)} = -p \log p - (1 - p) \log(1 - p),
\end{align*}

(49) (50) (51)

where \( p = n_S/(n_S + n_B) \) in the considered cell. When DTLogic is set to None (default), the PDE-Foam cell splitting algorithm (see Sec. 8.4.2) is used. This option is available only for classification and is an alternative to the classification with two separate foams or one single foam (see Sec. 8.4.3).

- **FillFoamWithOrigWeights** – **Event weights for boosting**

Since the PDE-Foam cells are filled with events after the splitting, it is in principle possible to use different event weights for the filling than for the foam build-up. The option FillFoamWithOrigWeights was created to choose either the original or the full event weight (including the boost weight) to be filled into the foam cells after the build-up. This option is only relevant for boosting, because for non-boosted classifiers the boost weights are always 1. When setting FillFoamWithOrigWeights=T, one would only boost the foam geometry, instead of the cell content. This would slow down the boosting process, because the boost weights are ignored in the classification. In most cases studied FillFoamWithOrigWeights=T leads to worse classification results than FillFoamWithOrigWeights=F for the same number of boosts. However, when using stronger boosting by choosing AdaBoostBeta accordingly, filling the original weights into the foam can improve the performance.

The PDE-Foam algorithm exhibits stable performance with respect to variations in most of the parameter settings. However, optimisation of the parameters is required for small training samples (< 10^4 events) in combination with many observables (> 5). In such cases, VolFrac should be increased until an acceptable performance is reached. Moreover, in cases where the classification time is not critical, one of the Kernel methods should be applied to further improve the classification performance. For large training samples (> 10^5) and if the training time is not critical, nActiveCells should be increased to improve the classification performance.

### 8.4.3 Classification

To classify an event in a \( d \)-dimensional phase space as being either of signal or of background type, a local estimator of the probability that this event belongs to either class can be obtained from the foam’s hyper-rectangular cells. The foams are created and filled based on samples of signal and background training events. For classification two possibilities are implemented. One foam can be used to separate the S/B probability density or two separate foams are created, one for the signal events and one for background events.
1) Separate signal and background foams

If the option SigBgSeparate=True is set, the method PDE-Foam treats the signal- and background distributions separately and performs the following steps to form the two foams and to calculate the classifier discriminator for a given event:

1. **Setup of training trees.** Two binary search trees are created and filled with the $d$-dimensional observable vectors of all signal and background training events, respectively.

2. **Initialisation phase.** Two independent foams for signal and background are created.

3. **Growing phase.** The growing is performed independently for the two foams. The density of events is estimated as the number of events found in the corresponding tree that are contained in the sampling box divided by the volume of the box (see VolFrac option). The geometries of the final foams reflect the distribution of the training samples: phase-space regions where the density of events is constant are combined in large cells, while in regions with large gradients in density many small cells are created.

4. **Filling phase.** Both for the signal and background foam each active cell is filled with the number of training events, $n_S$ (signal) or $n_B$ (background), contained in the corresponding cell volume, taking into account the event weights $w_i$: $n_S = \sum_{\text{sig. cell}}^{} w_i$, $n_B = \sum_{\text{bg. cell}}^{} w_i$.

5. **Evaluation phase.** The estimator for a given event is evaluated based on the number of events stored in the foam cells. The two corresponding foam cells that contain the event are found. The number of events ($n_S$ and $n_B$) is read from the cells. The estimator $y_{\text{PDE-Foam}}(i)$ is then given by

$$y_{\text{PDE-Foam}}(i) = \frac{n_S/V_S}{n_B/V_B + n_S/V_S}, \tag{52}$$

where $V_S$ and $V_B$ are the respective cell volumes. The statistical error of the estimator is calculated as:

$$\sigma_{y_{\text{PDE-Foam}}}(i) = \sqrt{\left(\frac{n_S\sqrt{n_B}}{(n_S + n_B)^2}\right)^2 + \left(\frac{n_B\sqrt{n_S}}{(n_S + n_B)^2}\right)^2}. \tag{53}$$

Note that the so defined discriminant approximates the probability for an event from within the cell to be of class signal, if the weights are normalised such that the total number of weighted signal events is equal to the total number of weighted background events. This can be enforced with the normalisation mode “EqualNumEvents” (cf. Sec. 3.1.4).

Steps 1-4 correspond to the training phase of the method. Step 5 is performed during the testing phase. In contrast to the PDE-RS method the memory consumption and computation time for the testing phase does not depend on the number of training events.

Two separate foams for signal and background allow the foam algorithm to adapt the foam geometries to the individual shapes of the signal and background event distributions. It is therefore well suited for cases where the shapes of the two distributions are very different.
2) Single signal and background foam

If the option \texttt{SigBgSeparate=False} is set (default), the PDE-Foam method creates only one foam, which holds directly the estimator instead of the number of signal and background events. The differences with respect to separate signal and backgrounds foams are:

1. \textit{Setup of training trees}. Fill only one binary search tree with both signal events and background events. This is possible, since the binary search tree has the information whether a event is of signal or background type.

2. \textit{Initialisation phase}. Only one foam is created. The cells of this foam will contain the estimator \( y_{\text{PDE-Foam}}(i) \) (see eq. (52)).

3. \textit{Growing phase}. The splitting algorithm in this case minimises the variance of the estimator density \( \sigma_{\rho}/\langle \rho \rangle \) across each cell. The estimator density \( \rho \) is sampled as the number of weighted signal events \( n_S \) over the total number of weighted events \( n_S + n_B \) in a small box around the sampling points:

\[
\rho = \frac{n_S}{n_S + n_B} \frac{1}{\text{VolFrac}}
\]

In this case the geometries of the final foams reflect the distribution of the estimator density in the training sample: phase-space regions where the signal to background ratio is constant are combined in large cells, while in regions where the signal-to-background ratio changes rapidly many small cells are created.

4. \textit{Filling phase}. Each active cell is filled with the estimator given as the ratio of weighted signal events to the total number of weighted events in the cell:

\[
y_{\text{PDE-Foam}}(i) = \frac{n_S}{n_S + n_B}.
\]

The statistical error of the estimator (53) also is stored in the cell.

5. \textit{Evaluation phase}. The estimator for a given event is directly obtained as the discriminator that is stored in the cell which contains the event.

For the same total number of foam cells, the performance of the two implementations was found to be similar.

8.4.4 Regression

Two different methods are implemented for regression. In the first method, applicable for single targets only (\textit{mono-target regression}), the target value is stored in each cell of the foam. In the second method, applicable to any number of targets (\textit{multi-target regression}), the target values are stored in higher foam dimensions.

In \textit{mono-target regression} the density used to form the foam is given by the mean target density in a given box. The procedure is as follows.
1. **Setup of training trees.** A binary search tree is filled with all training events.

2. **Growing phase.** One \( n_{\text{var}} \)-dimensional foam is formed: the density \( \rho \) is given by the mean target value \( \langle t \rangle \) within the sampling box, divided by the box volume (given by the \texttt{VolFrac} option):

\[
\rho = \frac{\langle t \rangle}{\text{VolFrac}} \equiv \frac{\sum_{i=1}^{N_{\text{box}}} t_i}{N_{\text{box}} \cdot \text{VolFrac}},
\]

where the sum is over all \( N_{\text{box}} \) events within the sampling box, and \( t_i \) is the target value of event \( i \).

3. **Filling phase.** The cells are filled with their average target values, \( \langle t \rangle = \sum_{i=1}^{N_{\text{box}}} t^{(i)}/N_{\text{box}} \).

4. **Evaluation phase.** Estimate the target value for a given test event: find the corresponding foam cell in which the test event is situated and read the average target value \( \langle t \rangle \) from the cell.

For **multi-target regression** the target information is stored in additional foam dimensions. For a training sample with \( n_{\text{var}} \) (\( n_{\text{tar}} \)) input variables (regression targets), one would form a \( (n_{\text{var}} + n_{\text{tar}}) \)-dimensional foam. To compute a target estimate for event \( i \), one needs the coordinates of the cell centre \( C(i,k) \) in each foam dimension \( k \). The procedure is as follows.

1. **Setup of training trees.** A binary search tree is filled with all training events.

2. **Growing phase.** A \( (n_{\text{var}} + n_{\text{tar}}) \)-dimensional foam is formed: the event density \( \rho \) is estimated by the number of events \( N_{\text{box}} \) within a predefined box of the dimension \( (n_{\text{var}} + n_{\text{tar}}) \), divided by the box volume, whereas the box volume is given by the \texttt{VolFrac} option

\[
\rho = \frac{N_{\text{box}}}{\text{VolFrac}}.
\]

3. **Filling phase.** Each foam cell is filled with the number of corresponding training events.

4. **Evaluation phase.** Estimate the target value for a given test event: find the \( N_{\text{cells}} \) foam cells in which the coordinates \( (x_1, \ldots, x_{n_{\text{var}}}) \) of the event vector are situated. Depending on the \texttt{TargetSelection} option, the target value \( t_k \) \( (k = 1, \ldots, n_{\text{tar}}) \) is

   (a) the coordinate of the cell centre in direction of the target dimension \( n_{\text{var}} + k \) of the cell \( j \) \( (j = 1, \ldots, N_{\text{cells}}) \), which has the maximum event density

\[
t_k = C(j, n_{\text{var}} + k),
\]

   if \texttt{TargetSelection=}\texttt{Mpv}.

   (b) the mean cell centre in direction of the target dimension \( n_{\text{var}} + k \) weighted by the event densities \( d_{\text{ev}}(j) \) \( (j = 1, \ldots, N_{\text{cells}}) \) of the cells

\[
t_k = \frac{\sum_{j=1}^{N_{\text{cells}}} C(j, n_{\text{var}} + k) \cdot d_{\text{ev}}(j)}{\sum_{j=1}^{N_{\text{cells}}} d_{\text{ev}}(j)}
\]

   if \texttt{TargetSelection=}\texttt{Mean}. 

---
Kernel functions for regression

The kernel weighting methods have been implemented also for regression, taking into account the modified structure of the foam in case of multi-target regression.

8.4.5 Visualisation of the foam via projections to 2 dimensions

A projection function is used to visualise the foam in two dimensions. It is called via:

```c
TH2D *proj = foam->Project2( dim1, dim2, cell_value, kernel, nbin );
```

Code Example 48: Call of the projection function. The first two arguments are the dimensions one wishes to project on, the third specifies the quantity to plot (kValue, kValueError, kValueDensity, kMeanValue, kRms, kRmsOvMean, kCellVolume), and the fourth argument chooses the kernel (default value is NULL which means that no kernel is used). By the optional parameter nbin one can set the number of bins of the resulting histogram (default value is 50).

For each bin in the histogram the algorithm finds all cells which match the bin coordinate and fills the cell values, depending on cell_value, into the bin. This implies that in the case of more than two dimensions the cell values in the dimensions that are not visible are summed.

- **cell_value==kValue – projecting the cell value**
  
  When this option is given, the standard cell values are visualized. The concrete values depend on the foam type.

  In case of classification with separate signal and background foams or multi-target regression, the cell value is the number of events found in the phase space region of the cell.

  In the single foam approach (SigBgSeparate==False) the foam cells are filled with the discriminator. The value \( v(i) \), filled in the histogram is equal to the discriminator \( y(i) \) stored in cell \( i \), multiplied by the cell volume, excluding the dimensions \( \text{dim1} \) and \( \text{dim2} \)

  \[
  v(i) = y(i) \prod_{k=1 \atop k \neq \text{dim1}, k \neq \text{dim2}}^{d} L(i,k) , \tag{60}
  \]

  where \( L(i,k) \) is the length of the foam cell \( i \) in the direction \( k \) of a \( d \)-dimensional foam. This means that the average discriminator weighted with the cell size of the non-visualised dimensions is filled.

  In case of a mono-target regression foam (MultiTargetRegression==False) the foam cells are filled with the target, which is then directly filled into the histogram.

- **cell_value==kRms, cell_value==kRmsOvMean – projection of cell variances**
  
  The variance (RMS) and the mean of the event distribution are saved in every foam cell,
independent of the foam type. If cell_value==kRms, then the plotted value is the RMS of the event distribution in the cell. If cell_value==kRmsOvMean, then the RMS divided by the mean of the event distribution in the cell is filled into the histogram.

- **kernel** – *Using kernels for projecting*
  Instead of filling the cell values directly into the histogram, one can use a PDEFoam kernel estimator to interpolate between cell values. In this case the cell value (set by the cell_value option) is passed to the kernel estimator, whose output is filled into the histogram. See page 87 for more details on kernels available for PDE-Foam.

### 8.4.6 Variable ranking

In PDE-Foam the input variables are ranked according to the number of cuts made in each PDE-Foam dimension. The dimension (the input variable) for which the most cuts were done is ranked highest.

### 8.4.7 Performance

Like PDE-RS (see Sec. 8.3), this method is a powerful classification tool for problems with highly non-linearly correlated observables. Furthermore PDE-Foam is a fast responding classifier, because of its limited number of cells, independent of the size of the training samples.

An exception is the multi-target regression with Gauss kernel because the time scales with the number of cells squared. Also the training can be slow, depending on the number of training events and number of cells one wishes to create.

### 8.5 k-Nearest Neighbour (k-NN) Classifier

Similar to PDE-RS (cf. Sec. 8.3), the k-nearest neighbour method compares an observed (test) event to reference events from a training data set. However, unlike PDE-RS, which in its original form uses a fixed-sized multidimensional volume surrounding the test event, and in its augmented form resizes the volume as a function of the local data density, the k-NN algorithm is intrinsically adaptive. It searches for a fixed number of adjacent events, which then define a volume for the metric used. The k-NN classifier has best performance when the boundary that separates signal and background events has irregular features that cannot be easily approximated by parametric learning methods.

#### 8.5.1 Booking options

The k-NN classifier is booked via the command:

---

26 Variable ranking is available for PDE-Foam since TMVA version 4.1.1.
### Option Table 14: Configuration options reference for MVA method: k-NN

Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured.

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>nkNN</td>
<td>–</td>
<td>20</td>
<td>–</td>
<td>Number of k-nearest neighbors</td>
</tr>
<tr>
<td>BalanceDepth</td>
<td>–</td>
<td>6</td>
<td>–</td>
<td>Binary tree balance depth</td>
</tr>
<tr>
<td>ScaleFrac</td>
<td>–</td>
<td>0.8</td>
<td>–</td>
<td>Fraction of events used to compute variable width</td>
</tr>
<tr>
<td>SigmaFact</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Scale factor for sigma in Gaussian kernel</td>
</tr>
<tr>
<td>Kernel</td>
<td>–</td>
<td>Gaus</td>
<td>–</td>
<td>Use polynomial (=Poln) or Gaussian (=Gaus) kernel</td>
</tr>
<tr>
<td>Trim</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Use equal number of signal and background events</td>
</tr>
<tr>
<td>UseKernel</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Use polynomial kernel weight</td>
</tr>
<tr>
<td>UseWeight</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Use weight to count kNN events</td>
</tr>
<tr>
<td>UseLDA</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Use local linear discriminant - experimental feature</td>
</tr>
</tbody>
</table>

The configuration options for the k-NN classifier are listed in Option Table 14 (see also Sec. 6).

#### 8.5.2 Description and implementation

The k-NN algorithm searches for $k$ events that are closest to the test event. Closeness is thereby measured using a metric function. The simplest metric choice is the Euclidean distance

$$R = \left( \sum_{i=1}^{n_{\text{var}}} |x_i - y_i|^2 \right)^{\frac{1}{2}}.$$  \hspace{1cm} (61)

where $n_{\text{var}}$ is the number of input variables used for the classification, $x_i$ are coordinates of an event from a training sample and $y_i$ are variables of an observed test event. The $k$ events with the smallest values of $R$ are the *k-nearest neighbours*. The value of $k$ determines the size of the neighbourhood for which a probability density function is evaluated. Large values of $k$ do not capture the local behavior of the probability density function. On the other hand, small values of $k$ cause statistical
fluctuations in the probability density estimate. A case study with real data suggests that values of \( k \) between 10 and 100 are appropriate and result in similar classification performance when the training sample contains hundreds of thousands of events (and \( n_{\text{var}} \) is of the order of a few variables).

The classification algorithm finds \( k \)-nearest training events around a query point

\[
k = k_S + k_B,
\]

where \( k_S(B) \) is number of the signal (background) events in the training sample. The relative probability that the test event is of signal type is given by

\[
P_S = \frac{k_S}{k_S + k_B} = \frac{k_S}{k}.
\]

The choice of the metric governs the performance of the nearest neighbour algorithm. When input variables have different units a variable that has a wider distribution contributes with a greater weight to the Euclidean metric. This feature is compensated by rescaling the variables using a scaling fraction determined by the option ScaleFrac. Rescaling can be turned off by setting ScaleFrac to 0. The scaling factor applied to variable \( i \) is determined by the width \( w_i \) of the \( x_i \) distribution for the combined sample of signal and background events: \( w_i \) is the interval that contains the fraction ScaleFrac of \( x_i \) training values. The input variables are then rescaled by \( 1/w_i \), leading to the rescaled metric

\[
R_{\text{rescaled}} = \left( \sum_{i=1}^{d} \frac{1}{w_i^2} |x_i - y_i|^2 \right)^{\frac{1}{2}}.
\]

Figure 17 shows an example of event classification with the \( k \)-nearest neighbour algorithm.\(^{27}\)

---

\(^{27}\) The number of training events shown has been greatly reduced to illustrate the principle of the algorithm. In a real application a typical \( k \)-NN training sample should be ample.
The output of the k-nearest neighbour algorithm can be interpreted as a probability that an event is of signal type, if the numbers (better: sum of event weights) of signal and background events in the training sample are equal. This can be enforced via the `Trim` option. If set training events of the overabundant type are randomly removed until parity is achieved.

Like (more or less) all TMVA classifiers, the k-nearest neighbour estimate suffers from statistical fluctuations in the training data. The typically high variance of the k-NN response is mitigated by adding a weight function that depends smoothly on the distance from a test event. The current k-NN implementation uses a polynomial kernel

\[ W(x) = \begin{cases} 
(1 - |x|^3)^3 & \text{if } |x| < 1 , \\
0 & \text{otherwise}.
\end{cases} \tag{65} \]

If \( R_k \) is the distance between the test event and the \( k \)th neighbour, the events are weighted according to the formula:

\[ W_{S(B)} = \sum_{i=1}^{k_{S(B)}} W \left( \frac{R_i}{R_k} \right) , \tag{66} \]

where \( k_{S(B)} \) is number of the signal (background) events in the neighbourhood. The weighted signal probability for the test event is then given by

\[ P_S = \frac{W_S}{W_S + W_B} . \tag{67} \]

The kernel use is switched on/off by the option `UseKernel`.

**Regression**

The k-NN algorithm in TMVA also implements a simple multi-dimensional (multi-target) regression model. For a test event, the algorithm finds the k-nearest neighbours using the input variables, where each training event contains a regression value. The predicted regression value for the test event is the weighted average of the regression values of the k-nearest neighbours, cf. Eq. (45) on page 81.

**8.5.3 Ranking**

The present implementation of k-NN does not provide a ranking of the input variables.

**8.5.4 Performance**

The simplest implementation of the k-NN algorithm would store all training events in an array. The classification would then be performed by looping over all stored events and finding the k-nearest neighbours. As discussed in Sec. 4.3, such an implementation is impractical for large training samples. The k-NN algorithm therefore uses a `kd-tree` structure [12] that significantly improves the performance.
The TMVA implementation of the k-NN method is reasonably fast to allow classification of large data sets. In particular, it is faster than the adaptive PDE-RS method (cf. Sec. 8.3). Note that the k-NN method is not appropriate for problems where the number of input variables exceeds \( n_{\text{var}} \gtrsim 10 \). The neighbourood size \( R \) depends on \( n_{\text{var}} \) and the size of the training sample \( N \) as

\[
R_N \propto \frac{1}{n_{\text{var}} \sqrt{N}}.
\]

A large training set allows the algorithm to probe small-scale features that distinguish signal and background events.

8.6 H-Matrix discriminant

The origins of the H-Matrix approach dates back to works of Fisher and Mahalanobis in the context of Gaussian classifiers [22, 23]. It discriminates one class (signal) of a feature vector from another (background). The correlated elements of the vector are assumed to be Gaussian distributed, and the inverse of the covariance matrix is the H-Matrix. A multivariate \( \chi^2 \) estimator is built that exploits differences in the mean values of the vector elements between the two classes for the purpose of discrimination.

The H-Matrix classifier as it is implemented in TMVA is equal or less performing than the Fisher discriminant (see Sec. 8.7), and has been only included for completeness.

8.6.1 Booking options

The H-Matrix discriminant is booked via the command:

```
factory->BookMethod( Types::kHMatrix, "HMatrix", "<options>" );
```

**Code Example 50:** Booking of the H-Matrix classifier: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ':'. See Sec. 3.1.5 for more information on the booking.

No specific options are defined for this method beyond those shared with all the other methods (cf. Option Table 9 on page 71).

8.6.2 Description and implementation

For an event \( i \), each one \( \chi^2 \) estimator \( (\chi^2_{S(B)}) \) is computed for signal \( (S) \) and background \( (B) \), using estimates for the sample means \( (\overline{x}_{S(B),i}) \) and covariance matrices \( (C_{S(B)}) \) obtained from the
training data

$$\chi^2_U(i) = \sum_{k,\ell=1}^{n_{\text{var}}} (x_k(i) - \bar{x}_{U,k}) C^{-1}_{U,\ell} (x_{\ell}(i) - \bar{x}_{U,\ell}) ,$$

where $U = S, B$. From this, the discriminant

$$y_H(i) = \frac{\chi^2_B(i) - \chi^2_S(i)}{\chi^2_B(i) + \chi^2_S(i)} ,$$

is computed to discriminate between the signal and background classes.

### 8.6.3 Variable ranking

The present implementation of the H-Matrix discriminant does not provide a ranking of the input variables.

### 8.6.4 Performance

The TMVA implementation of the H-Matrix classifier has been shown to underperform in comparison with the corresponding Fisher discriminant (cf. Sec. 8.7), when using similar assumptions and complexity. It might therefore be considered to be depreciated.

### 8.7 Fisher discriminants (linear discriminant analysis)

In the method of Fisher discriminants [22] event selection is performed in a transformed variable space with zero linear correlations, by distinguishing the mean values of the signal and background distributions. The linear discriminant analysis determines an axis in the (correlated) hyperspace of the input variables such that, when projecting the output classes (signal and background) upon this axis, they are pushed as far as possible away from each other, while events of a same class are confined in a close vicinity. The linearity property of this classifier is reflected in the metric with which ”far apart” and ”close vicinity” are determined: the covariance matrix of the discriminating variable space.

#### 8.7.1 Booking options

The Fisher discriminant is booked via the command:

```cpp
factory->BookMethod( Types::kFisher, "Fisher", "<options>" );
```

**Code Example 51:** Booking of the Fisher discriminant: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ’:’. See Sec. 3.1.5 for more information on the booking.
### 8.7 Fisher discriminants (linear discriminant analysis)

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method</td>
<td>–</td>
<td>Fisher</td>
<td>Fisher, Mahalanobis</td>
<td>Discrimination method</td>
</tr>
</tbody>
</table>

**Option Table 15**: Configuration options reference for MVA method: *Fisher*. Values given are defaults. If predefined categories exist, the default category is marked by a ‘*star*’. The options in Option Table refopt:mva::methodbase on page pagerefopt:mva::methodbase can also be configured.

The configuration options for the Fisher discriminant are given in Option Table 15.

#### 8.7.2 Description and implementation

The classification of the events in signal and background classes relies on the following characteristics: the overall sample means $\bar{x}_k$ for each input variable $k = 1, \ldots, n_{\text{var}}$, the class-specific sample means $\bar{x}_{S(B),k}$, and total covariance matrix $C$ of the sample. The covariance matrix can be decomposed into the sum of a *within-* ($W$) and a *between-class matrix* ($B$). They respectively describe the dispersion of events relative to the means of their own class (within-class matrix), and relative to the overall sample means (between-class matrix)\(^28\).

The *Fisher coefficients*, $F_k$, are then given by

$$F_k = \sqrt{\frac{N_S N_B}{N_S + N_B}} \sum_{\ell=1}^{n_{\text{var}}} W_{k\ell}^{-1} (\bar{x}_{S,\ell} - \bar{x}_{B,\ell}) , \quad (71)$$

where $N_{S(B)}$ are the number of signal (background) events in the training sample. The Fisher discriminant $y_{Fi}(i)$ for event $i$ is given by

$$y_{Fi}(i) = F_0 + \sum_{k=1}^{n_{\text{var}}} F_k x_k(i) . \quad (72)$$

The offset $F_0$ centers the sample mean $\bar{y}_{Fi}$ of all $N_S + N_B$ events at zero.

Instead of using the within-class matrix, the Mahalanobis variant determines the Fisher coefficients

\(^28\)The within-class matrix is given by

$$W_{k\ell} = \sum_{U=S,B} (x_{U,k} - \bar{x}_{U,k}) (x_{U,\ell} - \bar{x}_{U,\ell}) = C_{S,k\ell} + C_{B,k\ell} ,$$

where $C_{S(B)}$ is the covariance matrix of the signal (background) sample. The between-class matrix is obtained by

$$B_{k\ell} = \frac{1}{2} \sum_{U=S,B} (\bar{x}_{U,k} - \bar{x}_k) (\bar{x}_{U,\ell} - \bar{x}_\ell) ,$$

where $\bar{x}_{S(B),k}$ is the average of variable $x_k$ for the signal (background) sample, and $\bar{x}_k$ denotes the average for the entire sample.
The Fisher discriminant analysis aims at simultaneously maximising the between-class separation while minimising the within-class dispersion. A useful measure of the discrimination power of a variable is therefore given by the diagonal quantity $B_{kk}/C_{kk}$, which is used for the ranking of the input variables.

### 8.7.4 Performance

In spite of the simplicity of the classifier, Fisher discriminants can be competitive with likelihood and nonlinear discriminants in certain cases. In particular, Fisher discriminants are optimal for Gaussian distributed variables with linear correlations (cf. the standard toy example that comes with TMVA).

On the other hand, no discrimination at all is achieved when a variable has the same sample mean for signal and background, even if the shapes of the distributions are very different. Thus, Fisher discriminants often benefit from suitable transformations of the input variables. For example, if a variable $x \in [-1, 1]$ has a a signal distributions of the form $x^2$, and a uniform background distributions, their mean value is zero in both cases, leading to no separation. The simple transformation $x \to |x|$ renders this variable powerful for the use in a Fisher discriminant.

### 8.8 Linear discriminant analysis (LD)

The linear discriminant analysis provides data classification using a linear model, where $\textit{linear}$ refers to the discriminant function $y(x)$ being linear in the parameters $\beta$

$$y(x) = x^T \beta + \beta_0 \, ,$$

where $\beta_0$ (denoted the $\textit{bias}$) is adjusted so that $y(x) \geq 0$ for signal and $y(x) < 0$ for background. It can be shown that this is equivalent to the Fisher discriminant, which seeks to maximise the ratio of between-class variance to within-class variance by projecting the data onto a linear subspace.

### 8.8.1 Booking options

The LD is booked via the command:
8.8 Linear discriminant analysis (LD)

Code Example 52: Booking of the linear discriminant: the first argument is a predefined enumerator, the second argument is a user-defined string identifier. No method-specific options are available. See Sec. 3.1.5 for more information on the booking.

No specific options are defined for this method beyond those shared with all the other methods (cf. Option Table 9 on page 71).

8.8.2 Description and implementation

Assuming that there are $m+1$ parameters $\beta_0, \cdots, \beta_m$ to be estimated using a training set comprised of $n$ events, the defining equation for $\beta$ is

$$ Y = X \beta, $$

where we have absorbed $\beta_0$ into the vector $\beta$ and introduced the matrices

$$ Y = \begin{pmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{pmatrix} \quad \text{and} \quad X = \begin{pmatrix} 1 & x_{11} & \cdots & x_{1m} \\ 1 & x_{21} & \cdots & x_{2m} \\ \vdots & \vdots & \ddots & \vdots \\ 1 & x_{n1} & \cdots & x_{nm} \end{pmatrix}, $$

where the constant column in $X$ represents the bias $\beta_0$ and $Y$ is composed of the target values with $y_i = 1$ if the $i$th event belongs to the signal class and $y_i = 0$ if the $i$th event belongs to the background class. Applying the method of least squares, we now obtain the normal equations for the classification problem, given by

$$ X^T X \beta = X^T Y \iff \beta = (X^T X)^{-1} X^T Y. $$

The transformation $(X^T X)^{-1} X^T$ is known as the Moore-Penrose pseudo inverse of $X$ and can be regarded as a generalisation of the matrix inverse to non-square matrices. It requires that the matrix $X$ has full rank.

If weighted events are used, this is simply taken into account by introducing a diagonal weight matrix $W$ and modifying the normal equations as follows:

$$ \beta = (X^T W X)^{-1} X^T W Y. $$

Considering two events $x_1$ and $x_2$ on the decision boundary, we have $y(x_1) = y(x_2) = 0$ and hence $(x_1 - x_2)^T \beta = 0$. Thus we see that the LD can be geometrically interpreted as determining the decision boundary by finding an orthogonal vector $\beta$. 
8.8.3 Variable ranking

The present implementation of LD provides a ranking of the input variables based on the coefficients of the variables in the linear combination that forms the decision boundary. The order of importance of the discriminating variables is assumed to agree with the order of the absolute values of the coefficients.

8.8.4 Regression with LD

It is straightforward to apply the LD algorithm to linear regression by replacing the binary targets \( y_i \in 0, 1 \) in the training data with the measured values of the function which is to be estimated. The resulting function \( y(x) \) is then the best estimate for the data obtained by least-squares regression.

8.8.5 Performance

The LD is optimal for Gaussian distributed variables with linear correlations (cf. the standard toy example that comes with TMVA) and can be competitive with likelihood and nonlinear discriminants in certain cases. No discrimination is achieved when a variable has the same sample mean for signal and background, but the LD can often benefit from suitable transformations of the input variables. For example, if a variable \( x \in [-1, 1] \) has a signal distribution of the form \( x^2 \) and a uniform background distribution, their mean value is zero in both cases, leading to no separation. The simple transformation \( x \rightarrow |x| \) renders this variable powerful for the use with LD.

8.9 Function discriminant analysis (FDA)

The common goal of all TMVA discriminators is to determine an optimal separating function in the multivariate space of all input variables. The Fisher discriminant solves this analytically for the linear case, while artificial neural networks, support vector machines or boosted decision trees provide nonlinear approximations with – in principle – arbitrary precision if enough training statistics is available and the chosen architecture is flexible enough.

The function discriminant analysis (FDA) provides an intermediate solution to the problem with the aim to solve relatively simple or partially nonlinear problems. The user provides the desired function with adjustable parameters via the configuration option string, and FDA fits the parameters to it, requiring the function value to be as close as possible to the real value (to 1 for signal and 0 for background in classification). Its advantage over the more involved and automatic nonlinear discriminators is the simplicity and transparency of the discrimination expression. A shortcoming is that FDA will underperform for involved problems with complicated, phase space dependent nonlinear correlations.
8.9 Function discriminant analysis (FDA)

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Formula</td>
<td>–</td>
<td>(0)</td>
<td>–</td>
<td>The discrimination formula</td>
</tr>
<tr>
<td>ParRanges</td>
<td>–</td>
<td>()</td>
<td>–</td>
<td>Parameter ranges</td>
</tr>
<tr>
<td>FitMethod</td>
<td>–</td>
<td>MINUIT</td>
<td>MC, GA, SA, MINUIT</td>
<td>Optimisation Method</td>
</tr>
<tr>
<td>Converger</td>
<td>–</td>
<td>None</td>
<td>None, MINUIT</td>
<td>FitMethod uses Converger to improve result</td>
</tr>
</tbody>
</table>

**Option Table 16**: Configuration options reference for MVA method: FDA. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured. The input variables in the discriminator expression are denoted \(x_0, x_1, \ldots\) (until \(n_{\text{var}} - 1\)), where the number follows the order in which the variables have been registered with the Factory; coefficients to be determined by the fit must be denoted \((0), (1), \ldots\) (the number of coefficients is free) in the formula; allowed is any functional expression that can be interpreted by a ROOT TFormula. See Code Example 54 for an example expression. The limits for the fit parameters (coefficients) defined in the formula expression are given with the syntax: "\((-1, 3); (2, 10); \ldots\)”, where the first interval corresponds to parameter \((0)\). The converger allows to use (presently only) Minuit fitting in addition to Monte Carlo sampling or a Genetic Algorithm. More details on this combination are given in Sec. 6.5. The various fitters are configured using the options given in Tables 5, 6, 7 and 8, for MC, Minuit, GA and SA, respectively.

**8.9.1 Booking options**

FDA is booked via the command:

```cpp
factory->BookMethod( Types::kFDA, "FDA", "<options>" );
```

**Code Example 53**: Booking of the FDA classifier: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ‘:’. See Sec. 3.1.5 for more information on the booking.

The configuration options for the FDA classifier are listed in Option Table 16 (see also Sec. 6). A typical option string could look as follows:
"Formula=(0)+(1)*x0+(2)*x1+(3)*x2+(4)*x3:\nParRanges=(-1,1);(-10,10);(-10,10);(-10,10);(-10,10):\nFitMethod=MINUIT:\nErrorLevel=1:PrintLevel=-1:FitStrategy=2:UseImprove:UseMinos:SetBatch"

Code Example 54: FDA booking option example simulating a linear Fisher discriminant (cf. Sec. 8.7). The top line gives the discriminator expression, where the $x_i$ denote the input variables and the $(j)$ denote the coefficients to be determined by the fit. Allowed are all standard functions and expressions, including the functions belonging to the ROOT TMath library. The second line determines the limits for the fit parameters, where the numbers of intervals given must correspond to the number of fit parameters defined. The third line defines the fitter to be used (here Minuit), and the last line is the fitter configuration.

8.9.2 Description and implementation

The parsing of the discriminator function employs ROOT’s TFormula class, which requires that the expression complies with its rules (which are the same as those that apply for the TTree::Draw command). For simple formula with a single global fit solution, Minuit will be the most efficient fitter. However, if the problem is complicated, highly nonlinear, and/or has a non-unique solution space, more involved fitting algorithms may be required. In that case the Genetic Algorithm combined or not with a Minuit converger should lead to the best results. After fit convergence, FDA prints the fit results (parameters and estimator value) as well as the discriminator expression used on standard output. The smaller the estimator value, the better the solution found. The normalised estimator is given by

$$\text{For classification: } \mathcal{E} = \frac{1}{W_S} \sum_{i=1}^{N_S} (F(x_i) - 1)^2 w_i + \frac{1}{W_B} \sum_{i=1}^{N_B} F^2(x_i) w_i,$$

$$\text{For regression: } \mathcal{E} = \frac{1}{W} \sum_{i=1}^{N} (F(x_i) - t_i)^2 w_i,$$

(79)

where for classification the first (second) sum is over the signal (background) training events, and for regression it is over all training events, $F(x_i)$ is the discriminator function, $x_i$ is the tuple of the $n_{var}$ input variables for event $i$, $w_i$ is the event weight, $t_i$ the tuple of training regression targets, $W_{S(B)}$ is the sum of all signal (background) weights in the training sample, and $W$ the sum over all training weights.

8.9.3 Variable ranking

The present implementation of FDA does not provide a ranking of the input variables.

8.9.4 Performance

The FDA performance depends on the complexity and fidelity of the user-defined discriminator function. As a general rule, it should be able to reproduce the discrimination power of any linear
Artificial Neural Networks (nonlinear discriminant analysis). To reach into the nonlinear domain, it is useful to inspect the correlation profiles of the input variables, and add quadratic and higher polynomial terms between variables as necessary. Comparison with more involved nonlinear classifiers can be used as a guide.

8.10 Artificial Neural Networks (nonlinear discriminant analysis)

An Artificial Neural Network (ANN) is most generally speaking any simulated collection of interconnected neurons, with each neuron producing a certain response at a given set of input signals. By applying an external signal to some (input) neurons the network is put into a defined state that can be measured from the response of one or several (output) neurons. One can therefore view the neural network as a mapping from a space of input variables $x_1, \ldots, x_{nvar}$ onto a one-dimensional (e.g. in case of a signal-versus-background discrimination problem) or multi-dimensional space of output variables $y_1, \ldots, y_{mvar}$. The mapping is nonlinear if at least one neuron has a nonlinear response to its input.

In TMVA four neural network implementations are available to the user. The first was adapted from a FORTRAN code developed at the Université Blaise Pascal in Clermont-Ferrand,\textsuperscript{29} the second is the ANN implementation that comes with ROOT. The third is a newly developed neural network (denoted MLP) that is faster and more flexible than the other two and is the recommended neural network to use with TMVA. The fourth implementation is a highly optimized one which provides functionality to perform the training on multi-core and GPU architectures (see 8.11.1). This implementation has been developed specifically for the training of very complex neural networks with several hidden layers, so called deep neural networks. All four neural networks are feed-forward multilayer perceptrons.

**The Clermont-Ferrand neural network**

The Clermont-Ferrand neural network is booked via the command:

```cpp
factory->BookMethod( Types::kCFMlpANN, "CF_ANN", "<options>" );
```

**Code Example 55:** Booking of the Clermont-Ferrand neural network: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the options string. Individual options are separated by a `:`. See Sec. 3.1.5 for more information on the booking.

The configuration options for the Clermont-Ferrand neural net are given in Option Table 17. Since CFMlpANN is less powerful than the other neural networks in TMVA, it is only kept for backward compatibility and we do not recommend to use it when starting a new analysis.

\textsuperscript{29}The original Clermont-Ferrand neural network has been used for Higgs search analyses in ALEPH, and background fighting in rare $B$-decay searches by the BABAR Collaboration. For the use in TMVA the FORTRAN code has been converted to C++.
Option Table 17: Configuration options reference for MVA method: \texttt{CFMlpANN}. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured. See Sec. 8.10.2 for a description of the network architecture configuration.

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCycles</td>
<td>–</td>
<td>3000</td>
<td>–</td>
<td>Number of training cycles</td>
</tr>
</tbody>
</table>
| HiddenLayers | –    | N,N-1   | –                 | Specification of hidden layer architec-

| ValidationFraction | – | 0.5 | – | Fraction of events in training tree used for cross validation |
| LearningMethod     | – | Stochastic | Stochastic, Batch, SteepestDescent, RibierePolak, FletcherReeves, BFGS | Learning method |

Option Table 18: Configuration options reference for MVA method: \texttt{TMlpANN}. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured. See Sec. 8.10.2 for a description of the network architecture configuration.

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCycles</td>
<td>–</td>
<td>200</td>
<td>–</td>
<td>Number of training cycles</td>
</tr>
</tbody>
</table>
| HiddenLayers | –    | N,N-1   | –                 | Specification of hidden layer architec-

| ValidationFraction | – | 0.5 | – | Fraction of events in training tree used for cross validation |
| LearningMethod     | – | Stochastic | Stochastic, Batch, SteepestDescent, RibierePolak, FletcherReeves, BFGS | Learning method |

The ROOT neural network (class \texttt{TMultiLayerPerceptron})

This neural network interfaces the ROOT class \texttt{TMultiLayerPerceptron} and is booked through the Factory via the command line:

```
factory->BookMethod( Types::kTMlpANN, "TMlp_ANN", "<options>" );
```

Code Example 56: Booking of the ROOT neural network: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. See Sec. 3.1.5 for more information on the booking.

The configuration options for the ROOT neural net are given in Option Table 18.
The MLP neural network

The MLP neural network is booked through the Factory via the command line:

```cpp
factory->BookMethod( Types::kMLP, "MLP_ANN", "<options>" );
```

**Code Example 57**: Booking of the MLP neural network: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the options string. See Sec. 3.1.5 for more information on the booking.

The configuration options for the MLP neural net are given in Option Tables 19 and 20. The option `UseRegulator` is of special interest since it invokes the Bayesian extension of the MLP, which is described in Section .

The TMVA implementation of MLP supports random and importance event sampling. With event sampling enabled, only a fraction (set by the option `Sampling`) of the training events is used for the training of the MLP. Values in the interval $[0, 1]$ are possible. If the option `SamplingImportance` is set to 1, the events are selected randomly, while for a value below 1 the probability for the same events to be sampled again depends on the training performance achieved for classification or regression. If for a given set of events the training leads to a decrease of the error of the test sample, the probability for the events of being selected again is multiplied with the factor given in `SamplingImportance` and thus decreases. In the case of an increased error of the test sample, the probability for the events to be selected again is divided by the factor `SamplingImportance` and thus increases. The probability for an event to be selected is constrained to the interval $[0, 1]$. For each set of events, the importance sampling described above is performed together with the overtraining test.

Event sampling is performed until the fraction specified by the option `SamplingEpoch` of the total number of epochs (`NCycles`) has been reached. Afterwards, all available training events are used for the training. Event sampling can be turned on and off for training and testing events individually with the options `SamplingTraining` and `SamplingTesting`.

The aim of random and importance sampling is foremost to speed-up the training for large training samples. As a side effect, random or importance sampling may also increase the robustness of the training algorithm with respect to convergence in a local minimum.

Since it is typically not known beforehand how many epochs are necessary to achieve a sufficiently good training of the neural network, a convergence test can be activated by setting `ConvergenceTests` to a value above 0. This value denotes the number of subsequent convergence tests which have to fail (i.e. no improvement of the estimator larger than `ConvergenceImprove`) to consider the training to be complete. Convergence tests are performed at the same time as overtraining tests. The test frequency is given by the parameter `TestRate`.

It is recommended to set the option `VarTransform=Norm`, such that the input (and in case of regression the output as well) is normalised to the interval $[-1, 1]$. 
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCycles</td>
<td>–</td>
<td>500</td>
<td>–</td>
<td>Number of training cycles</td>
</tr>
<tr>
<td>HiddenLayers</td>
<td>–</td>
<td>N,N-1</td>
<td>–</td>
<td>Specification of hidden layer architecture</td>
</tr>
<tr>
<td>NeuronType</td>
<td>–</td>
<td>sigmoid</td>
<td>–</td>
<td>Neuron activation function type</td>
</tr>
<tr>
<td>RandomSeed</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Random seed for initial synapse weights (0 means unique seed for each run; default value '1')</td>
</tr>
<tr>
<td>EstimatorType</td>
<td>–</td>
<td>MSE</td>
<td>MSE, CE, linear, sigmoid, tanh, radial</td>
<td>MSE (Mean Square Estimator) for Gaussian Likelihood or CE(Cross-Entropy) for Bernoulli Likelihood</td>
</tr>
<tr>
<td>NeuronInputType</td>
<td>–</td>
<td>sum</td>
<td>sum, sqsum, abssum</td>
<td>Neuron input function type</td>
</tr>
<tr>
<td>TrainingMethod</td>
<td>–</td>
<td>BP</td>
<td>BP, GA, BFGS</td>
<td>Train with Back-Propagation (BP), BFGS Algorithm (BFGS), or Genetic Algorithm (GA - slower and worse)</td>
</tr>
<tr>
<td>LearningRate</td>
<td>–</td>
<td>0.02</td>
<td>–</td>
<td>ANN learning rate parameter</td>
</tr>
<tr>
<td>DecayRate</td>
<td>–</td>
<td>0.01</td>
<td>–</td>
<td>Decay rate for learning parameter</td>
</tr>
<tr>
<td>TestRate</td>
<td>–</td>
<td>10</td>
<td>–</td>
<td>Test for overtraining performed at each #th epochs</td>
</tr>
<tr>
<td>EpochMonitoring</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Provide epoch-wise monitoring plots according to TestRate (caution: causes big ROOT output file!)</td>
</tr>
<tr>
<td>Sampling</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Only 'Sampling' (randomly selected) events are trained each epoch</td>
</tr>
<tr>
<td>SamplingEpoch</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Sampling is used for the first 'SamplingEpoch' epochs, afterwards, all events are taken for training</td>
</tr>
<tr>
<td>SamplingImportance</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>The sampling weights of events in epochs which successful (worse estimator than before) are multiplied with SamplingImportance, else they are divided.</td>
</tr>
<tr>
<td>SamplingTraining</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>The training sample is sampled</td>
</tr>
<tr>
<td>SamplingTesting</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>The testing sample is sampled</td>
</tr>
<tr>
<td>ResetStep</td>
<td>–</td>
<td>50</td>
<td>–</td>
<td>How often BFGS should reset history</td>
</tr>
<tr>
<td>Tau</td>
<td>–</td>
<td>3</td>
<td>–</td>
<td>LineSearch size step</td>
</tr>
<tr>
<td>BPMode</td>
<td>–</td>
<td>sequential</td>
<td>sequential, batch</td>
<td>Back-propagation learning mode: sequential or batch</td>
</tr>
</tbody>
</table>

**Option Table 19:** Configuration options reference for MVA method: MLP. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured. See Sec. 8.10.2 for a description of the network architecture configuration. Continuation in Table 20.
8.10 Artificial Neural Networks (nonlinear discriminant analysis)

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BatchSize</td>
<td>–</td>
<td>-1</td>
<td>–</td>
<td>Batch size: number of events/batch, only set if in Batch Mode, -1 for Batch-Size=number_of_events</td>
</tr>
<tr>
<td>ConvergenceImprove</td>
<td>–</td>
<td>1e-30</td>
<td>–</td>
<td>Minimum improvement which counts as improvement (&lt;0 means automatic convergence check is turned off)</td>
</tr>
<tr>
<td>ConvergenceTests</td>
<td>–</td>
<td>-1</td>
<td>–</td>
<td>Number of steps (without improvement) required for convergence (&lt;0 means automatic convergence check is turned off)</td>
</tr>
<tr>
<td>UseRegulator</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Use regulator to avoid over-training</td>
</tr>
<tr>
<td>UpdateLimit</td>
<td>–</td>
<td>10000</td>
<td>–</td>
<td>Maximum times of regulator update</td>
</tr>
<tr>
<td>CalculateErrors</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Calculates inverse Hessian matrix at the end of the training to be able to calculate the uncertainties of an MVA value</td>
</tr>
<tr>
<td>WeightRange</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Take the events for the estimator calculations from small deviations from the desired value to large deviations only over the weight range</td>
</tr>
</tbody>
</table>

Option Table 20: Configuration options reference for MVA method: MLP. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured. See Sec. 8.10.2 for a description of the network architecture configuration. See also Table 19.

8.10.1 Description and implementation

The behaviour of an artificial neural network is determined by the layout of the neurons, the weights of the inter-neuron connections, and by the response of the neurons to the input, described by the neuron response function \( \rho \).

Multilayer Perceptron

While in principle a neural network with \( n \) neurons can have \( n^2 \) directional connections, the complexity can be reduced by organising the neurons in layers and only allowing direct connections from a given layer to the following layer (see Fig. 18). This kind of neural network is termed multi-layer perceptron; all neural net implementations in TMVA are of this type. The first layer of a multilayer perceptron is the input layer, the last one the output layer, and all others are hidden layers. For a classification problem with \( n_{\text{var}} \) input variables the input layer consists of \( n_{\text{var}} \) neurons that hold the input values, \( x_1, \ldots, x_{n_{\text{var}}} \), and one neuron in the output layer that holds the output variable, the neural net estimator \( y_{\text{ANN}} \).
For a regression problem the network structure is similar, except that for multi-target regression each of the targets is represented by one output neuron. A weight is associated to each directional connection between the output of one neuron and the input of another neuron. When calculating the input value to the response function of a neuron, the output values of all neurons connected to the given neuron are multiplied with theses weights.

**Neuron response function**

The neuron response function $\rho$ maps the neuron input $i_1, \ldots, i_n$ onto the neuron output (Fig. 19). Often it can be separated into a $\mathbb{R}^n \mapsto \mathbb{R}$ synapse function $\kappa$, and a $\mathbb{R} \mapsto \mathbb{R}$ neuron activation function $\sigma$. However, this separation is not always possible. In this document, it is simpler to use a single function $\rho$ which is the result of chaining $\kappa$ and $\sigma$. The equation for $\rho$ is given by:

$$\rho = \sigma(\kappa(i_1, \ldots, i_n))$$

Figure 18: Multilayer perceptron with one hidden layer.

Figure 19: Single neuron $j$ in layer $\ell$ with $n$ input connections. The incoming connections carry a weight of $w_{ij}^{(\ell-1)}$. 
function $\alpha$, so that $\rho = \alpha \circ \kappa$. The functions $\kappa$ and $\alpha$ can have the following forms:

$$
\kappa: (y_1^{(l)}, \ldots, y_n^{(l)} | w_{0j}^{(l)}, \ldots, w_{nj}^{(l)}) \rightarrow \begin{cases} 
  w_{0j}^{(l)} + \sum_{i=1}^{n} y_i^{(l)} w_{ij}^{(l)} & \text{Sum,} \\
  w_{0j}^{(l)} + \sum_{i=1}^{n} (y_i^{(l)} w_{ij}^{(l)})^2 & \text{Sum of squares,} \\
  w_{0j}^{(l)} + \sum_{i=1}^{n} |y_i^{(l)} w_{ij}^{(l)}| & \text{Sum of absolutes,}
\end{cases}
$$

(80)

$$
\alpha: x \rightarrow \begin{cases} 
  x & \text{Linear,} \\
  \frac{1}{1 + e^{-\kappa x}} & \text{Sigmoid,} \\
  \frac{e^{x} - e^{-x}}{e^{x} + e^{-x}} & \text{Tanh,} \\
  e^{-x^2/2} & \text{Radial.}
\end{cases}
$$

(81)

8.10.2 Network architecture

The number of hidden layers in a network and the number of neurons in these layers are configurable via the option HiddenLayers. For example the configuration "HiddenLayers=N-1,N+10,3" creates a network with three hidden layers, the first hidden layer with $n_{var} - 1$ neurons, the second with $n_{var} + 10$ neurons, and the third with 3 neurons.

When building a network two rules should be kept in mind. The first is the theorem by Weierstrass, which if applied to neural nets, ascertains that for a multilayer perceptron a single hidden layer is sufficient to approximate a given continuous correlation function to any precision, provided that a sufficiently large number of neurons is used in the hidden layer. If the available computing power and the size of the training data sample suffice, one can increase the number of neurons in the hidden layer until the optimal performance is reached.

It is likely that the same performance can be achieved with a network of more than one hidden layer and a potentially much smaller total number of hidden neurons. This would lead to a shorter training time and a more robust network.

8.10.3 Training of the neural network

Back-propagation (BP)

The most common algorithm for adjusting the weights that optimise the classification performance of a neural network is the so-called back propagation. It belongs to the family of supervised learning methods, where the desired output for every input event is known. Back propagation is used by all neural networks in TMVA. The output of a network (here for simplicity assumed to have a single hidden layer with a Tanh activation function, and a linear activation function in the output layer)
is given by
\[ y_{\text{ANN}}(2) = \sum_{j=1}^{n_h} y_j(2) \cdot w_{j1}^{(2)} = \sum_{j=1}^{n_h} \tanh \left( \sum_{i=1}^{n_{\text{var}}} x_i w_{ij}^{(1)} \right) \cdot w_{j1}^{(2)}, \]  
(82)

where \( n_{\text{var}} \) and \( n_h \) are the number of neurons in the input layer and in the hidden layer, respectively, \( w_{ij}^{(1)} \) is the weight between input-layer neuron \( i \) and hidden-layer neuron \( j \), and \( w_{j1}^{(2)} \) is the weight between the hidden-layer neuron \( j \) and the output neuron. A simple sum was used in Eq. (82) for the synapse function \( \kappa \).

During the learning process the network is supplied with \( N \) training events \( x_a = (x_1, \ldots, x_{n_{\text{var}}}) \), \( a = 1, \ldots, N \). For each training event \( a \) the neural network output \( y_{\text{ANN},a} \) is computed and compared to the desired output \( \hat{y}_a \in \{1, 0\} \) (in classification 1 for signal events and 0 for background events). An error function \( E \), measuring the agreement of the network response with the desired one, is defined by
\[ E(x_1, \ldots, x_N | w) = \sum_{a=1}^{N} E_a(x_a | w) = \sum_{a=1}^{N} \frac{1}{2} (y_{\text{ANN},a} - \hat{y}_a)^2, \]  
(83)

where \( w \) denotes the ensemble of adjustable weights in the network. The set of weights that minimises the error function can be found using the method of steepest or gradient descent, provided that the neuron response function is differentiable with respect to the input weights. Starting from a random set of weights \( w^{(\rho)} \) the weights are updated by moving a small distance in \( w \)-space into the direction \(- \nabla_w E\) where \( E \) decreases most rapidly
\[ w^{(\rho+1)} = w^{(\rho)} - \eta \nabla_w E, \]  
(84)

where the positive number \( \eta \) is the learning rate.

The weights connected with the output layer are updated by
\[ \Delta w_{j1}^{(2)} = -\eta \sum_{a=1}^{N} \frac{\partial E_a}{\partial w_{j1}^{(2)}} = -\eta \sum_{a=1}^{N} (y_{\text{ANN},a} - \hat{y}_a) y_j^{(2)} w_{j1}^{(2)} , \]  
(85)

and the weights connected with the hidden layers are updated by
\[ \Delta w_{ij}^{(1)} = -\eta \sum_{a=1}^{N} \frac{\partial E_a}{\partial w_{ij}^{(1)}} = -\eta \sum_{a=1}^{N} (y_{\text{ANN},a} - \hat{y}_a) y_j^{(2)} (1 - y_j^{(2)}) w_{j1}^{(2)} x_i,a , \]  
(86)

where we have used \( \tanh' x = \tanh x (1 - \tanh x) \). This method of training the network is denoted bulk learning, since the sum of errors of all training events is used to update the weights. An alternative choice is the so-called online learning, where the update of the weights occurs at each event. The weight updates are obtained from Eqs. (85) and (86) by removing the event summations. In this case it is important to use a well randomised training sample. Online learning is the learning method implemented in TMVA.
**BFGS**

The Broyden-Fletcher-Goldfarb-Shannon (BFGS) method \[24\] differs from back propagation by the use of second derivatives of the error function to adapt the synapse weight by an algorithm which is composed of four main steps.

1. Two vectors, \(D\) and \(Y\) are calculated. The vector of weight changes \(D\) represents the evolution between one iteration of the algorithm \((k-1)\) to the next \((k)\). Each synapse weight corresponds to one element of the vector. The vector \(Y\) is the vector of gradient errors.

\[
D^{(k)}_i = w^{(k)}_i - w^{(k-1)}_i, \quad (87)
\]
\[
Y^{(k)}_i = g^{(k)}_i - g^{(k-1)}_i, \quad (88)
\]

where \(i\) is the synapse index, \(g_i\) is the \(i\)-th synapse gradient, \(w_i\) is the weight of the \(i\)-th synapse, and \(k\) denotes the iteration counter.

2. Approximate the inverse of the Hessian matrix, \(H^{-1}\), at iteration \(k\) by

\[
H^{-1(k)} = \frac{D \cdot D^T \cdot (1 + Y^T \cdot H^{-1(k-1)} \cdot Y)}{Y^T \cdot D} - D \cdot Y^T \cdot H + H \cdot Y \cdot D^T + H^{-1(k-1)}, \quad (89)
\]

where superscripts \((k)\) are implicit for \(D\) and \(Y\).

3. Estimate the vector of weight changes by

\[
D^{(k)} = -H^{-1(k)} \cdot Y^{(k)}. \quad (90)
\]

4. Compute a new vector of weights by applying a line search algorithm. In the line search the error function is locally approximated by a parabola. The algorithm evaluates the second derivatives and determines the point where the minimum of the parabola is expected. The total error is evaluated for this point. The algorithm then evaluates points along the line defined by the direction of the gradient in weights space to find the absolute minimum. The weights at the minimum are used for the next iteration. The learning rate can be set with the option Tau. The learning parameter, which defines by how much the weights are changed in one epoch along the line where the minimum is suspected, is multiplied with the learning rate as long as the training error of the neural net with the changed weights is below the one with unchanged weights. If the training error of the changed neural net were already larger for the initial learning parameter, it is divided by the learning rate until the training error becomes smaller. The iterative and approximate calculation of \(H^{-1(k)}\) turns less accurate with an increasing number of iterations. The matrix is therefore reset to the unit matrix every ResetStep steps.

The advantage of the BFGS method compared to BG is the smaller number of iterations. However, because the computing time for one iteration is proportional to the squared number of synapses, large networks are particularly penalised.

\[30\] The synapse gradient is estimated in the same way as in the BP method (with initial gradient and weights set to zero).
8.10.4 Variable ranking

The MLP neural network implements a variable ranking that uses the sum of the weights-squared of the connections between the variable’s neuron in the input layer and the first hidden layer. The importance $I_i$ of the input variable $i$ is given by

$$I_i = \overline{x}_i^2 \sum_{j=1}^{n_h} (w_{ij}^{(1)})^2 , \quad i = 1, \ldots, n_{\text{var}} , \quad (91)$$

where $\overline{x}_i$ is the sample mean of input variable $i$.

8.10.5 Bayesian extension of the MLP

Achieving a good test performance with MLP is a tradeoff between using a network architecture that is flexible enough to allow for the modelling of complex functions and the avoidance of overtraining. Avoiding overtraining by limiting the flexibility of the network in the first place by keeping the number of hidden units/layers small is a very rigid approach which will typically hurt the performance of the neural network. The Bayesian extension of MLP offers a means to allow for more complex network architectures while at the same time regularizing the model complexity adaptively to avoid unwanted overfitting effects. The price to pay is an increase in computation time. The extension is enabled with the option UseRegulator and should typically be used in difficult setting where the problem calls for a network architecture with many hidden units and/or several layers. Enabling the Bayesian extension of the MLP will add an additional term to the network error function $E(w)$ (which is equivalent to minus the log-likelihood of the training data given the network model):

$$\tilde{E}(w) = E(w) + \alpha |w|^2 \quad (92)$$

The penalty term is proportional to the squared norm of the weight ensemble $w$ of the network and effectively penalizes large weights which results in a reduced model complexity. The metaparameter $\alpha$ controls the amount of regularization applied and thus ultimatvely controls the level of model complexity the network can exhibit. This is called a Bayesian extension since it is equivalent to introducing a Gaussian prior for the network weights, whose width is controlled by the parameter $\alpha$. Since choosing $\alpha$ manually would only shift the problem of picking the correct network complexity to picking the right $\alpha$, TMVA automatically infers the best value of $\alpha$ from the training data by doing a complexity analysis based on the so-called evidence approximation. The user can therefore choose to train a large network which will then automatically be regularized to a degree that optimally fits the data.

8.10.6 Performance

In the tests we have carried out so far, the MLP and ROOT networks performed equally well, however with a clear speed advantage for the MLP. The Clermont-Ferrand neural net exhibited worse classification performance in these tests, which is partly due to the slow convergence of its training (at least 10k training cycles are required to achieve approximately competitive results).
8.11 Deep Learning

The Deep Learning module in TMVA provides the support to build several deep learning architectures such as Deep Neural Networks (DNN), Convolutional networks (CNN) and Recurrent networks (RNN). The purpose of this module is to provide optimized implementations of these networks that can be efficiently trained on modern multi-core and GPU hardware architectures. For this, the implementation provides two backends for training and evaluating deep learning models.

The CPU backend uses multithreading to perform the training in parallel on multi-core CPU architectures and it can make use of efficient multithreaded BLAS implementations such as OpenBLAS and the Intel TBB library.

The GPU backend can be used to perform the training on CUDA-capable GPU architectures. In the case of Convolutional and Recurrent network models the implementation uses the low level CuDNN library of CUDA for optimal performances.

Building the CPU and GPU Backends To be able to use the multithreaded CPU backend, an implementation of the BLAS library as well as the Intel TBB library must be available on the system and detected by CMake. Note that the BLAS implementation must be multithreaded in order to exploit the multithreading capabilities of the underlying hardware. The Intel TBB library is used by the ROOT multi-threading libraries and it can be also part of ROOT build. Note that the IMT CMake flag must be set to enable multithreading in ROOT. If a BLAS implementation is not available, but ROOT is built with support for the MathMore library that is based on Gnu Scientific Library (GSL), the CBLAS implementation provided by GSL will be used. If neither BLAS or CBLAS are found, then the native ROOT matrix package will be used for the matrix multiplications needed to evaluate and train deep learning models. If ROOT has been built without IMT (multi-threading support) the execution will be serial instead of parallel.

For the GPU backend, a CUDA installation is required on the system and must be successfully detected by CMake. This requires the user to set the CUDA CMake flag when installing Root. Furthermore, for using CNN and RNN networks on GPU the CUDA cuDNN library must be installed and detected by CMAKE (CUDNN CMAKE flag).

If the user tries to use a backend that is not installed (e.g GPU), the training of the DNN method will be performed using the alternative one (e.g. CPU).

8.11.1 Deep Neural Networks

A deep neural network (DNN) is an artificial neural network (see Section 8.10) with several hidden layers and a large number of neurons in each layer. Recent developments in machine learning have shown that these networks are capable of learning complex, non-linear relations when trained on a sufficiently large amount of training data. For an article describing the application of DNNs to a classification problem in particle physics see the article by Baldi, Sadowski and Whiteson [48].
The deep neural network implementation provides an optimized implementation of feed-forward multilayer perceptrons that can be efficiently trained on modern multi-core and GPU architectures.

Note that previous ROOT/TMVA versions (version < 6.22) provide for Deep Neural Networks a method \texttt{TMVA::kDNN}, implementing only fully connected layers. Since ROOT version 6.18 it is recommended to use the new \texttt{TMVA::kDL} method, which supports not only dense layer as in the previous implementation, but also new type of layers as convolutional and recurrent, allowing to build convolutional and recurrent networks, as we will see in the next paragraphs.

8.11.2 Training of Deep Learning Models

As common for deep neural networks, this implementation uses several methods to train the network. As example we report here the formula for the \textit{stochastic batch gradient descend} (SGD). This means that in each training step the weights $W_{i,j}^k$ and bias terms $\theta_i^k$ of a given layer $k$ are updated using

$$W_{i,j}^k \rightarrow W_{i,j}^k - \alpha \frac{\partial J(x_b, y_b)}{\partial W_{i,j}^k}$$

$$\theta_i^k \rightarrow \theta_i^k - \alpha \frac{\partial J(x_b, y_b)}{\partial \theta_i^k}$$

Here $J(x_b, y_b)$ is the value of the loss function corresponding to the randomly chosen input batch $x_b$ and expected output $y_b$. If regularization is applied, the loss may as well contain contributions from the weight terms $W_{i,j}^k$ of each layer. This implementation also supports training with momentum. In this case training updates take the form:

$$\Delta W_{i,j}^k \rightarrow p \Delta W_{i,j}^k + (1.0 - p) \frac{\partial J(x_b, y_b)}{\partial W_{i,j}^k}$$

$$W_{i,j}^k \rightarrow W_{i,j}^k - \alpha \Delta W_{i,j}^k$$

$$\Delta \theta_i^k \rightarrow p \Delta \theta_i^k + (1.0 - p) \frac{\partial J(x_b, y_b)}{\partial \theta_i^k}$$

$$\theta_i^k \rightarrow \theta_i^k - \alpha \Delta \theta_i^k$$

Note that for $p = 0$ the standard stochastic batch gradient descent method is obtained.

Other optimizer algorithms such as ADAM, which is the default one since ROOT version 6.18, ADAGRAD, RMSPROP and ADADELTA are available as optimization algorithms, through the \texttt{Optimizer} option of the training strategy. For a detailed description of the various method see [49].

The training batch $(x_b, y_b)$ is chosen randomly and without replacement from the set of training samples. The number of training steps necessary to traverse the whole training set one time is referred to as an \textit{epoch}. 
The training of the deep neural network is performed in one or multiple training phases. A training phase ends when the test error did not decrease for a user-specified number of epochs.

**Validation and Evaluation Set**  The user should note that this implementation splits the TMVA training set in a sub-training set used to compute the gradient and update the weights and a validation set used to compute the error and to check for convergence. To properly evaluate the performance, the model is later evaluated on a completely separate data set, the TMVA test set.

### 8.11.3 Booking of the DL Method

The booking of the DL method is similar to any other method in TMVA. The method specific options are passed in the form of an option string \(<\text{options}>\) to the method.

```cpp
factory->BookMethod(dataloader, TMVA::Types::kDL, "DNN", <options>);
```

**Code Example 58:** Booking of a deep neural network: The first argument is a `TMVA::DataLoader` object, which holds training and test data. The second argument is the predefined enumerator object that represents the deep neural network implementation. The third argument is a string holding a user defined name for the method. The fourth argument is the option string holding the options for the method. See Sec. 3.1.5 for more information on the booking.

Through the option string, several configuration parameters can be specified. The complete set of options that can be set in the options string are summarized in Table 21. Some of the option such as the layout, define the models and the layers of the network and will be described in greater details later, when describing the different deep learning models that can be built.

The **Architecture** parameter is used to specify the backend used for the training. The multi-core CPU and GPU backends must be enabled during the installation of Root (CMAKE flags `tmva-cpu` and `tmva-gpu`).

The method used to initialize the weight matrices before beginning the training is specified using the **WeightInitialization** argument. Possible values are `XAVIER`, which initializes the weights with zero-mean Gaussian distributed random values, and `XAVIERUNIFORM`, which initializes the weights with uniformly distributed values.

### 8.11.4 Network Layout

The structure of the deep learning model is specified by the **Layout** parameter of the option string. This string specifies the type of layer (e.g. dense, convolutional, recurrent ), the activation function and some specific layer parameters such as the number of neurons of each layer of the network or the filter sizes for the convolutional layers. Different layers are separated by ' ' characters while
Option Table 21: Configuration options reference for MVA method: DL.

layer parameter such as activation function and number of neurons are separated by ‘|’ characters. Let’s see in detail how to build the layout string for the different models.

**Dense Layer Layout**  For example the layout string

```
Layout=TANH|128,TANH|128,TANH|128,LINEAR
```

defines a network with three hidden layers with 128 neurons and $\tanh$ activation functions. The neurons of the output layer are inferred automatically from the number of classes (in the case of classification) or the number of target variables (in the case of regression). The available activation functions are given in Table 22.

### 8.11.5 Convolutional Networks

Convolutional Neural Networks are a popular variation of the feed-forward networks described in the previous section. By making the explicit assumption that the input is an image, they manage to outperform conventional networks by significantly reducing the number of learnable parameters through a feature known as parameter sharing [50]. Instead of connecting every neuron of the current layer with every neuron of the previous layer as in conventional dense layers, we only learn the parameters for a set of a small kernels (typically 3x3 or 5x5) which slide over the input. The size of each kernel, as well as the number of pixels for each vertical or horizontal shift, are user defined meta-parameters. An important detail is that while the model assumes spatial invariance with respect to the height and width, no such assumption is made with respect to the mage depth. This asymmetry implies that at each receptive field, the kernel is connected with every slice of the...
### Option Table 22: DL Activation Functions

<table>
<thead>
<tr>
<th>Name</th>
<th>Activation Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>IDENTITY</td>
<td>( f(x) = x )</td>
</tr>
<tr>
<td>RELU</td>
<td>( f(x) = \begin{cases} x, &amp; \text{if } x \geq 0 \ 0, &amp; \text{otherwise} \end{cases} )</td>
</tr>
<tr>
<td>TANH</td>
<td>( f(x) = \text{tanh}(x) )</td>
</tr>
<tr>
<td>SIGMOID</td>
<td>( f(x) = \frac{1}{1+\exp(-x)} )</td>
</tr>
<tr>
<td>SOFTSIGN</td>
<td>( f(x) = \frac{x}{1+</td>
</tr>
<tr>
<td>GAUSS</td>
<td>( f(x) = \exp(-x^2) )</td>
</tr>
<tr>
<td>SYMMRELU</td>
<td>( f(x) = \begin{cases} x, &amp; \text{if } x \geq 0 \ -x, &amp; \text{otherwise} \end{cases} )</td>
</tr>
</tbody>
</table>

In order to reduce overfitting, we sometimes include a pooling layer between subsequent convolutional layers. This layer simply downsamples the input, typically by outputting the maximum value within its receptive field. This does not only reduce the number of neurons, but also helps reduce overfitting. For more details the reader might find this paper [?] informative. The meta-parameters of this layer type are a subset of those of a convolutional layer, and are described in the section below.

#### Convolutional Network Layout

A typical convolution network will first include several convolutional layers with different meta-parameters, each of them possibly followed by max-pooling layers to reduce overfitting. The last layers are fully connected (dense), in order to facilitate a soft-max activation similarly to traditional ANNs. An example layout configuration is depicted in the snippet below:

```
Layout=CONV|12|3|3|1|1|1|1|RELU, CONV|12|3|3|1|1|1|1|RELU, MAXPOOL|2|2|2|2,
RESHAPE|FLAT, DENSE|64|RELU, DENSE|1|LINEAR
```

Here we define a layout consisting of 2 convolutional layers, the last of which is followed by a pooling layer. After those layers, we include 2 fully connected layers where the last one consists of a single neuron (binary classification). We also employ a flattening reshape layer necessary to connect the multi-dimensional output, produced by convolutional layers, to a dense layer.

Figure ?? visually explains the layout parameterization of the first convolutional layer.

#### Convolutional Layer Layout

The parameters of the Convolutional layers are defined in the Layout string and they are separated by the '|' delimiter character. Note that all of them must be
The parameters are the following, in order of occurrence:

- **Layer Type (string)**: CONV for defining a 2D convolutional layer.
- **Number of kernels (integer)**, e.g. 12 in the above example.
- **Filter height (integer)**.
- **Filter width (integer)**.
- **Vertical stride (integer)**. Note that the stride must be a divisor of the input image size.
- **Horizontal stride (integer)**.
- **Vertical padding (integer)**. The padding value has to be a valid one, i.e. it provides an output image size which is an integer and not fractional. See the formula below defining the output size.
- **Horizontal padding (integer)**.
- **Activation function (string)**: See Table 22 for the list of possible activation functions.

The Padding and stride parameter values must be consistent and have values that used with the given input image sizes and filter sizes result in integer output sizes and not fractional. The formula that defines the output size is:

\[
o = \frac{i - f + 2 \times p}{s} + 1\]  

(99)

where \(o\) is the horizontal (or vertical) output size, \(i\), \(f\), \(p\), \(s\) are the corresponding input, filter, padding and stride sizes. In the example above with \(f = 3\), \(s = 1\), \(p = 1\) in both dimensions, the output image size will be equal to the input image size.
MaxPool Layer Layout  The MaxPool layer is typically used in combination with convolutional layers in a CNN to down-sample the images. The MaxPool layer computes the maximum of the input in the provided rectangular pool size. The parameters describing the MaxPool layer are:

- Layer Type (string) : MAXPOOL for 2D max pooling layer.
- Pool height dimension (integer) : e.g. 2 in the example above.
- Pool width dimension (integer).
- Vertical stride (integer).
- Horizontal stride (integer).

Note that the output images after a maxpooling will be reduced(down-sampled) according to the following formula:

\[ o = \frac{i - f}{s} + 1 \]  \hspace{1cm} (100)

Reshape Layer  The reshape layer is used to flatten the image that is the output from a convolutional layer or a maxpool layer in a format that can be input by a dense layer. After a convolution layer the output data will have a dimension equal to batch size × number of kernels × output height × output width \((B \times C \times H \times W)\). The reshape will reformat the data to a 2-dimensional tensor (i.e. a matrix) with batch size as number of rows and number of kernels × height × width as number of columns \((B \times CHW)\) that can be processed by a dense layer. The reshape layer can also perform the inverse operation (de-flattening).

The parameters of the reshape layers are different depending on the flattening or de-flattening case. In the flattening case the output depth, height and width can be omitted:

- Layer Type (string) : RESHAPE.
- 1st output dimension (integer), e.g. output depth.
- 2nd output dimension (integer).
- 3rd output dimension (integer).
- string type: FLAT for a flattening layer. In that case no need to specify the output dimension.

Batch Normalization Layer  The batch normalization layer normalizes the input by re-centering and re-scaling using the average and the standard deviation of the input batch. It is described in this paper [51]. The parameters for this layer are:

- Layer type: BNORM.
- Momentum $\rho$, used for the running average (floating). Optional, default is $\rho = 0.99$.
- $\epsilon$ parameter used for computing batchnorm (see formula in [51]). Optional, default is $\epsilon = 0.001$.

### 8.11.6 Recurrent Neural Networks

Recurrent neural network is a class of artificial neural networks where connections between nodes form a directed graph along a temporal sequence. This allows it to exhibit temporal dynamic behavior. Different types of recurrent networks exist. This implementation provides the support for 3 types of recurrent layers:

- Simple Recurrent layer (Vanilla RNN) as described for example in par. 10.2 of [52].
- LSTM (Long Short-Term Memory) layer as defined in [53]
- GRU (Gate Recurrent unit) layer as defined in [54]

Here is an example of defining a recurrent neural networks composed of one LSTM layer followed by a dense layer for classification:

```
InputLayout=10|3:
Layout=LSTM|12|3|10|0|1|RESHAP|FLAT,DENSE|64|RELU,DENSE|1|LINEAR
```

The layer layout parameters for recurrent layers are

- Layer Type (string) : RNN (Vanilla RNN), LSTM, GRU.
- Layer state size (integer).
- Input size (integer)
- Time steps size (integer), i.e. number of recurrent cells.
- Remember state option (boolean), default is false (0). This corresponds to the stateful option in Keras.
- Return the full output time sequence (boolean): default is false (0), i.e. return only output of last recurrent cell.
- Reset gate after option (boolean) : default is false. This is an option that applies only for GRU which defines a slightly different implementation.. This option is forced to be true when using the GPU implementation, since it is the way cuDNN implements GRU. Note that with this option the GRU computation is slightly faster.
8.11.7 Input Layout

When using a convolutional or a recurrent network it is needed to provide an input layout string defining the format of the input, i.e. its shape. In the case of convolutional networks the size of the input image (e.g. number of channels, image height, image width) must be defined and in the case of recurrent networks we the time and the feature dimensions must be provided. In the case of simple dense layer there is no need for an input layout because the total number of input event features can be deduced internally by TMVA.

The input layout shape is defined as following; here we give the example for input image with 3 channels, height = 32, width = 32:

\[
\text{InputLayout}=3|32|32
\]

The input layout parameters (input tensor dimensions) are separated as in the case of the layer layout by the ‘|’ character. For a RNN it is enough to provide an input layout of 2 dimensions: time and feature size. For example for an RNN with an input dimension of 3 features in 10 time steps, we will have an input layout as following

\[
\text{InputLayout}=10|3
\]

8.11.8 Training Strategy

The training of deep neural networks is performed in one or several consecutive training phases that are specified using the TrainingStrategy argument of the ⟨options⟩ string passed to the BookMethod call. A single phase is specified by the parameters described in Table 23. Different training phases are separated by ‘|’ characters. As an example, a valid training strategy string would be the following:

\[
\text{TrainingStrategy} = \text{LearningRate}=1e-1, \text{MaxEpoch}=20
| \text{LearningRate}=1e-2, \text{MaxEpoch}=30
| \text{LearningRate}=1e-3, \text{MaxEpoch}=40
\]

This training strategy trains the network in three training phases each with a lower learning rate. This is particular useful if using the SGD optimizers which requires a manual adaptation of the learning rate. When using instead an optimizer, such as ADAM, the learning rate adapts automatically and there is no such need to define several training strategies.

Dropout is a regularization technique that with a certain probability sets neuron activations to zero. This probability can be set for all layers at once by giving a single floating point value or a value for each hidden layer of the network separated by ‘+’ signs. Probabilities should be given by a value in the interval \([0, 1]\).
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BatchSize</td>
<td>–</td>
<td>30</td>
<td>–</td>
<td>Batch size used for this training phase.</td>
</tr>
<tr>
<td>ConvergenceSteps</td>
<td>–</td>
<td>100</td>
<td>–</td>
<td>Convergence criterion: Number of training epochs without improvement in test error to be performed before ending the training phase</td>
</tr>
<tr>
<td>MaxEpochs</td>
<td>–</td>
<td>2000</td>
<td>–</td>
<td>Maximum number of training epochs to exit before convergence.</td>
</tr>
<tr>
<td>TestRepetitions</td>
<td>–</td>
<td>7</td>
<td>–</td>
<td>Interval for evaluation of the test set error in epochs.</td>
</tr>
<tr>
<td>Optimizer</td>
<td>–</td>
<td>ADAM</td>
<td>ADAM, SGD, ADAGRAD, RMSPROP, ADADELTA</td>
<td>Type of optimizer used for this training strategy</td>
</tr>
<tr>
<td>DropConfig</td>
<td>–</td>
<td>0.0</td>
<td>–</td>
<td>Dropout fractions for each layer separated by ‘+’ signs or a single number if the same for all layers.</td>
</tr>
<tr>
<td>LearningRate</td>
<td>–</td>
<td>$10^{-5}$</td>
<td>–</td>
<td>Learning rate $\alpha$ to use for the current training phase.</td>
</tr>
<tr>
<td>Momentum</td>
<td>–</td>
<td>0.0</td>
<td>–</td>
<td>The momentum $p$ to use for the training. See section 8.11.2.</td>
</tr>
<tr>
<td>Regularization</td>
<td>–</td>
<td>NONE</td>
<td>NONE, L1, L2</td>
<td>Type of regularization</td>
</tr>
<tr>
<td>WeightDecay</td>
<td>–</td>
<td>0.0</td>
<td>–</td>
<td>Scaling factor applied to the L1 or L2 norms of the weight matrices when using regularization.</td>
</tr>
</tbody>
</table>

**Option Table 23:** Configuration options reference for MVA method: *DL*. 
8.12 Support Vector Machine (SVM)

In the early 1960s a linear support vector method has been developed for the construction of separating hyperplanes for pattern recognition problems [39, 40]. It took 30 years before the method was generalised to nonlinear separating functions [41, 42] and for estimating real-valued functions (regression) [43]. At that moment it became a general purpose algorithm, performing classification and regression tasks which can compete with neural networks and probability density estimators. Typical applications of SVMs include text categorisation, character recognition, bio-informatics and face detection.

The main idea of the SVM approach to classification problems is to build a hyperplane that separates signal and background vectors (events) using only a minimal subset of all training vectors (support vectors). The position of the hyperplane is obtained by maximizing the margin (distance) between it and the support vectors. The extension to nonlinear SVMs is performed by mapping the input vectors onto a higher dimensional feature space in which signal and background events can be separated by a linear procedure using an optimally separating hyperplane. The use of kernel functions eliminates thereby the explicit transformation to the feature space and simplifies the computation.

The implementation of the newly introduced regression is similar to the approach in classification. It also maps input data into higher dimensional space using previously chosen support vectors. Instead of separating events of two types, it determines the hyperplane with events of the same value (which is equal to the mean from all training events). The final value is estimated based on the distance to the hyperplane which is computed by the selected kernel function.

8.12.1 Booking options

The SVM classifier is booked via the command:

```c
factory->BookMethod( TMVA::Types::kSVM, "SVM", "<options>");
```

**Code Example 59**: Booking of the SVM classifier: the first argument is a unique type enumerator, the second is a user-defined name which must be unique among all booked classifiers, and the third argument is the configuration option string. Individual options are separated by a ‘:’. For options that are not set in the string default values are used. See Sec. 3.1.5 for more information on the booking.

The configuration options for the SVM classifier are given in Option Table 24.

8.12.2 Description and implementation

A detailed description of the SVM formalism can be found, for example, in Ref. [44]. Here only a brief introduction of the TMVA implementation is given.
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gamma</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>RBF kernel parameter: Gamma (size of the Kernel)</td>
</tr>
<tr>
<td>C</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Cost parameter</td>
</tr>
<tr>
<td>Tol</td>
<td>–</td>
<td>0.01</td>
<td>–</td>
<td>Tolerance parameter</td>
</tr>
<tr>
<td>MaxIter</td>
<td>–</td>
<td>1000</td>
<td>–</td>
<td>Maximum number of training loops</td>
</tr>
</tbody>
</table>

**Option Table 24**: Configuration options reference for MVA method: SVM. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured.

**Linear SVM**

Consider a simple two-class classifier with oriented hyperplanes. If the training data is linearly separable, a vector-scalar pair \((\vec{w}, b)\) can be found that satisfies the constraints

\[
y_i (\vec{x}_i \cdot \vec{w} + b) - 1 \geq 0, \quad \forall_i, \tag{101}
\]

where \(\vec{x}_i\) are the input vectors, \(y_i\) the desired outputs \((y_i = \pm 1)\), and where the pair \((\vec{w}, b)\) defines a hyperplane. The decision function of the classifier is \(f(\vec{x}_i) = \text{sign}(\vec{x}_i \cdot \vec{w} + b)\), which is +1 for all points on one side of the hyperplane and −1 for the points on the other side.

Intuitively, the classifier with the largest margin will give better separation. The margin for this linear classifier is just \(2/|\vec{w}|\). Hence to maximise the margin, one needs to minimise the cost function \(W = |\vec{w}|^2/w\) with the constraints from Eq. (101).

At this point it is beneficial to consider the significance of different input vectors \(\vec{x}_i\). The training events lying on the margins, which are called the support vectors (SV), are the events that contribute to defining the decision boundary (see Fig. 21). Hence if the other events are removed from the training sample and the classifier is retrained on the remaining events, the training will result in the same decision boundary. To solve the constrained quadratic optimisation problem, we first reformulate it in terms of a Lagrangian

\[
\mathcal{L}(\vec{w}, b, \vec{\alpha}) = \frac{1}{2} |\vec{w}|^2 - \sum_i \alpha_i \left( y_i ((\vec{x}_i \cdot \vec{w}) + b) - 1 \right) \tag{102}
\]

where \(\alpha_i \geq 0\) and the condition from Eq. (101) must be fulfilled. The Lagrangian \(\mathcal{L}\) is minimised with respect to \(\vec{w}\) and \(b\) and maximised with respect to \(\vec{\alpha}\). The solution has an expansion in terms of a subset of input vectors for which \(\alpha_i \neq 0\) (the support vectors):

\[
\vec{w} = \sum_i \alpha_i y_i \vec{x}_i, \tag{103}
\]

because \(\partial \mathcal{L} / \partial b = 0\) and \(\partial \mathcal{L} / \partial \vec{w} = 0\) hold at the extremum. The optimisation problem translates to finding the vector \(\vec{\alpha}\) which maximises

\[
\mathcal{L}(\vec{\alpha}) = \sum_i \alpha_i - \frac{1}{2} \sum_{ij} \alpha_i \alpha_j y_i y_j \vec{x}_i \cdot \vec{x}_j. \tag{104}
\]
Both the optimisation problem and the final decision function depend only on scalar products between input vectors, which is a crucial property for the generalisation to the nonlinear case.

**Nonseparable data**

The above algorithm can be extended to non-separable data. The classification constraints in Eq. (101) are modified by adding a “slack” variable $\xi_i$ to it ($\xi_i = 0$ if the vector is properly classified, otherwise $\xi_i$ is the distance to the decision hyperplane)

$$y_i(x_i \cdot \vec{w} + b) + 1 + \xi_i \geq 0, \quad \xi_i \geq 0, \quad \forall i.$$  \hspace{1cm} (105)

This admits a certain amount of misclassification. The training algorithm thus minimises the modified cost function

$$W = \frac{1}{2} |\vec{w}|^2 + C \sum_i \xi_i,$$  \hspace{1cm} (106)

describing a trade-off between margin and misclassification. The cost parameter $C$ sets the scale by how much misclassification increases the cost function (see Tab. 24).

**Nonlinear SVM**

The SVM formulation given above can be further extended to build a nonlinear SVM which can classify nonlinearly separable data. Consider a function $\Phi : \mathbb{R}^{n_{var}} \rightarrow \mathcal{H}$, which maps the training
data from $\mathbb{R}^{n_{\text{var}}}$, where $n_{\text{var}}$ is the number of discriminating input variables, to some higher dimensional space $\mathcal{H}$. In the $\mathcal{H}$ space the signal and background events can be linearly separated so that the linear SVM formulation can be applied. We have seen in Eq. (104) that event variables only appear in the form of scalar products $\vec{x}_i \cdot \vec{x}_j$, which become $\Phi(\vec{x}_i) \cdot \Phi(\vec{x}_j)$ in the higher dimensional feature space $\mathcal{H}$. The latter scalar product can be approximated by a kernel function

$$K(\vec{x}_i, \vec{x}_j) \approx \Phi(\vec{x}_i) \cdot \Phi(\vec{x}_j), \quad (107)$$

which avoids the explicit computation of the mapping function $\Phi(\vec{x})$. This is desirable because the exact form of $\Phi(\vec{x})$ is hard to derive from the training data. Most frequently used kernel functions are

$$K(\vec{x}, \vec{y}) = (\vec{x} \cdot \vec{y} + \theta)^d \quad \text{Polynomial},$$
$$K(\vec{x}, \vec{y}) = \exp\left(-|\vec{x} - \vec{y}|^2 / 2\sigma^2\right) \quad \text{Gaussian},$$
$$K(\vec{x}, \vec{y}) = \tanh(\kappa(\vec{x} \cdot \vec{y}) + \theta) \quad \text{Sigmoidal}, \quad (108)$$

of which currently only the Gaussian Kernel is implemented in TMVA. It was shown in Ref. [43] that a suitable function kernel must fulfill Mercer’s condition

$$\int K(\vec{x}, \vec{y})g(\vec{x})g(\vec{y})d\vec{x}d\vec{y} \geq 0, \quad (109)$$

for any function $g$ such that $\int g^2(\vec{x})d\vec{x}$ is finite. While Gaussian and polynomial kernels are known to comply with Mercer’s condition, this is not strictly the case for sigmoidal kernels. To extend the linear methodology to nonlinear problems one substitutes $\vec{x}_i \cdot \vec{x}_j$ by $K(\vec{x}_i, \vec{x}_j)$ in Eq. (104). Due to Mercer’s conditions on the kernel, the corresponding optimisation problem is a well defined convex quadratic programming problem with a global minimum. This is an advantage of SVMs compared to neural networks where local minima occur.

For regression problems, the same algorithm is used as for classification with the exception that instead of dividing events based on their type (signal/background), it separates them based on the value (larger/smaller than average). In the end, it does not return the sigmoid of the distance between the event and the hyperplane, but the distance itself – increased by the average target value.

**Implementation**

The TMVA implementation of the Support Vector Machine follows closely the description given in the literature. It employs a sequential minimal optimisation (SMO) [45] to solve the quadratic problem. Acceleration of the minimisation is achieved by dividing a set of vectors into smaller subsets [46]. The number of training subsets is controlled by option $\text{NSubSets}$. The SMO method drives the subset selection to the extreme by selecting subsets of two vectors (for details see Ref. [44]). The pairs of vectors are chosen, using heuristic rules, to achieve the largest possible improvement (minimisation) per step. Because the working set is of size two, it is straightforward to write down the analytical solution. The minimisation procedure is repeated recursively until the minimum is found. The SMO algorithm has proven to be significantly faster than other methods and has become the most common minimisation method used in SVM implementations. The precision of
the minimisation is controlled by the tolerance parameter Tol (see Tab. 24). The SVM training
time can be reduced by increasing the tolerance. Most classification problems should be solved with
less then 1000 training iterations. Interrupting the SVM algorithm using the option MaxIter may
thus be helpful when optimising the SVM training parameters. MaxIter can be released for the
final classifier training.

8.12.3 Variable ranking

The present implementation of the SVM classifier does not provide a ranking of the input variables.

8.12.4 Performance

The TMVA SVM algorithm comes currently only with the Gaussian kernel function. With sufficient
training statistics, the Gaussian kernel allows to approximate any separating function in the input
space. It is crucial for the performance of the SVM to appropriately tune the kernel parameters and
the cost parameter $C$. In case of a Gaussian, the kernel is tuned via option Gamma which is related
to the width $\sigma$ by $\Gamma = 1/(2\sigma^2)$. The optimal tuning of these parameters is specific to the problem
and must be done by the user.

The SVM training time scales with $n^2$, where $n$ is the number of vectors (events) in the training
data set. The user is therefore advised to restrict the sample size during the first rough scan of the
kernel parameters. Also increasing the minimisation tolerance helps to speed up the training.

SVM is a nonlinear general purpose classification and regression algorithm with a performance
similar to neural networks (Sec. 8.10) or to a multidimensional likelihood estimator (Sec. 8.3).

8.13 Boosted Decision and Regression Trees

A decision (regression) tree is a binary tree structured classifier (regressor) similar to the one
sketched in Fig. 22. Repeated left/right (yes/no) decisions are taken on one single variable at
a time until a stop criterion is fulfilled. The phase space is split this way into many regions that are
eventually classified as signal or background, depending on the majority of training events that end
up in the final leaf node. In case of regression trees, each output node represents a specific value of
the target variable. The boosting (see Sec. 7) of a decision (regression) tree extends this concept
from one tree to several trees which form a forest. The trees are derived from the same training
ensemble by reweighting events, and are finally combined into a single classifier (regressor) which
is given by a (weighted) average of the individual decision (regression) trees. Boosting stabilizes
the response of the decision trees with respect to fluctuations in the training sample and is able to
considerably enhance the performance w.r.t. a single tree. In the following, we will use the term
decision tree for both, decision- and regression trees and we refer to regression trees only if both
types are treated differently.

31The target variable is the variable the regression “function” is trying to estimate.
8.13.1 Booking options

The boosted decision (regression) tree (BDT) classifier is booked via the command:

```
factory->BookMethod( Types::kBDT, "BDT", "<options>" );
```

**Code Example 60:** Booking of the BDT classifier: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a `:`. See Sec. 3.1.5 for more information on the booking.

Several configuration options are available to customize the BDT classifier. They are summarized in Option Tables 25 and 27 and described in more detail in Sec. 8.13.2.
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NTrees</td>
<td>–</td>
<td>800</td>
<td>–</td>
<td>Number of trees in the forest</td>
</tr>
<tr>
<td>MaxDepth</td>
<td>–</td>
<td>3</td>
<td>–</td>
<td>Max depth of the decision tree allowed</td>
</tr>
<tr>
<td>MinNodeSize</td>
<td>–</td>
<td>5%</td>
<td>–</td>
<td>Minimum percentage of training events required in a leaf node (default: Classification: 5%, Regression: 0.2%)</td>
</tr>
<tr>
<td>nCuts</td>
<td>–</td>
<td>20</td>
<td>–</td>
<td>Number of grid points in variable range used in finding optimal cut in node splitting</td>
</tr>
<tr>
<td>BoostType</td>
<td>–</td>
<td>AdaBoost</td>
<td>AdaBoost, RealAdaBoost, Bagging, AdaBoostR2, Grad</td>
<td>Boosting type for the trees in the forest (note: AdaCost is still experimental)</td>
</tr>
<tr>
<td>AdaBoostR2Loss</td>
<td>–</td>
<td>Quadratic</td>
<td>Linear, Quadratic, Exponential</td>
<td>Type of Loss function in AdaBoostR2</td>
</tr>
<tr>
<td>UseBaggedGrad</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Use only a random subsample of all events for growing the trees in each iteration. (Only valid for GradBoost)</td>
</tr>
<tr>
<td>Shrinkage</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>Learning rate for GradBoost algorithm</td>
</tr>
<tr>
<td>AdaBoostBeta</td>
<td>–</td>
<td>0.5</td>
<td>–</td>
<td>Learning rate for AdaBoost algorithm</td>
</tr>
<tr>
<td>UseRandomisedTrees</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Determine at each node splitting the cut variable only as the best out of a random subset of variables (like in RandomForests)</td>
</tr>
<tr>
<td>UseNvars</td>
<td>–</td>
<td>2</td>
<td>–</td>
<td>Size of the subset of variables used with RandomisedTree option</td>
</tr>
<tr>
<td>UsePoissonNvars</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Interpret UseNvars not as fixed number but as mean of a Possion distribution in each split with RandomisedTree option</td>
</tr>
<tr>
<td>BaggedSampleFraction</td>
<td>–</td>
<td>0.6</td>
<td>–</td>
<td>Relative size of bagged event sample to original size of the data sample (used whenever bagging is used (i.e. UseBaggedGrad, Bagging,)</td>
</tr>
<tr>
<td>UseYesNoLeaf</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Use Sig or Bkg categories, or the purity=$S/(S+B)$ as classification of the leaf node -&gt; Real-AdaBoost</td>
</tr>
</tbody>
</table>

**Option Table 25:** Configuration options reference for MVA method: BDT. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured. The table is continued in Option Table 27.
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NegWeightTreatment</td>
<td>–</td>
<td>InverseBoostNegWeights, IgnoreNegWeightsInTraining, PairNegWeightsGlobal, Pray</td>
<td>to treat events with negative weights in the BDT training (particular the boosting): IgnoreInTraining; Boost With inverse boostweight; Pair events with negative and positive weights in training sample and <em>annihilate</em> them (experimental)</td>
<td></td>
</tr>
<tr>
<td>NodePurityLimit</td>
<td>–</td>
<td>0.5</td>
<td></td>
<td>In boosting/pruning, nodes with purity &gt; NodePurityLimit are signal; background otherwise.</td>
</tr>
<tr>
<td>SeparationType</td>
<td>–</td>
<td>GiniIndex</td>
<td>CrossEntropy, GiniIndex, GiniIndexWithLaplace, MisClassificationError, SDivSqrtSPlusB, RegressionVariance</td>
<td>Separation criterion for node splitting</td>
</tr>
<tr>
<td>DoBoostMonitor</td>
<td>–</td>
<td>False</td>
<td></td>
<td>Create control plot with ROC integral vs tree number</td>
</tr>
<tr>
<td>UseFisherCuts</td>
<td>–</td>
<td>False</td>
<td></td>
<td>Use multivariate splits using the Fisher criterion</td>
</tr>
<tr>
<td>MinLinCorrForFisher</td>
<td>–</td>
<td>0.8</td>
<td></td>
<td>The minimum linear correlation between two variables demanded for use in Fisher criterion in node splitting</td>
</tr>
<tr>
<td>UseExclusiveVars</td>
<td>–</td>
<td>False</td>
<td></td>
<td>Variables already used in fisher criterion are not anymore analysed individually for node splitting</td>
</tr>
<tr>
<td>DoPreseletion</td>
<td>–</td>
<td>False</td>
<td></td>
<td>and and apply automatic pre-selection for 100% efficient signal (bkg) cuts prior to training</td>
</tr>
<tr>
<td>RenormByClass</td>
<td>–</td>
<td>False</td>
<td></td>
<td>Individually re-normalize each event class to the original size after boosting</td>
</tr>
<tr>
<td>SigToBkgFraction</td>
<td>–</td>
<td>1</td>
<td></td>
<td>Sig to Bkg ratio used in Training (similar to NodePurityLimit, which cannot be used in real adaboost)</td>
</tr>
</tbody>
</table>

Option Table 26: Continuation of Option Table 25.
### Option Table 27: Some deprecated options for the BDT Method that are for the moment still kept for compatibility.

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>PruneMethod</td>
<td>–</td>
<td>NoPruning</td>
<td>NoPruning, ExpectedError, CostComplexity</td>
<td>Note: for BDTs use small trees (e.g. MaxDepth=3) and NoPruning; Pruning: Method used for pruning (removal) of statistically insignificant branches</td>
</tr>
<tr>
<td>PruneStrength</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>Pruning strength</td>
</tr>
<tr>
<td>PruningValFraction</td>
<td>–</td>
<td>0.5</td>
<td>–</td>
<td>Fraction of events to use for optimizing automatic pruning.</td>
</tr>
<tr>
<td>nEventsMin</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>deprecated: Use MinNodeSize (in % of training events) instead</td>
</tr>
<tr>
<td>GradBaggingFraction</td>
<td>–</td>
<td>0.6</td>
<td>–</td>
<td>deprecated: Use <em>BaggedSampleFraction</em> instead: Defines the fraction of events to be used in each iteration, e.g. when UseBaggedGrad=kTRUE.</td>
</tr>
<tr>
<td>UseNTrainEvents</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>deprecated: Use <em>BaggedSampleFraction</em> instead: Number of randomly picked training events used in randomised (and bagged) trees</td>
</tr>
<tr>
<td>NNodesMax</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>deprecated: Use MaxDepth instead to limit the tree size</td>
</tr>
</tbody>
</table>
8.13.2 Description and implementation

Decision trees are well known classifiers that allow a straightforward interpretation as they can be visualized by a simple two-dimensional tree structure. They are in this respect similar to rectangular cuts. However, whereas a cut-based analysis is able to select only one hypercube as region of phase space, the decision tree is able to split the phase space into a large number of hypercubes, each of which is identified as either “signal-like” or “background-like”, or attributed a constant event (target) value in case of a regression tree. For classification trees, the path down the tree to each leaf node represents an individual cut sequence that selects signal or background depending on the type of the leaf node.

A shortcoming of decision trees is their instability with respect to statistical fluctuations in the training sample from which the tree structure is derived. For example, if two input variables exhibit similar separation power, a fluctuation in the training sample may cause the tree growing algorithm to decide to split on one variable, while the other variable could have been selected without that fluctuation. In such a case the whole tree structure is altered below this node, possibly resulting also in a substantially different classifier response.

This problem is overcome by constructing a forest of decision trees and classifying an event on a majority vote of the classifications done by each tree in the forest. All trees in the forest are derived from the same training sample, with the events being subsequently subjected to so-called boosting (see 7), a procedure which modifies their weights in the sample. Boosting increases the statistical stability of the classifier and is able to drastically improve the separation performance compared to a single decision tree. However, the advantage of the straightforward interpretation of the decision tree is lost. While one can of course still look at a limited number of trees trying to interpret the training result, one will hardly be able to do so for hundreds of trees in a forest. Nevertheless, the general structure of the selection can already be understood by looking at a limited number of individual trees. In many cases, the boosting performs best if applied to trees (classifiers) that, taken individually, have not much classification power. These so called “weak classifiers” are small trees, limited in growth to a typical tree depth of as small as two, depending on the how much interaction there is between the different input variables. By limiting the tree depth during the tree building process (training), the tendency of overtraining for simple decision trees which are typically grown to a large depth and then pruned, is almost completely eliminated.

8.13.3 Boosting, Bagging and Randomising

The different “boosting” algorithms (in the following we will call also bagged and and randomised trees “boosted”) available for decision trees in TMVA are currently:

- AdaBoost (Discrete AdaBoost, see Sec. 7.1), RealAdaBoost (see below and [27]) and AdaBoostR2(see Sec. 35) for regression
- Gradient Boost (see Sec. 7.2)
- Bagging (see Sec. 7.3)
Randomised Trees, like the Random Forests of L. Breiman [30]. Each tree is grown in such a way that at each split only a random subset of all variables is considered. Moreover, each tree in the forest can be grown using only a (resampled) subset of the original training events. The size of the subset as well as the number of variables considered at each split can be set using the options BaggedSampleFraction (this requires the option UseBaggedBoost=True) and UseNVars.

A possible modification of Eq. (30) for the result of the combined classifier from the forest is to use the training purity\textsuperscript{32} in the leaf node as respective signal or background weights rather than relying on the binary decision. This is then called Real-AdaBoost can be chosen as one of the boost algorithms. For other boosting algorithms rather than AdaBoost, the same averaging of the individual trees can be chosen using the option UseYesNoLeaf=False.

Training (Building) a decision tree

The training, building or growing of a decision tree is the process that defines the splitting criteria for each node. The training starts with the root node, where an initial splitting criterion for the full training sample is determined. The split results in two subsets of training events that each go through the same algorithm of determining the next splitting iteration. This procedure is repeated until the whole tree is built. At each node, the split is determined by finding the variable and corresponding cut value that provides the best separation between signal and background. The node splitting stops once it has reached the minimum number of events which is specified in the BDT configuration (option nEventsMin). The leaf nodes are classified as signal or background according to the class the majority of events belongs to. If the option UseYesNoLeaf is set the end-nodes are classified in the same way. If UseYesNoLeaf is set to false the end-nodes are classified according to their purity.

A variety of separation criteria can be configured (option SeparationType see Option Table 27) to assess the performance of a variable and a specific cut requirement. Because a cut that selects predominantly background is as valuable as one that selects signal, the criteria are symmetric with respect to the event classes. All separation criteria have a maximum where the samples are fully mixed, i.e., at purity $p = 0.5$, and fall off to zero when the sample consists of one event class only. Tests have revealed no significant performance disparity between the following separation criteria:

- **Gini Index** [default], defined by $p \cdot (1 - p)$;
- **Cross entropy**, defined by $-p \cdot \ln(p) - (1 - p) \cdot \ln(1 - p)$;
- **Misclassification error**, defined by $1 - \max(p, 1 - p)$;
- **Statistical significance**, defined by $S/\sqrt{S + B}$;

\textsuperscript{32}The purity of a node is given by the ratio of signal events to all events in that node. Hence pure background nodes have zero purity.
• *Average squared error*, defined by $1/N \cdot \sum^N_{i=1} (y_i - \hat{y})^2$ for regression trees where $y$ is the regression target of each event in the node and $\hat{y}$ is its mean value over all events in the node (which would be the estimate of $y$ that is given by the node).

Since the splitting criterion is always a cut on a single variable, the training procedure selects *the* variable and cut value that optimises the *increase* in the separation index between the parent node and the sum of the indices of the two daughter nodes, weighted by their relative fraction of events. The cut values are optimised by scanning over the variable range with a granularity that is set via the option `nCuts`. The default value of `nCuts=20` proved to be a good compromise between computing time and step size. Finer stepping values did not increase noticeably the performance of the BDTs. However, a truly optimal cut, given the training sample, is determined by setting `nCuts=-1`. This invokes an algorithm that tests all possible cuts on the training sample and finds the best one. The latter is of course “slightly” slower than the coarse grid.

In principle, the splitting could continue until each leaf node contains only signal or only background events, which could suggest that perfect discrimination is achievable. However, such a decision tree would be strongly overtrained. To avoid overtraining a decision tree must be *pruned*.

**Pruning a decision tree**

Pruning is the process of cutting back a tree from the bottom up after it has been built to its maximum size. Its purpose is to remove statistically insignificant nodes and thus reduce the overtraining of the tree. For simple decision trees it has been found to be beneficial to first grow the tree to its maximum size and then cut back, rather than interrupting the node splitting at an earlier stage. This is because apparently insignificant splits can nevertheless lead to good splits further down the tree. For Boosted Decision trees however, as the boosting algorithms perform best on weak classifiers, pruning is unnecessary as one should rather drastically limit the tree depth far stronger than any pruning algorithm would do afterwards. Hence, while pruning algorithms are still implemented in TMVA, they are obsolete as they should not be used.

**8.13.4 Variable ranking**

A ranking of the BDT input variables is derived by counting how often the variables are used to split decision tree nodes, and by weighting each split occurrence by the separation gain-squared it has achieved and by the number of events in the node [32]. This measure of the variable importance can be used for a single decision tree as well as for a forest.

**8.13.5 Performance**

Only limited experience has been gained so far with boosted decision trees in HEP. In the literature decision trees are sometimes referred to as the best “out of the box” classifiers. This is because little tuning is required in order to obtain reasonably good results. This is due to the simplicity of the
method where each training step (node splitting) involves only a one-dimensional cut optimisation. Decision trees are also insensitive to the inclusion of poorly discriminating input variables. While for artificial neural networks it is typically more difficult to deal with such additional variables, the decision tree training algorithm will basically ignore non-discriminating variables as for each node splitting only the best discriminating variable is used. However, the simplicity of decision trees has the drawback that their theoretically best performance on a given problem is generally inferior to other techniques like neural networks. This is seen for example using the academic training samples included in the TMVA package. For this sample, which has equal RMS but shifted mean values for signal and background and linear correlations between the variables only, the Fisher discriminant provides theoretically optimal discrimination results. While the artificial neural networks are able to reproduce this optimal selection performance the BDTs always fall short in doing so. However, in other academic examples with more complex correlations or real life examples, the BDTs often outperform the other techniques. This is because either there are not enough training events available that would be needed by the other classifiers, or the optimal configuration (i.e. how many hidden layers, which variables) of the neural network has not been specified. We have only very limited experience at the time with the regression, hence cannot really comment on the performance in this case.

8.14 Predictive learning via rule ensembles (RuleFit)

This classifier is a TMVA implementation of Friedman-Popescu’s RuleFit method described in [34]. Its idea is to use an ensemble of so-called rules to create a scoring function with good classification power. Each rule \( r_i \) is defined by a sequence of cuts, such as

\[
\begin{align*}
  r_1(x) &= I(x_2 < 100.0) \cdot I(x_3 > 35.0), \\
  r_2(x) &= I(0.45 < x_4 < 1.00) \cdot I(x_1 > 150.0), \\
  r_3(x) &= I(x_3 < 11.00),
\end{align*}
\]

where the \( x_i \) are discriminating input variables, and \( I(\ldots) \) returns the truth of its argument. A rule applied on a given event is non-zero only if all of its cuts are satisfied, in which case the rule returns 1.

The easiest way to create an ensemble of rules is to extract it from a forest of decision trees (cf. Sec. 8.13). Every node in a tree (except the root node) corresponds to a sequence of cuts required to reach the node from the root node, and can be regarded as a rule. Hence for the tree illustrated in Fig. 22 on page 130 a total of 8 rules can be formed. Linear combinations of the rules in the ensemble are created with coefficients (rule weights) calculated using a regularised minimisation procedure [35]. The resulting linear combination of all rules defines a score function (see below) which provides the RuleFit response \( y_{RF}(x) \).

In some cases a very large rule ensemble is required to obtain a competitive discrimination between signal and background. A particularly difficult situation is when the true (but unknown) scoring function is described by a linear combination of the input variables. In such cases, e.g., a Fisher discriminant would perform well. To ease the rule optimisation task, a linear combination of the
input variables is added to the model. The minimisation procedure will then select the appropriate coefficients for the rules and the linear terms. More details are given in Sec. 8.14.2 below.

### 8.14.1 Booking options

The RuleFit classifier is booked via the command:

```c++
factory->BookMethod( Types::kRuleFit, "RuleFit", "<options>" );
```

**Code Example 61**: Booking of RuleFit: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ‘:’. See Sec. 3.1.5 for more information on the booking.

The RuleFit configuration options are given in Option Table 28.

### 8.14.2 Description and implementation

As for all TMVA classifiers, the goal of the rule learning is to find a classification function \( y_{RF}(x) \) that optimally classifies an event according to the tuple of input observations (variables) \( x \). The classification function is written as

\[
y_{RF}(x) = a_0 + \sum_{m=1}^{M_R} a_m f_m(x),
\]

where the set \( \{ f_m(x) \}_{M_R} \) forms an ensemble of base learners with \( M_R \) elements. A base learner may be any discriminating function derived from the training data. In our case, they consist of rules and linear terms as described in the introduction. The complete model then reads

\[
y_{RF}(x) = a_0 + \sum_{m=1}^{M_R} a_m r_m(x) + \sum_{i=1}^{n_{\text{var}}} b_i x_i.
\]

To protect against outliers, the variables in the linear terms are modified to

\[
x'_i = \min(\delta^+_i, \max(\delta^-_i)) ,
\]

where \( \delta^+_i \) are the lower and upper \( \beta \) quantiles\(^{33}\) of the variable \( x_i \). The value of \( \beta \) is set by the option **LinQuantile**. If the variables are used “as is”, they may have an unequal \textit{a priori} influence relative to the rules. To counter this effect, the variables are normalised

\[
x'_i \rightarrow \sigma_r \cdot x'_i / \sigma_i,
\]

where \( \sigma_r \) and \( \sigma_i \) are the estimated standard deviations of an ensemble of rules and the variable \( x'_i \), respectively.

\(^{33}\)Quantiles are points taken at regular intervals from the PDF of a random variable. For example, the 0.5 quantile corresponds to the median of the PDF.
8.14 Predictive learning via rule ensembles (RuleFit)

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GDTau</td>
<td>–</td>
<td>-1</td>
<td>–</td>
<td>Gradient-directed (GD) path: default fit cut-off</td>
</tr>
<tr>
<td>GDTauPrec</td>
<td>–</td>
<td>0.01</td>
<td>–</td>
<td>GD path: precision of tau</td>
</tr>
<tr>
<td>GDStep</td>
<td>–</td>
<td>0.01</td>
<td>–</td>
<td>GD path: step size</td>
</tr>
<tr>
<td>GDSteps</td>
<td>–</td>
<td>10000</td>
<td>–</td>
<td>GD path: number of steps</td>
</tr>
<tr>
<td>GDErrScale</td>
<td>–</td>
<td>1.1</td>
<td>–</td>
<td>Stop scan when error &gt; scale*errmin</td>
</tr>
<tr>
<td>LinQuantile</td>
<td>–</td>
<td>0.025</td>
<td>–</td>
<td>Quantile of linear terms (removes outliers)</td>
</tr>
<tr>
<td>GDPathEveFrac</td>
<td>–</td>
<td>0.5</td>
<td>–</td>
<td>Fraction of events used for the path search</td>
</tr>
<tr>
<td>GDValidEveFrac</td>
<td>–</td>
<td>0.5</td>
<td>–</td>
<td>Fraction of events used for the validation</td>
</tr>
<tr>
<td>fEventsMin</td>
<td>–</td>
<td>0.1</td>
<td>–</td>
<td>Minimum fraction of events in a split-table node</td>
</tr>
<tr>
<td>fEventsMax</td>
<td>–</td>
<td>0.9</td>
<td>–</td>
<td>Maximum fraction of events in a split-table node</td>
</tr>
<tr>
<td>nTrees</td>
<td>–</td>
<td>20</td>
<td>–</td>
<td>Number of trees in forest</td>
</tr>
<tr>
<td>ForestType</td>
<td>–</td>
<td>AdaBoost</td>
<td>AdaBoost, Random</td>
<td>Method to use for forest generation (AdaBoost or RandomForest)</td>
</tr>
<tr>
<td>RuleMinDist</td>
<td>–</td>
<td>0.001</td>
<td>–</td>
<td>Minimum distance between rules</td>
</tr>
<tr>
<td>MinImp</td>
<td>–</td>
<td>0.01</td>
<td>–</td>
<td>Minimum rule importance accepted</td>
</tr>
<tr>
<td>Model</td>
<td>–</td>
<td>ModRuleLinear</td>
<td>ModRule, ModRuleLinear, ModLinear</td>
<td>Model to be used</td>
</tr>
<tr>
<td>RuleFitModule</td>
<td>–</td>
<td>RFTMVA</td>
<td>RFTMVA, RFFriedman</td>
<td>Which RuleFit module to use</td>
</tr>
<tr>
<td>RFWorkDir</td>
<td>–</td>
<td>./rulefit</td>
<td>–</td>
<td>Friedman’s RuleFit module (RFF): working dir</td>
</tr>
<tr>
<td>RFNrules</td>
<td>–</td>
<td>2000</td>
<td>–</td>
<td>RFF: Maximum number of rules</td>
</tr>
<tr>
<td>RFNendnodes</td>
<td>–</td>
<td>4</td>
<td>–</td>
<td>RFF: Average number of end nodes</td>
</tr>
</tbody>
</table>

Option Table 28: Configuration options reference for MVA method: RuleFit. Values given are defaults. If predefined categories exist, the default category is marked by a ‘⋆’. The options in Option Table 9 on page 71 can also be configured.

**Rule generation**

The rules are extracted from a forest of decision trees. There are several ways to generate a forest. In the current RuleFit implementation, each tree is generated using a fraction of the training sample. The fraction depends on which method is used for generating the forest. Currently two methods are supported (selected by option ForestType): AdaBoost and Random Forest. The first method
is described in Sec. 8.13.2. In that case, the whole training set is used for all trees. The diversity is obtained through using different event weights for each tree. For a random forest, though, the diversity is created by training each tree using random sub-samples. If this method is chosen, the fraction is calculated from the training sample size \( N \) (signal and background) using the empirical formula \[36]\)

\[
f = \min(0.5, (100.0 + 6.0 \cdot \sqrt{N})/N).
\]

By default, \texttt{AdaBoost} is used for creation of the forest. In general it seems to perform better than the random forest.

The topology of each tree is controlled by the parameters \texttt{fEventsMin} and \texttt{fEventsMax}. They define a range of fractions which are used to calculate the minimum number of events required in a node for further splitting. For each tree, a fraction is drawn from a uniform distribution within the given range. The obtained fraction is then multiplied with the number of training events used for the tree, giving the minimum number of events in a node to allow for splitting. In this way both large trees (small fraction) giving complex rules and small trees (large fraction) for simple rules are created. For a given forest of \( N_t \) trees, where each tree has \( n_t \) leaf nodes, the maximum number of possible rules is

\[
M_{R,\text{max}} = \sum_{i=1}^{N_t} 2(n_{t,i} - 1).
\]

To prune similar rules, a distance is defined between two \textit{topologically equal} rules. Two rules are topologically equal if their cut sequences follow the same variables only differing in their cut values. The rule distance used in TMVA is then defined by

\[
\delta^2_R = \sum_i \frac{\delta^2_{i,L} + \delta^2_{i,U}}{\sigma_i^2},
\]

where \( \delta_{i,L(U)} \) is the difference in lower (upper) limit between the two cuts containing the variable \( x_i \), \( i = 1, \ldots, n_{\text{var}} \). The difference is normalised to the RMS-squared \( \sigma^2_i \) of the variable. Similar rules with a distance smaller than \texttt{RuleMinDist} are removed from the rule ensemble. The parameter can be tuned to improve speed and to suppress noise. In principle, this should be achieved in the fitting procedure. However, pruning the rule ensemble using a distance cut will reduce the fitting time and will probably also reduce the number of rules in the final model. Note that the cut should be used with care since a too large cut value will deplete the rule ensemble and weaken its classification performance.

\textbf{Fitting}

Once the rules are defined, the coefficients in Eq. (111) are fitted using the training data. For details, the fitting method is described in [35]. A brief description is provided below to motivate the corresponding RuleFit options.

A \textit{loss function} \( L(y_{\text{RF}}(x)|\hat{y}) \), given by the “squared-error ramp” [35]

\[
L(y_{\text{RF}}|\hat{y}) = (\hat{y} - H(y_{\text{RF}}))^2,
\]

\[117]
where \( H(y) = \max(-1, \min(y_{RF}, 1)) \), quantifies the “cost” of misclassifying an event of given true class \( \hat{y} \). The risk \( R \) is defined by the expectation value of \( L \) with respect to \( x \) and the true class. Since the true distributions are generally not known, the average of \( N \) training events is used as an estimate

\[
R = \frac{1}{N} \sum_{i=1}^{N} L(y_{RF}(x_i) | \hat{y}_i).
\]

(118)

A line element in the parameter space of the rule weights (given by the vector \( a \) of all coefficients) is then defined by

\[
a(\epsilon + \delta\epsilon) = a(\epsilon) + \delta\epsilon \cdot g(\epsilon),
\]

(119)

where \( \delta\epsilon \) is a positive small increment and \( g(\epsilon) \) is the negative derivative of the estimated risk \( R \), evaluated at \( a(\epsilon) \). The estimated risk-gradient is evaluated using a sub-sample (\( GDPathEveFrac \)) of the training events.

Starting with all weights set to zero, the consecutive application of Eq. (119) creates a path in the \( a \) space. At each step, the procedure selects only the gradients \( g_k \) with absolute values greater than a certain fraction (\( \tau \)) of the largest gradient. The fraction \( \tau \) is an \emph{a priori} unknown quantity between 0 and 1. With \( \tau = 0 \) all gradients will be used at each step, while only the strongest gradient is selected for \( \tau = 1 \). A measure of the “error” at each step is calculated by evaluating the risk (Eq. 118) using the validation sub-sample (\( GDValidEveFrac \)). By construction, the risk will always decrease at each step. However, for the validation sample the value will increase once the model starts to be overtrained. Currently, the fitting is crudely stopped when the error measure is larger than \( GDErrScale \) times the minimum error found. The number of steps is controlled by \( GDNSteps \) and the step size (\( \delta\epsilon \) in Eq. 119) by \( GDStep \).

If the selected \( \tau \) (\( GDTau \)) is a negative number, the best value is estimated by means of a scan. In such a case several paths are fitted in parallel, each with a different value of \( \tau \). The number of paths created depend on the required precision on \( \tau \) given by \( GDTauPrec \). By only selecting the paths being “close enough” to the minimum at each step, the speed for the scan is kept down. The path leading to the lowest estimated error is then selected. Once the best \( \tau \) is found, the fitting proceeds until a minimum is found. A simple example with a few scan points is illustrated in Fig. 23.

### 8.14.3 Variable ranking

Since the input variables are normalised, the ranking of variables follows naturally from the coefficients of the model. To each rule \( m \) (\( m = 1, \ldots, M_R \)) can be assigned an importance defined by

\[
I_m = |a_m| \sqrt{s_m(1.0 - s_m)},
\]

(120)

where \( s_m \) is the \textit{support} of the rule with the following definition

\[
s_m = \frac{1}{N} \sum_{n=1}^{N} r_m(x_n).
\]

(121)

The support is thus the average response for a given rule on the data sample. A large support implies that many events pass the cuts of the rule. Hence, such rules cannot have strong discriminating
Figure 23: An example of a path scan in two dimensions. Each point represents an $\epsilon$ in Eq. (119) and each step is given by $\delta\epsilon$. The direction along the path at each point is given by the vector $g$. For the first few points, the paths $\tau(1, 2, 3)$ are created with different values of $\tau$. After a given number of steps, the best path is chosen and the search is continued. It stops when the best point is found. That is, when the estimated error-rate is minimum.

power. On the other hand, rules with small support only accept few events. They may be important for these few events they accept, but they are not in the overall picture. The definition (120) for the rule importance suppresses rules with both large and small support.

For the linear terms, the definition of importance is

$$I_i = |b_i| \cdot \sigma_i,$$

so that variables with small overall variation will be assigned a small importance.

A measure of the variable importance may then be defined by

$$J_i = I_i + \sum_{m|x_i \in r_m} I_m/q_m,$$

where the sum is over all rules containing the variable $x_i$, and $q_m$ is the number of variables used in the rule $r_m$. This is introduced in order to share the importance equally between all variables in rules with more than one variable.

8.14.4 Friedman’s module

By setting RuleFitModule to RFFriedman, the interface to Friedman’s RuleFit is selected. To use this module, a separate setup is required. If the module is selected in a run prior to setting up the
environment, TMVA will stop and give instructions on how to proceed. A command sequence to setup Friedman’s RuleFit in a UNIX environment is:

```bash
~> mkdir rulefit
~> cd rulefit
~> wget http://www-stat.stanford.edu/~jhf/r-rulefit/linux/rf_go.exe
~> chmod +x rf_go.exe
```

**Code Example 62:** The first line creates a working directory for Friedman's module. In the third line, the binary executable is fetched from the official web-site. Finally, it is made sure that the module is executable.

As of this writing, binaries exists only for Linux and Windows. Check J. Friedman’s home page at [http://www-stat.stanford.edu/~jhf](http://www-stat.stanford.edu/~jhf) for updated information. When running this module from TMVA, make sure that the option RFWorkDir is set to the proper working directory (default is `./rulefit`). Also note that only the following options are used: Model, RFWorkDir, RFNrules, RFNendnodes, GDNSteps, GDStep and GDErrScale. The options RFNrules and RFNendnodes correspond in the package by Friedman-Popescu to the options max.rules and tree.size, respectively. For more details, the reader is referred to Friedman’s RuleFit manual [36].

**Technical note**

The module `rf_go.exe` communicates with the user by means of both ASCII and binary files. This makes the input/output from the module machine dependant. TMVA reads the output from `rf_go.exe` and produces the normal machine independent weight (or class) file. This can then be used in other applications and environments.

### 8.14.5 Performance

Rule ensemble based learning machines are not yet well known within the HEP community, although they start to receive some attention [37]. Apart from RuleFit [34] other rule ensemble learners exists, such as SLIPPER [38].

The TMVA implementation of RuleFit follows closely the original design described in Ref. [34]. Currently the performance is however slightly less robust than the one of the Friedman-Popescu package. Also, the experience using the method is still scarce at the time of this writing.

To optimise the performance of RuleFit several strategies can be employed. The training consists of two steps, rule generation and rule ensemble fitting. One approach is to modify the complexity of the generated rule ensemble by changing either the number of trees in the forest, or the complexity of each tree. In general, large tree ensembles with varying trees sizes perform better than short non-complex ones. The drawback is of course that fitting becomes slow. However, if the fitting performs well, it is likely that a large amount of rules will have small or zero coefficients. These can
be removed, thus simplifying the ensemble. The fitting performance can be improved by increasing
the number of steps along with using a smaller step size. Again, this will be at the cost of speed
performance although only at the training stage. The setting for the parameter $\tau$ may greatly affect
the result. Currently an automatic scan is performed by default. In general, it should find the
optimum $\tau$. If in doubt, the user may set the value explicitly. In any case, the user is initially
advised to use the automatic scan option to derive the best path.

9 The PyMVA Methods

PyMVA is the interface for third-party MVA tools based on Python. It is created to make powerful
external libraries easily accessible with a direct integration into the TMVA workflow. All PyMVA
methods provide the same plug-and-play mechanisms than the TMVA methods described in section
8.

Because the base method of PyMVA is inherited from the TMVA base method, all options from
the Option Table 9 apply for PyMVA methods as well.

9.1 Keras

Keras (www.keras.io) is a high-level wrapper for the machine learning frameworks Theano (www.
deeplearning.net/software/theano/) and TensorFlow (www.tensorflow.org), which are mainly used
to set up deep neural networks.

9.1.1 Training and Testing Data

Please note that data defined as testing data in the dataloader is used for validation of the model
after each epoch. Such as stated in section 8.11.2, a proper evaluation of the performance should
be done on a completely independent dataset.

9.1.2 Booking Options

The Keras method is booked mainly similar to other TMVA methods such as shown in Code
Example 63. The different part is the definition of the neural network model itself. These options
are not set in the options string of TMVA, but these are defined in Python and exported to a model
file, which is later on loaded by the method. The definition of such a model using the Keras API is
explained in detail in section 9.1.3. The settings in the method’s option string manage the training
process.

The full list of possible settings that can be used in the option string is presented in Table 29.
Code Example 63: Booking of the PyKeras method

```cpp
factory->BookMethod(dataloader, TMVA::Types::kPyKeras, "Keras", <options>);
```

**Option Table 29:** Configuration options reference for PyMVA method PyKeras.

<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>FilenameModel</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>Filename of the neural network model defined by Keras</td>
</tr>
<tr>
<td>FilenameTrainedModel</td>
<td>—</td>
<td>Placed in weights directory of dataloader</td>
<td>—</td>
<td>Filename of the model with trained weights. By default, the file is placed in the weights directory of the dataloader, but you can define with this option a custom path to the model file with trained weights.</td>
</tr>
<tr>
<td>BatchSize</td>
<td>—</td>
<td>100</td>
<td>—</td>
<td>Batch size used for a single gradient step during training</td>
</tr>
<tr>
<td>NumEpochs</td>
<td>—</td>
<td>10</td>
<td>—</td>
<td>Number of training epochs</td>
</tr>
<tr>
<td>Verbose</td>
<td>—</td>
<td>1</td>
<td>—</td>
<td>Verbosity during training</td>
</tr>
<tr>
<td>ContinueTraining</td>
<td>—</td>
<td>false</td>
<td>—</td>
<td>Continue training by loading weights from FilenameTrainedModel at beginning of training</td>
</tr>
<tr>
<td>SaveBestOnly</td>
<td>—</td>
<td>true</td>
<td>—</td>
<td>Store only weights with smallest validation loss</td>
</tr>
<tr>
<td>TriesEarlyStopping</td>
<td>—</td>
<td>-1</td>
<td>—</td>
<td>Number of epochs with no improvement in validation loss after which training will be stopped. The default or a negative number deactivates this option.</td>
</tr>
<tr>
<td>LearningRateSchedule</td>
<td>—</td>
<td>' '</td>
<td>—</td>
<td>Set new learning rate during training at specific epochs, e.g., set new learning rate at epochs 50 and 70 with '50, 0.01;70, 0.005'. By default, there is no schedule set.</td>
</tr>
</tbody>
</table>

9.1.3 Model Definition

The model of the neural network is defined in Python using the Keras API, stored to a file and then loaded by the Keras method of PyMVA. This section gives an detailed example how this can be done for a classification task. The final reference to build a Keras model is the Keras documentation itself (www.keras.io). As well, ROOT ships with TMVA tutorials, which include examples for the usage of the PyKeras method for binary classification, multiclass classification and regression.
Code Example 64 shows as an example a simple three-layer neural network, which can be applied for any classification task. After running this script, the model file, e.g. model.h5 in this example, has to be set as the option FilenameModel of the method.

```python
from keras.models import Sequential
from keras.layers.core import Dense

# Define model architecture
model = Sequential()
model.add(Dense(64, init='glorot_normal', activation='relu',
                input_dim=num_input_variables))
model.add(Dense(num_output_classes, init='glorot_uniform',
                activation='softmax'))

# Set loss function and optimizer algorithm
model.compile(loss='categorical_crossentropy', optimizer='Adam',
              metrics=['accuracy',])

# Store model to file
model.save('model.h5')
```

**Code Example 64:** Definition of a classification model in Python using Keras. The placeholders `num_input_variables` and `num_output_classes` have to be set accordingly to the specific task.

### 9.1.4 Variable Ranking

A variable ranking is not supported.

### 10 Combining MVA Methods

In challenging classification or regression problems with a high demand for optimisation, or when treating feature spaces with strongly varying properties, it can be useful to combined MVA methods. There is large room for creativity inherent in such combinations. For TMVA we distinguish three classes of combinations:

1. **Boosting** MVA methods,
2. **Categorising** MVA methods,
3. Building committees of MVA methods.

While the first two combined methods are available since TMVA 4.1, the third one is still under development. The MVA **booster** is a generalisation of the Boosted Decision Trees approach (see
Sec. 8.13) to any method in TMVA. *Category methods* allow the user to specify sub-regions of phase space, which are assigned by requirements on input or spectator variables, and which define disjoint sub-populations of the training sample, thus improving the modelling of the training data. Finally, *Committee methods* allow one to input MVA methods into other MVA methods, a procedure that can be arbitrarily chained.

All of these combined methods are of course MVA methods themselves, treated just like any other method in TMVA for training, testing, evaluation and application. This also allows to categorise a committee method, for example.

### 10.1 Boosted classifiers

Since generalised boosting is not yet available for regression in TMVA, we restrict the following discussion to classification applications. A boosted classifier is a combination of a collection of classifiers of the same type trained on the same sample but with different events weights.\(^{34}\) The response of the final classifier is a weighted response of each individual classifier in the collection. The boosted classifier is potentially more powerful and more stable with respect to statistical fluctuations in the training sample. The latter is particularly the case for bagging as “boost” algorithm (cf. Sec. 7.3, page 68).

The following sections do not apply to decision trees. We refer to Sec. 8.13 (page 129) for a description of boosted decision trees. In the current version of TMVA only the AdaBoost and Bagging algorithms are implemented for the boost of arbitrary classifiers. The boost algorithms are described in detail in Sec. 7 on page 65.

#### 10.1.1 Booking options

To book a boosted classifier, one needs to add the booster options to the regular classifier’s option string. The minimal option required is the number of boost iterations *Boost_Num*, which must be set to a value larger than zero. Once the Factory detects a *Boost_Num* > 0 in the option string it books a boosted classifier and passes all boost options (recognised by the prefix *Boost_*) to the Boost method and the other options to the regular classifier.

```
factory->BookMethod( TMVA::Types::kLikelihood, "BoostedLikelihood", 
                      "Boost_Num=10:Boost_Type=Bagging:Spline=2:NSmooth=5:NAvEvtPerBin=50" );
```

**Code Example 65:** Booking of the boosted classifier: the first argument is the predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. All options with the prefix *Boost_* (in this example the first two options) are passed on to the boost method, the other options are provided to the regular classifier (which in this case is Likelihood). Individual options are separated by a ‘:’. See Sec. 3.1.5 for more information on the booking.

\(^{34}\)The Boost method is at the moment only applicable to classification problems.
<table>
<thead>
<tr>
<th>Option</th>
<th>Array</th>
<th>Default</th>
<th>Predefined Values</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boost_Num</td>
<td>–</td>
<td>100</td>
<td>–</td>
<td>Number of times the classifier is boosted</td>
</tr>
<tr>
<td>Boost_MonitorMethod</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Write monitoring histograms for each boosted classifier</td>
</tr>
<tr>
<td>Boost_DetailedMonitoring</td>
<td>–</td>
<td>False</td>
<td>–</td>
<td>Produce histograms for detailed boost-wise monitoring</td>
</tr>
<tr>
<td>Boost_Type</td>
<td>–</td>
<td>AdaBoost</td>
<td>AdaBoost, Bagging, HighEdgeGauss, HighEdgeCoPara</td>
<td>Boosting type for the classifiers</td>
</tr>
<tr>
<td>Boost_BaggedSampleFraction</td>
<td>–</td>
<td>0.6</td>
<td>–</td>
<td>Relative size of bagged event sample to original size of the data sample (used whenever bagging is used)</td>
</tr>
<tr>
<td>Boost_MethodWeightType</td>
<td>–</td>
<td>ByError</td>
<td>ByError, Average, ByROC, ByOverlap, LastMethod</td>
<td>How to set the final weight of the boosted classifiers</td>
</tr>
<tr>
<td>Boost_RecalculateMVACut</td>
<td>–</td>
<td>True</td>
<td>–</td>
<td>Recalculate the classifier MVA Signal-like cut at every boost iteration</td>
</tr>
<tr>
<td>Boost_AdaBoostBeta</td>
<td>–</td>
<td>1</td>
<td>–</td>
<td>The ADA boost parameter that sets the effect of every boost step on the events’ weights</td>
</tr>
<tr>
<td>Boost_Transform</td>
<td>–</td>
<td>step</td>
<td>step, linear, log, gauss</td>
<td>Type of transform applied to every boosted method linear, log, step</td>
</tr>
<tr>
<td>Boost_RandomSeed</td>
<td>–</td>
<td>0</td>
<td>–</td>
<td>Seed for random number generator used for bagging</td>
</tr>
</tbody>
</table>

Option Table 30: Boosting configuration options. These options can be simply added to a simple classifier’s option string or used to form the option string of an explicitly booked boosted classifier.

The boost configuration options are given in Option Table 30.

The options most relevant for the boost process are the number of boost iterations, Boost_Num, and the choice of the boost algorithm, Boost_Type. In case of Boost_Type=AdaBoost, the option Boost_Num describes the maximum number of boosts. The algorithm is iterated until an error rate of 0.5 is reached or until Boost_Num iterations occurred. If the algorithm terminates after to few iterations, the number might be extended by decreasing the $\beta$ variable (option Boost_AdaBoostBeta). Within the AdaBoost algorithm a decision must be made how to classify an event, a task usually done by the user. For some classifiers it is straightforward to set a cut on the MVA response to define signal-like events. For the others, the MVA cut is chosen that the error rate is minimised. The option Boost_RecalculateMVACut determines whether this cut should be recomputed for every boosting iteration. In case of Bagging as boosting algorithm the number of boosting iterations always reaches Boost_Num.
10.1 Boosted classifiers

By default boosted classifiers are combined as a weighted average with weights computed from the misclassification error (option `Boost.MethodWeightType=ByError`). It is also possible to use the arithmetic average instead (`Boost.MethodWeightType=Average`).

10.1.2 Boostable classifiers

The boosting process was originally introduced for simple classifiers. The most commonly boosted classifier is the decision tree (DT – cf. Sec. 8.13, page 129). Decision trees need to be boosted a few hundred times to effectively stabilise the BDT response and achieve optimal performance.

Another simple classifier in the TMVA package is the Fisher discriminant (cf. Sec. 8.7, page 98 – which is equivalent to the linear discriminant described in Sec. 8.8). Because the output of a Fisher discriminant represents a linear combination of the input variables, a linear combination of different Fisher discriminants is again a Fisher discriminant. Hence linear boosting cannot improve the performance. It is nevertheless possible to effectively boost a linear discriminant by applying the linear combination not on the discriminant’s output, but on the actual classification results provided. This corresponds to a “non-linear” transformation of the Fisher discriminant output according to a step function. The Boost method in TMVA also features a fully non-linear transformation that is directly applied to the classifier response value. Overall, the following transformations are available:

- `linear`: no transformation is applied to the MVA output,
- `step`: the output is $-1$ below the step and $+1$ above (default setting),
- `log`: logarithmic transformation of the output.

The macro `Boost.C` (residing in the `macros (test)` directory for the sourceforge (ROOT) version of TMVA) provides examples for the use of these transformations to boost a Fisher discriminant. We point out that the performance of a boosted classifier strongly depends on its characteristics as well as on the nature of the input data. A careful adjustment of options is required if AdaBoost is applied to an arbitrary classifier, since otherwise it might even lead to a worse performance than for the unboosted method.

10.1.3 Monitoring tools

The current GUI provides figures to monitor the boosting process. Plotted are the boost weights, the classifier weights in the boost ensemble, the classifier error rates, and the classifier error rates using unboosted event weights. In addition, when the option `Boost.MonitorMethod=T` is set, monitoring histograms are created for each classifier in the boost ensemble. The histograms generated during the boosting process provide useful insight into the behaviour of the boosted classifiers and help

---

35 Note that in the TMVA standard example, which uses linearly correlated, Gaussian-distributed input variables for signal and background, a single Fisher discriminant already provides the theoretically maximum separation power. Hence on this example, no further gain can be expected by boosting, no matter what “tricks” are applied.
to adjust to the optimal number of boost iterations. These histograms are saved in a separate folder in the output file, within the folder of MethodBoost/<Title>/ Besides the specific classifier monitoring histograms, this folder also contains the MVA response of the classifier for the training and testing samples.

10.1.4 Variable ranking

The present boosted classifier implementation does not provide a ranking of the input variables.

10.2 Category Classifier

The Category method allows the user to separate the training data (and accordingly the application data) into disjoint sub-populations exhibiting different properties. The separation into phase space regions is done by applying requirements on the input and/or spectator variables (variables defined as spectators are not used as direct inputs to MVA methods). It thus reaches beyond a simple subclassification of the original feature space. In each of these disjoint regions (each event must belong to one and only one region), an independent training is performed using the most appropriate MVA method, training options and set of training variables in that zone. The division into categories in presence of distinct sub-populations reduces the correlations between the training variables, improves the modelling, and hence increases the classification and regression performance. Another common application is when variables are not available in some phase space regions. For example, a sub-detector may only cover a fraction of the training data sample. By defining two categories the variables provided by this detector could be excluded from the training in the phase space region outside of its fiducial volume. Finally, an advantage of categorisation, often exploited in likelihood fits, also lies in the gain of signal significance when the fraction of signal events in the event sample differs between the categories.

We point out that the explicit use of the Category method is not mandatory. The categorisation functionality can be achieved by hand by training independent MVAs in each of the disjoint regions. Our experience however shows that often the corresponding bookkeeping is considered too cumbersome and the required categorisation is not performed, thus leading to performance loss. The Category method is straightforward to implement and should help in almost all analyses to better describe the training sample. Performance comparison with the corresponding non-categorised method should guide the user in his/her choice.

For the current release, the Category method is only implemented for classification. Extension to regression is however straightforward and should follow soon.

\[^{36}\text{This is particularly the case for projective likelihood methods, even when including prior decorrelation of the input data (cf. Sec. 8.2).}\]

\[^{37}\text{In the above example with the reduced detector fiducial acceptance, it often occurs that non-physical values ("−99") are assigned to the variables for events where they are not available. Inserting such meaningless (apart from indicating the category) values into an MVA penalises its performance.}\]
10.2 Category Classifier

10.2.1 Booking options

The Category method is booked via the command:

```cpp
TMVA::IMethod* category = factory->BookMethod( TMVA::Types::kCategory,
                                              "Category",
                                              "<options>" );
```

**Code Example 66**: Booking of the Category method: the first argument is a predefined enumerator, the second argument is a user-defined string identifier, and the third argument is the configuration options string. Individual options are separated by a ‘:’. See Sec. 3.1.5 for more information on the booking.

The categories and sub-classifiers are defined as follows:

```cpp
TMVA::MethodCategory* mcategory = dynamic_cast<TMVA::MethodCategory*>(category);
mcategory->AddCategory( "<cut>",
                        "<variables>",
                        TMVA::Types::<enumerator>,
                        "<sub-classifier name>",
                        "<sub-classifier options>" );
```

**Code Example 67**: Adding a category to the Category method: the first argument is the cut that defines the category, the second defines is the set of variables used to train this sub-classifier, the third argument is the predefined enumerator of the sub-classifier, the fourth argument is a user-defined string identifier of the sub-classifier, and the last argument sets the configuration options of the sub-classifier. Individual variables and options are both separated by a ‘:’. See Sec. 10.2.2 for further information on cuts and variables. The dynamic cast is required to allow access to specific Category members.

There are no options implemented for the Category method itself, apart from those common to all methods (cf. Option Table 9 on page 71). The available options for the sub-classifiers are found in the corresponding sections of this Users Guide.

The following example illustrates how sub-classifiers are added:
mcategory->AddCategory( "abs(eta)<=1.3",
    "var1:var2:var3:var4:",
    TMVA::Types::kFisher,
    "Category_Fisher",
    "H:!V:Fisher" );

mcategory->AddCategory( "abs(eta)>1.3",
    "var1:var2:var3:",
    TMVA::Types::kFisher,
    "Category_Fisher",
    "H:!V:Fisher" );

10.2.2 Description and implementation

The Category method is implemented in an entirely transparent way, conserving full flexibility for
the user on which input variables, MVA method and options to use in each category, and how the
category should be defined.

The categories are defined via cuts on the input and/or spectator variables. These cuts can be
arbitrary boolean expressions or may involve any number of variables. In case of particularly
intricate category definitions it may turn out advantageous for the user to pre-define and pre-fill
category variables, and use those as spectators in TMVA to set an event’s category. It is required
that the cuts are defined in such a way that they create disjoint event samples. TMVA will issue a
fatal error in case of an ambiguous category definition.

The variables used for the training of each of the sub-classifiers can be chosen from the variables
and spectators that have been defined in the TMVA script. The individual variables have to be
separated by a ‘:’. The distinction between input variables and spectators is necessary to allow per-
formance comparisons between categorised and non-categorised methods within the same training
and evaluation job (spectator variables would not be used by non-categorised methods).

The advantage offered by the Category classifier is illustrated with the following academic example.
We consider four uncorrelated and Gaussian distributed input variables for training. The mean
values of the variables are shifted between signal and background, hence providing classification
information. In addition, all mean values (signal and background) depend on a spectator variable
eta: for abs(eta)>1.3 (abs(eta)<=1.3) they have a positive (negative) offset. Hence, the Gauss-
sians are broader and the variables receive an effective correlation when integrating over eta. The
distributions for signal and background of one out of the four variables is shown for both abs(eta)
categories in Fig. 24. Figure 25 shows the ROC curves (cf. Sec. 2.6) of the categorised and non-categorised Fisher and Projective Likelihood classifiers. Optimum performance is recovered by the Category classifier.

10.2.3 Variable ranking

Due to the nature of the Category classifier it does not provide a general variable ranking. As far as available, the sub-classifiers will provide ranking for each category.

10.2.4 Performance

The performance of the Category method depends entirely on the judicious choice of the categories, properly separating events with different properties, and on the performance of the sub-methods within these categories. Control by performance comparison with the corresponding non-categorised methods allows to measure the performance gain, and to assess whether categorisation improves the classification.

11 Which MVA method should I use for my problem?

There is obviously no generally valid answer to this question. To guide the user, we have attempted a coarse assessment of various MVA properties in Table 6. Simplicity is a virtue, but only if it is not at the expense of significant loss of discrimination power. A properly trained Neural Network with the problem dependent optimal architecture or a Support Vector Machine with the appropriate kernel size should theoretically give superior performance compared to a more robust “out of the box” Boosted Decision Trees, which in practice often outperforms the former.

To assess whether a linear discriminant analysis (LDA) could be sufficient for a classification (regression) problem, the user is advised to analyse the correlations among the discriminating variables (among the variables and regression target) by inspecting scatter and profile plots (it is not enough to print the correlation coefficients, which by definition are linear only). Using an LDA greatly reduces the number of parameters to be adjusted and hence allow smaller training samples. It usually is robust with respect to generalisation to larger data samples. For moderately intricate problems, the function discriminant analysis (FDA) with some added nonlinearity may be found sufficient. It is always useful to cross-check its performance against several of the sophisticated nonlinear methods to see how much can be gained over the use of the simple and very transparent FDA.

For problems that require a high degree of optimisation and allow to use a large number of input

38Note, in an earlier version we noted the problem of overtraining in BDTs when using large trees and trying to regulate them using pruning. This however is overcome completely when using trees with limited depth (say 2 to 4) and minimal leaf node size (some % of training sample size) instead of pruning, which in addition gives superior performance.
Figure 24: Examples for the usage of categories. The variable var4 depends on $\text{abs}(\eta)$: for $\text{abs}(\eta) \leq 1.3$ (left plot), the signal and background Gaussian distributions are both shifted to lower values, while for $\text{abs}(\eta) > 1.3$ (right plot) shifts to larger values have been applied. Ignoring this dependence, broadens the inclusive Gaussian distributions and generates variable correlations (here a coefficient of +0.4 between var3 and var4) due to the existence of distinct subsamples. The Category classifier treats the two cases independently, thus recovering optimal separation.

Figure 25: ROC curves for categorised and non-categorised Fisher and Projective Likelihood classifiers applied to an academic four-Gaussian example with $\eta$-dependent mean values distinguishing two categories (cf. Fig. 24 for the category-dependent signal and background distributions of one of the four input variables). The categorised (“...Cat”) classifiers recover optimum performance (their red and black curves are superimposed).
TABLE 6: Assessment of MVA method properties. The symbols stand for the attributes “good” (⋆⋆), “fair” (⋆) and “bad” (◦). “Curse of dimensionality” refers to the “burden” of required increase in training statistics and processing time when adding more input variables. See also comments in the text. The FDA method is not listed here since its properties depend on the chosen function.

variables, complex nonlinear methods like neural networks, the support vector machine, boosted decision trees and/or RuleFit are more appropriate.

Very involved multi-dimensional variable correlations with strong nonlinearities are usually best mapped by the multidimensional probability density estimators such as PDE-RS, k-NN and PDE-Foam, requiring however a reasonably low number of input variables.

For RuleFit classification we emphasise that the TMVA implementation differs from Friedman-Popescu’s original code [34], with slightly better robustness and out-of-the-box performance for the latter version. In particular, the behaviour of the original code with respect to nonlinear correlations and the curse of dimensionality would have merited two stars. We also point out that the excellent performance for by majority linearly correlated input variables is achieved somewhat artificially by adding a Fisher-like term to the RuleFit classifier (this is the case for both implementations, cf. Sec. 8.14 on page 137).

In Fig 6 (page 12) we have shown the example of ROC-curves obtained from various linear and non-linear classifiers on a simple toy Monte Carlo sample of linear correlated Gaussian distribution functions. Here all classifiers capable of dealing with such type of correlations are equally performing. Only the projective likelihood method, which ignores all correlations, performs significantly worse (it recovers optimum performance after prior decorrelation of the variables). For such a problem, a Fisher (aka linear) discriminant is optimal by construction, and the non-linear methods effectively reduce in the training to linear discriminants achieving competitive, but somewhat less robust

40 An interface to Friedman-Popescu’s original code can be requested from the TMVA authors. See Sec. 8.14.4.
performance.

For another academic toy Monte Carlo example however, which exhibits strongly non-linear correlations in form of a “Schachbrett” (chess board with two-dimensional Gaussian-distributed signal and background spots – see Fig. 26), the limitations of the linear classifiers, being unable to capture the features of the data, become manifest. Non-linear classifiers, once appropriately optimised such as choosing a sufficiently complex network architecture for the MLP, or the proper kernel functions in the SVM, all non-linear methods reproduce or approximate the theoretical limit of this classification problem (indicated by the thick red line in the ROCc curve in the lower plot of Fig. 26).

12 TMVA implementation status summary for classification and regression

All TMVA methods are fully operational for user analysis, requiring training, testing, evaluating and reading for the application to unknown data samples. Additional features are optional and – despite our attempts to provide a fully transparent analysis – not yet uniformly available. A status summary is given in Table 7 and annotated below.

Although since TMVA 4 the framework supports multi-dimensional MVA outputs it has not yet been implemented for classification. For regression, only a few methods are fully multi-target capable so far (see Table 7).

Individual event-weight support is now commonly realised, only missing (and not foreseen to be provided) for the less recommended neural network CFMlpANN. Support of negative event weights occurring, e.g., in NLO MC requires more scrutiny as discussed in Sec. 3.1.2 on page 18.

Ranking of the input variables cannot be defined in a straightforward manner for all MVA methods. Transparent and objective variable ranking through performance comparison of the MVA method under successive elimination of one input variable at a time is under consideration (so far only realised for the naive-Bayes likelihood classifier).

Standalone C++ response classes (not required when using the Reader application) are generated by the majority of the classifiers, but not yet for regression analysis. The missing ones for PDE-RS, PDE-Foam, k-NN, Cuts and CFMlpANN will only be considered on explicit request.

The availability of help messages, which assist the user with the performance tuning and which are printed on standard output when using the booking option ‘H’, is complete.

Finally, custom macros are provided for some MVA methods to analyse specific properties, such as the fidelity of likelihood reference distributions or the neural network architecture, etc. More macros can be added upon user request.
Figure 26: The top left plot shows the two-dimensional Gaussian-distributed signal (blue) and background (red) distributions for the “Schachbrett” toy Monte Carlo sample. In the top right plot is drawn as an example the Support Vector Machine’s classification response in terms of the weights given to events in the two dimensional plane. Dark red signifies large MVA output values, i.e. signal like event regions. The bottom plot gives the ROC curves obtained with several TMVA classifiers for this example after training. The theoretically best ROC curve is known and shown by the thick red outer line. While all the non-linear classification methods (after carefully tuning their parameters) are able to approximate the theoretical limit, the linear methods strongly underperform.
Table 7: Status of the methods with respect to various TMVA features. See text for comments. Note that the column “Standalone response class” only refers to classification. It is yet unavailable for regression.

<table>
<thead>
<tr>
<th>Method</th>
<th>ClassIFIcation</th>
<th>Regression</th>
<th>Multi-class/target</th>
<th>Treats event weights</th>
<th>Variable weights</th>
<th>Standalone response class</th>
<th>Help</th>
<th>CustomMVA method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cut optimisation</td>
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(*) Not a generic TMVA method – Interface to ROOT class TMultiLayerPerceptron.

- Classification: Methods available for classification.
- Regression: Methods available for regression.
- Multi-class/target: Methods available for multi-class or target classification.
- Treats event weights: Methods available for treating event weights.
- Variable weights: Methods available for variable weights.
- Standalone response class: Methods available for standalone response class.
- Help: Methods available for help.
- CustomMVA method: Methods available for custom MVA method.
13 Conclusions and Plans

TMVA is a toolkit that unifies highly customisable multivariate (MVA) classification and regression algorithms in a single framework thus ensuring convenient use and an objective performance assessment. It is designed for machine learning applications in high-energy physics, but not restricted to these. Source code and library of TMVA-v.3.5.0 and higher versions are part of the standard ROOT distribution kit (v5.14 and higher). The newest TMVA development version can be downloaded from Sourceforge.net at http://tmva.sourceforge.net.

This Users Guide introduced the main steps of a TMVA analysis allowing a user to optimise and perform her/his own multivariate classification or regression. Let us recall the main features of the TMVA design and purpose:

- TMVA works in transparent factory mode to allow an unbiased performance assessment and comparison: all MVA methods see the same training and test data, and are evaluated following the same prescription.

- A complete TMVA analysis consists of two steps:
  1. **Training**: the ensemble of available and optimally customised MVA methods are trained and tested on independent signal and background data samples; the methods are evaluated and the most appropriate (performing and concise) ones are selected.
  2. **Application**: selected trained MVA methods are used for the classification of data samples with unknown signal and background composition, or for the estimate of unknown target values (regression).

- A Factory class object created by the user organises the customisation and interaction with the MVA methods for the training, testing and evaluation phases of the TMVA analysis. The training results together with the configuration of the methods are written to result (“weight”) files in XML format.

- Standardised outputs during the Factory running, and dedicated ROOT macros allow a refined assessment of each method’s behaviour and performance for classification and regression.

- Once appropriate methods have been chosen by the user, they can be applied to data samples with unknown classification or target values. Here, the interaction with the methods occurs through a Reader class object created by the user. A method is booked by giving the path to its weight file resulting from the training stage. Then, inside the user’s event loop, the MVA response is returned by the Reader for each of the booked MVA method, as a function of the event values of the discriminating variables used as input for the classifiers. Alternatively, for classification, the user may request from the Reader the probability that a given event belongs to the signal hypothesis and/or the event’s Rarity.

- In parallel to the XML files, TMVA generates standalone C++ classes after the training, which can be used for classification problems (feature not available yet for regression). Such classes are available for all classifiers except for cut optimisation, PDE-RS, PDE-Foam, k-NN and the old CFMlpANN.
We give below a summary of the TMVA methods, outlining the current state of their implementation, their advantages and shortcomings.

- **Rectangular Cut Optimisation**
  The current implementation is mature. It includes speed-optimised range searches using binary trees, and three optimisation algorithms: Monte Carlo sampling, a Genetic Algorithm and Simulated Annealing. In spite of these tools, optimising the cuts for a large number of discriminating variables remains challenging. The user is advised to reduce the available dimensions to the most significant variables (e.g., using a principal component analysis) prior to optimising the cuts.

- **Likelihood**
  Automatic non-parametric probability density function (PDF) estimation through histogram smoothing and interpolation with various spline functions and quasi-unbinned kernel density estimators is implemented. The PDF description can be individually tuned for each input variable.

- **PDE-RS**
  The multidimensional probability density estimator (PDE) approach is in an advanced development stage featuring adaptive range search, several kernel estimation methods, and speed optimised range search using event sorting in binary trees. It has also been extended to regression.

- **PDE-Foam**
  This new multidimensional PDE algorithm uses self-adapting phase-space binning and is a fast realisation of PDE-RS in fixed volumes, which are determined and optimised during the training phase. Much work went into the development of PDE-Foam. It has been thoroughly tested, and can be considered a mature method. PDE-Foam performs classification and regression analyses.

- **k-NN**
  The k-Nearest Neighbour classifier is also in a mature state, featuring both classification and regression. The code has been well tested and shows satisfactory results. With scarce training statistics it may slightly underperform in comparison with PDE-RS, whereas it is significantly faster in the application to large data samples.

- **Fisher and H-Matrix**
  Both are mature algorithms, featuring linear discrimination for classification only. Higher-order correlations are taken care of by FDA (see below).

- **Linear Discriminant (LD)**
  LD is equivalent to Fisher but providing both classification and linear regression.

- **Function Discriminant Analysis (FDA)**
  FDA is a mature algorithm, which has not been extensively used yet. It extends the linear discriminant to moderately non-linear correlations that are fit to the training data.
• **Artificial Neural Networks**

Significant work went into the implementation of fast feed-forward multilayer perceptron algorithms into TMVA. Two external ANNs have been integrated as fully independent methods, and another one has been newly developed for TMVA, with emphasis on flexibility and speed. The performance of the latter ANN (MLP) has been cross checked against the Stuttgart ANN (using as an example \( \tau \) identification in ATLAS), and was found to achieve competitive performance. The MLP ANN also performs multi-target regression.

• **Support Vector Machine**

SVM is a relatively new multivariate analysis algorithm with a strong statistical background. It performs well for nonlinear discrimination and is insensitive to overtraining. Optimisation is straightforward due to a low number of adjustable parameters (only two in the case of Gaussian kernel). The response speed is slower than for a not-too-exhaustive neural network, but comparable with other nonlinear methods. SVM is being extended to multivariate regression.

• **Boosted Decision Trees**

The BDT implementation has received constant attention over the years of its development. The current version includes additional features like bagging or gradient boosting, and manual or automatic pruning of statistically insignificant nodes. It is a highly performing MVA method that also applies to regression problems.

• **RuleFit**

The current version has the possibility to run either the original program written by J. Friedman [34] or an independent TMVA implementation. The TMVA version has been improved both in speed and performance and achieves almost equivalent results with respect to the original one, requiring however somewhat more tuning.

The new framework introduced with TMVA 4 provides the flexibility to combine MVA methods in a general fashion. Exploiting these capabilities for classification and regression however requires to create so-called committee methods for each combination. So far, we provide a generalised Boost method, allowing to boost any classifier by simply setting the variable Boost_Num in the configuration options to a positive number (plus possible adjustment of other configuration parameters). The result is a potentially powerful committee method unifying the excellent properties of boosting with MVA methods that already represent highly optimised algorithms.

Boosting is not the only combination the new framework allows us to establish. TMVA now also supports the categorisation of classification according to the phase space, which allows a better modelling and hence simplifies the mapping of the feature space. This particularly improves simple methods, such as projective likelihood and linear discriminants.

**Acknowledgements**

The fast growth of TMVA would not have been possible without the contribution and feedback from many developers (also co-authors of this Users Guide) and users to whom we are indebted. We thank in particular the CERN Summer students Matt Jachowski (Stanford U.) for the implementation of TMVA’s MLP neural
network, and Yair Mahalalel (Tel Aviv U.) for a significant improvement of PDE-RS. The Support Vector Machine has been contributed to TMVA by Andrzej Zemla and Marcin Wolter (IFJ PAN Krakow), and the k-NN method has been written by Rustem Ospanov (Texas U.). We are grateful to Lucian Ancu, Doug Applegate, Kregg Arms, René Brun and the ROOT team, Andrea Bulgarelli, Marc Escalier, Zhiyi Liu, Colin Mclean, Elzbieta Richter-Was, Alfio Rizzo, Lydia Roos, Vincent Tisserand, Alexei Volk, Jiahang Zhong for helpful feedback and bug reports. Thanks also to Lucian Ancu for improving the plotting macros.
A  More Classifier Booking Examples

The Code Examples 69 and 70 give a (non-exhaustive) collection of classifier bookings with appropriate default options. They correspond to the example training job TMVAClassification.C.

```
// Cut optimisation using Monte Carlo sampling
factory->BookMethod( TMVA::Types::kCuts, "Cuts",
    "!H:!V:FitMethod=MC:EffSel:SampleSize=200000:VarProp=FSmart" );

// Cut optimisation using Genetic Algorithm
factory->BookMethod( TMVA::Types::kCuts, "CutsGA",
    Steps=30:Cycles=3:PopSize=400:SC_steps=10:SC_rate=5:SC_factor=0.95" );

// Cut optimisation using Simulated Annealing algorithm
factory->BookMethod( TMVA::Types::kCuts, "CutsSA",
    "!H:!V:FitMethod=SA:EffSel:MaxCalls=150000:KernelTemp=IncAdaptive:"n
    InitialTemp=1e+6:MinTemp=1e-6:Eps=1e-10:UseDefaultScale" );

// Likelihood classification (naive Bayes) with Spline PDF parametrisation
factory->BookMethod( TMVA::Types::kLikelihood, "Likelihood",
    "!H:!V:TransformOutput:PDFInterpol=Spline2:NSmoothSig[0]=20:"n
    NSmoothBkg[0]=20:NSmoothBkg[1]=10:NSmooth=1:NAvEvtPerBin=50" );

// Likelihood with decorrelation of input variables
factory->BookMethod( TMVA::Types::kLikelihood, "LikelihoodD",
    "!H:!V:TransformOutput:PDFInterpol=Spline2:NSmoothSig[0]=20:"n
    NSmoothBkg[0]=20:NSmooth=5:NAvEvtPerBin=50:VarTransform=Decorrelate" );

// Likelihood with unbinned kernel estimator for PDF parametrisation
factory->BookMethod( TMVA::Types::kLikelihood, "LikelihoodKDE",
    "!H:!V:TransformOutput:PDFInterpol=KDE:KDEtype=Gauss:KDEiter=Adaptive:"n
    KDEFineFactor=0.3:KDEborder=None:NAvEvtPerBin=50" );
```

Code Example 69: Examples for booking MVA methods in TMVA for application to classification and – where available – to regression problems. The first argument is a unique type enumerator (the available types can be looked up in src/Types.h), the second is a user-defined name (must be unique among all booked classifiers), and the third a configuration option string that is specific to the classifier. For options that are not set in the string default values are used. The syntax of the options should become clear from the above examples. Individual options are separated by a ‘:’. Boolean variables can be set either explicitly as MyBoolVar=True/False, or just via MyBoolVar/!MyBoolVar. All concrete option variables are explained in the tools and classifier sections of this Users Guide. The list is continued in Code Example 70.
// Probability density estimator range search method (multi-dimensional)
factory->BookMethod( TMVA::Types::kPDERS, "PDERS", 
   "!H:V:NormTree=T:VolumeRangeMode=Adaptive:KernelEstimator=Gauss:
   GaussSigma=0.3:NEventsMin=400:NEventsMax=600" );

// Multi-dimensional PDE using self-adapting phase-space binning
factory->BookMethod( TMVA::Types::kPDEFoam, "PDEFoam", 
   "H:V:SigBgSeparate=F:TailCut=0.001:VolFrac=0.0333:nActiveCells=500:\n   nSampl=2000:nBin=5:CutNmin=T:Nmin=100:Kernel=None:Compress=T" );

// k-Nearest Neighbour method (similar to PDE-RS)
factory->BookMethod( TMVA::Types::kKNN, "KNN", 
   "H:nkNN=20:ScaleFrac=0.8:SigmaFact=1.0:Kernel=Gaus:UseKernel=F:\n   UseWeight=T:!Trim" );

// H-matrix (chi-squared) method
factory->BookMethod( TMVA::Types::kHMatrix, "HMatrix", "!H:!V" );

// Fisher discriminant (also creating Rarity distribution of MVA output)
factory->BookMethod( TMVA::Types::kFisher, "Fisher", 
   "H:!V:Fisher:CreateMVAPdfs:PDFInterpolMVAPdf=Spline2:NbinsMVAPdf=60:\n   NsmoothMVAPdf=10" );

// Fisher discriminant with Gauss-transformed input variables
factory->BookMethod( TMVA::Types::kFisher, "FisherG", "VarTransform=Gauss" );

// Fisher discriminant with principle-value-transformed input variables
factory->BookMethod( TMVA::Types::kFisher, "FisherG", "VarTransform=PCA" );

// Boosted Fisher discriminant
factory->BookMethod( TMVA::Types::kFisher, "BoostedFisher", 
   "Boost_Num=20:Boost_Transform=log:\n   Boost_Type=AdaBoost:Boost_AdaBoostBeta=0.2" );

// Linear discriminant (same as Fisher, but also performing regression)
factory->BookMethod( TMVA::Types::kLD, "LD", "H:!V:VarTransform=None" );

// Function discrimination analysis (FDA), fitting user-defined function
factory->BookMethod( TMVA::Types::kFDA, "FDA_MT", 
   "H:!V:Formula=(0)+(1)*x0+(2)*x1+(3)*x2+(4)*x3:\n   ParRanges=(-1,1);(-10,10);(-10,10);(-10,10);(-10,10):FitMethod=MINUIT:\n   ErrorLevel=1:PrintLevel=-1:FitStrategy=2:UseImprove:UseMinos:SetBatch" );

// Artificial Neural Network (Multilayer perceptron) - TMVA version
factory->BookMethod( TMVA::Types::kMLP, "MLP",
   "H:!V:NeuronType=tanh:VarTransform=N:NCycles=600:HiddenLayers=N+5:\
   TestRate=5" );

// NN with BFGS quadratic minimisation
factory->BookMethod( TMVA::Types::kMLP, "MLPBFGS",
   "H:!V:NeuronType=tanh:VarTransform=N:NCycles=600:HiddenLayers=N+5:\
   TestRate=5:TrainingMethod=BFGS" );

// NN (Multilayer perceptron) - ROOT version
factory->BookMethod( TMVA::Types::kTMlpANN, "TMlpANN",
   "!H:!V:NCycles=200:HiddenLayers=N+1,N:LearningMethod=BFGS:
   ValidationFraction=0.3" );

// NN (Multilayer perceptron) - ALEPH version (depreciated)
factory->BookMethod( TMVA::Types::kCFMlpANN, "CFMlpANN",
   "!H:!V:NCycles=2000:HiddenLayers=N+1,N" );

// Support Vector Machine
factory->BookMethod( TMVA::Types::kSVM, "SVM", "Gamma=0.25:Tol=0.001" );

// Boosted Decision Trees with adaptive boosting
factory->BookMethod( TMVA::Types::kBDT, "BDT",
   "!H:!V:NTrees=400:nEventsMin=400:MaxDepth=3:BoostType=AdaBoost:\
   SeparationType=GiniIndex:nCuts=20:PruneMethod=NoPruning" );

// Boosted Decision Trees with gradient boosting
factory->BookMethod( TMVA::Types::kBDT, "BDTG",
   "!H:!V:NTrees=1000:BoostType=Grad:Shrinkage=0.30:UseBaggedGrad:
   GradBaggingFraction=0.6:SeparationType=GiniIndex:nCuts=20:\
   PruneMethod=CostComplexity:PruneStrength=50:NNodesMax=5" );

// Boosted Decision Trees with bagging
factory->BookMethod( TMVA::Types::kBDT, "BDTB",
   "!H:!V:NTrees=400:BoostType=Bagging:SeparationType=GiniIndex:\
   nCuts=20:PruneMethod=NoPruning" );

// Predictive learning via rule ensembles (RuleFit)
factory->BookMethod( TMVA::Types::kRuleFit, "RuleFit",
   "H:1V:RuleFitModule=RFTMVA:Model=ModRuleLinear:MinImp=0.001:\
   RuleMinDist=0.001:NTrees=20:fEventsMin=0.01:fEventsMax=0.5:\
   GD Tau=-1.0:GDTauPrec=0.01:GDStep=0.01:GDNsteps=10000:GDErrScale=1.02" );

Code Example 71: Continuation from Code Example 70.
References


[8] To our information, the Rarity has been originally defined by F. Le Diberder in an unpublished Mark II internal note. In a single dimension, as defined in Eq. (13), it is equivalent to the $\mu$-transform developed in: M. Pivk, “Etude de la violation de CP dans la désintégration $B^0 \rightarrow h^+h^-$ ($h = \pi,K$) auprès du détecteur BABAR à SLAC”, PhD thesis (in French), http://tel.archives-ouvertes.fr/documents/archives0/00/00/29/91/index.html (2003).


[33] We use the ROOT class TParallelCoord by B. Dalla Piazza, 2007.


