Data Analysis With TMVA

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MVA-Literature /Software Packages... a biased selection

Literature:
- C.M.Bishop, “Pattern Recognition and Machine Learning”, Springer 2006

Software packages for Multivariate Data Analysis/Classification
- individual classifier software
  - e.g. “JETNET” C.Peterson, T. Rognvaldsson, L.Loennblad
    and many other packages
- attempts to provide “all inclusive” packages
    [http://www.hep.caltech.edu/~narsky/spr.html](http://www.hep.caltech.edu/~narsky/spr.html)
    [http://tmva.sf.net](http://tmva.sf.net) or every ROOT distribution (development moved from SourceForge to ROOT repository)
  - “R”: a huge data analysis library: [http://www.r-project.org/](http://www.r-project.org/)

Conferences: PHYSTAT, ACAT,...
Event Classification

Suppose data sample of two types of events: with class labels \textit{Signal} and \textit{Background} (will restrict here to two class cases. Many classifiers can in principle be extended to several classes, otherwise, analyses can be staged)

- how to set the decision boundary to select events of type \textit{S}?
- we have discriminating variables \(x_1, x_2, \ldots\)

How can we decide what to uses?

Low variance (stable), high bias methods
High variance, small bias methods

Once decided on a class of boundaries, how to find the “optimal” one?
Regression

- how to estimate a “functional behaviour” from a given set of ‘known measurements’?
- assume for example “D”-variables that somehow characterize the shower in your calorimeter.

- seems trivial? → The human brain has very good pattern recognition capabilities!
- what if you have many input variables?

constant?  
linear?  
non-linear?

Energy  
f(x)  
Cluster Size  
x  

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Regression → model functional behaviour

- Assume for example “D”-variables that somehow characterize the shower in your calorimeter.
  - Monte Carlo or testbeam
    - data sample with measured cluster observables
    + known particle energy
    = calibration function (energy == surface in D+1 dimensional space)

1-D example

- better known: (linear) regression → fit a known analytic function
  - e.g. the above 2-D example → reasonable function would be: \( f(x) = ax^2 + by^2 + c \)
- what if we don’t have a reasonable “model”? → need something more general:
  - e.g. piecewise defined splines, kernel estimators, decision trees to approximate \( f(x) \)
Event Classification

- Each event, if **Signal** or **Background**, has “D” measured variables.
- Find a mapping from D-dimensional input/observable/“feature” space to one dimensional output → class labels

\[ y(x): \mathbb{R}^n \rightarrow \mathbb{R}: \]

\[ y(x): \text{“test statistic” in D-dimensional space of input variables} \]

\[ \text{distributions of } y(x): \text{PDF}_S(y) \text{ and PDF}_B(y) \]

\[ \text{used to set the selection cut!} \]

\[ \rightarrow \text{efficiency and purity} \]

\[ y(x)=\text{const: surface defining the decision boundary.} \]

\[ \text{overlap of PDF}_S(y) \text{ and PDF}_B(y) \rightarrow \text{separation power, purity} \]
Classification ↔ Regression

Classification:
- Each event, if Signal or Background, has “D” measured variables.
- $y(x): \mathbb{R}^D \rightarrow \mathbb{R}$: “test statistic” in D-dimensional space of input variables
- $y(x) =$ const: surface defining the decision boundary.

Regression:
- Each event has “D” measured variables + one function value (e.g. cluster shape variables in the ECAL + particles energy)
- $y(x): \mathbb{R}^D \rightarrow \mathbb{R}$
  → find
- $y(x) =$ const → hyperplanes where the target function is constant
Now, $y(x)$ needs to be build such that it best approximates the target, not such that it best separates signal from bkgr.
Event Classification

\( y(x): \mathbb{R}^n \rightarrow \mathbb{R} \): the mapping from the “feature space” (observables) to one output variable

PDF\(_B(y)\), PDF\(_S(y)\): normalised distribution of \( y=y(x) \) for background and signal events (i.e. the “function” that describes the shape of the distribution)

with \( y=y(x) \) one can also say \( \text{PDF}_B(y(x)), \text{PDF}_S(y(x)) \):

Probability densities for background and signal

now let’s assume we have an unknown event from the example above for which \( y(x) = 0.2 \)

\( \rightarrow \text{PDF}_B(y(x)) = 1.5 \) and \( \text{PDF}_S(y(x)) = 0.45 \)

let \( f_S \) and \( f_B \) be the fraction of signal and background events in the sample, then:

\[
\frac{f_S \text{PDF}_S(y)}{f_S \text{PDF}_S(y) + f_B \text{PDF}_B(y)} = P(C = S | y)
\]

is the probability of an event with measured \( x = \{x_1, \ldots, x_D\} \) that gives \( y(x) \) to be of type signal
Neyman-Pearson Lemma

Likelihood Ratio: $y(x) = \frac{P(x \mid S)}{P(x \mid B)}$

Neyman-Pearson:
The Likelihood ratio used as “selection criterium” $y(x)$ gives for each selection efficiency the best possible background rejection.

i.e. it maximises the area under the “Receiver Operation Characteristics” (ROC) curve

varying $y(x)$>“cut” moves the working point (efficiency and purity) along the ROC curve

how to choose “cut” $\rightarrow$ need to know prior probabilities (S, B abundances)

- measurement of signal cross section: maximum of $S/\sqrt{(S+B)}$ or equiv. $\sqrt{\varepsilon \cdot p}$
- discovery of a signal (typically: $S<<B$): maximum of $S/\sqrt{B}$
- precision measurement:
- trigger selection:
  - high purity (p)
  - high efficiency ($\varepsilon$)
Efficiency or Purity?

if discriminating function \( y(x) = \text{“true likelihood ratio”} \)

\( \Rightarrow \) optimal working point for specific analysis lies somewhere on the ROC curve

\( y(x) \neq \text{“true likelihood ratio”} \) differently, point

\( \Rightarrow \ y'(x) \) might be better for a specific working point than \( y''(x) \) and vice versa

- application of “cost matrix” \( \Rightarrow \) favour one over the other in the training process

  - e.g. cancer diagnostic:
    - consequences of a Type2 error much more severe (death due to missing treatment)
    - compared to Type1 error (unnecessary distress)

\( \text{Type-1 error small} \quad \text{Type-2 error large} \)

\( \text{Type-1 error large} \quad \text{Type-2 error small} \)

"limit" in ROC curve given by likelihood ratio which one of those two is the better??
The previous slide was basically the idea of “Multivariate Analysis” (MVA)
rem: What about “standard cuts”? 

Finding \( y(x) : \mathbb{R}^n \rightarrow \mathbb{R} \)

- given a certain type of model class \( y(x) \)
- in an automatic way using “known” or “previously solved” events
  - i.e. learn from known “patterns”
- such that the resulting \( y(x) \) has good generalization properties when applied to “unknown” events (regression: fits well the target function “in between” the known training events

→ that is what the “machine” is supposed to be doing: supervised machine learning

Of course… there’s no magic, we still need to:

- choose the discriminating variables
- choose the class of models (linear, non-linear, flexible or less flexible)
- tune the “learning parameters” → bias vs. variance trade off
- check generalization properties
- consider trade off between statistical and systematic uncertainties
Event Classification

- Unfortunately, the true probability densities functions are typically unknown:
  → Neyman-Pearsons lemma doesn’t really help us directly

- Monte Carlo simulation or in general cases: set of known (already classified) “events”

- 2 different ways: Use these “training” events to:
  - estimate the functional form of $p(x|C)$: (e.g. the differential cross section folded with the detector influences) from which the likelihood ratio can be obtained
    → e.g. D-dimensional histogram, Kernel density estimators, …
  - find a “discrimination function” $y(x)$ and corresponding decision boundary (i.e. hyperplane* in the “feature space”: $y(x) = \text{const}$) that optimally separates signal from background
    → e.g. Linear Discriminator, Neural Networks, …

* hyperplane in the strict sense goes through the origin. Here I mean “affine set” to be precise
Nearest Neighbour and Kernel Density Estimator

- estimate probability density $P(x)$ in $D$-dimensional space: “events” distributed according to $P(x)$
- The only thing at our disposal is our “training data”
- Say we want to know $P(x)$ at “this” point “$x$”
- One expects to find in a volume $V$ around point “$x$” $N\int P(x)dx$ events from a dataset with $N$ events
- For the chosen a rectangular volume $\rightarrow$ K-events:

$$K = \sum_{n=1}^{N} k\left(\frac{x - x_n}{h}\right), \quad \text{with} \quad k(u) = \begin{cases} 1, & |u| \leq \frac{1}{2}, i = 1...D \\ 0, & \text{otherwise} \end{cases}$$

$K$ (from the “training data”) $\rightarrow$ estimate of average $P(x)$ in the volume $V$:

$$\int_{V} P(x) dx = K/N$$

- Classification: Determine $PDF_S(x)$ and $PDF_B(x)$
  $\rightarrow$ likelihood ratio as classifier!

Kernel Density estimator of the probability density

$$P(\mathbf{x}) = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{h^D} k\left(\frac{\mathbf{x} - \mathbf{x}_n}{h}\right)$$
Nearest Neighbour and Kernel Density Estimator

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- $K$ (from the “training data”) $\Rightarrow$ estimate of average $P(x)$ in the volume $V$: $\hat{P}(x)dx = K/N$

- **Regression:** If each events with $(x_1, x_2)$ carries a “function value” $f(x_1, x_2)$ (e.g. energy of incident particle) $\Rightarrow$

$$\frac{1}{N} \sum_{i}^{N} f(x_1^i, x_2^i) = \int_V \hat{f}(x_1, x_2) P(\bar{x})d\bar{x}$$

i.e.: the average function value
Nearest Neighbour and Kernel Density Estimator

- estimate probability density \( P(x) \) in \( D \)-dimensional space: “events” distributed according to \( P(x) \)

- The only thing at our disposal is our “training data”

- Say we want to know \( P(x) \) at “this” point “\( x \)"

- One expects to find in a volume \( V \) around point “\( x \)"
  \( \frac{N \times \int P(x)dx}{V} \) events from a dataset with \( N \) events

- For the chosen rectangular volume
  \( \rightarrow \) \( K \)-events:

\[
K = \sum_{n=1}^{N} k \left( \frac{x - x_n}{h} \right), \quad \text{with} \quad k(u) = \begin{cases} 
1, & |u_i| \leq \frac{1}{2}, i = 1 \ldots D \\
0, & \text{otherwise}
\end{cases}
\]

- determine \( K \) from the “training data” with signal and background mixed together

\( \rightarrow \) \( k\text{NN} \): \( k \)-Nearest Neighbours
relative number events of the various classes amongst the \( k \)-nearest neighbours

\[
y(x) = \frac{n_s}{K}
\]
Kernel Density Estimator

- Parzen Window: “rectangular Kernel” → discontinuities at window edges
  → smoother model for \( P(x) \) when using smooth Kernel Functions: e.g. Gaussian

\[
P(x) = \frac{1}{N} \sum_{n=1}^{N} \frac{1}{(2\pi h^2)^{1/2}} \exp \left( \frac{\|x - x_n\|^2}{2h^2} \right)
\]

\[\leftrightarrow\] probability density estimator

- place a “Gaussian” around each “training data point” and sum up their contributions at arbitrary points “\( x \)” \( \rightarrow P(x) \)
- \( h \): “size” of the Kernel \( \rightarrow \) “smoothing parameter”
- there is a large variety of possible Kernel functions

![Individual kernels and averaged kernels](image)
Kernel Density Estimator

\[ P(\mathbf{x}) = \frac{1}{N} \sum_{n=1}^{N} K_h(\mathbf{x} - \mathbf{x}_n) \] : a general probability density estimator using kernel K

- \( h \): “size” of the Kernel  \( \rightarrow \) “smoothing parameter”

- chosen size of the “smoothing-parameter” \( \rightarrow \) more important than kernel function

- \( h \) too small: overtraining
- \( h \) too large: not sensitive to features in \( P(\mathbf{x}) \)

for Gaussian kernels, the optimum in terms of MISE (mean integrated squared error) is given by: \( h_{\mathbf{x}_i} = (4/(3N))^{1/5} \sigma_{\mathbf{x}_i} \) with \( \sigma_{\mathbf{x}_i} = \text{RMS in variable } \mathbf{x}_i \)

- a drawback of Kernel density estimators:
  Evaluation for any test events involves ALL TRAINING DATA \( \rightarrow \) typically very time consuming

- binary search trees (i.e. Kd-trees) are typically used in kNN methods to speed up searching

\[ nP(\mathbf{x}) = \sum_{i=1}^{N} \frac{1}{N} K_h(\mathbf{x} - \mathbf{x}_n) \]

(Christopher M. Bishop)
“Curse of Dimensionality”

We all know:
Filling a D-dimensional histogram to get a mapping of the PDF is typically unfeasable due to lack of Monte Carlo events.

Shortcoming of nearest-neighbour strategies:

- in higher dimensional classification/regression cases the idea of looking at “training events” in a reasonably small “vicinity” of the space point to be classified becomes difficult:

consider: total phase space volume $V=1^D$
for a cube of a particular fraction of the volume:

$\text{edge length} = \left(\text{fraction of volume}\right)^{1/D}$

- In 10 dimensions: in order to capture 1% of the phase space
  $\rightarrow 63\%$ of range in each variable necessary $\rightarrow$ that’s not “local” anymore..

$\rightarrow$Therefore we still need to develop all the alternative classification/regression techniques
Naïve Bayesian Classifier  
“often called: (projective) Likelihood”

while Kernel methods or Nearest Neighbour classifiers try to estimate the full D-dimensional joint probability distributions

If correlations between variables are weak: \( P(x) \approx \prod_{i=0}^{D} P_i(x) \)

product of marginal PDFs  (1-dim “histograms”)

- One of the first and still very popular MVA-algorithm in HEP
- rather than making hard cuts on individual variables, allow for some “fuzzyness”: one very signal like variable may counterweigh another less signal like variable
- This is about the optimal method in the absence of correlations

\[
y(x_{\text{PDE,k}_{\text{event}}}) = \prod_{i \in \{\text{variables}\}} P_{i_{\text{signal}}}(x_{i,k_{\text{event}}}) \frac{\sum_{C \in \{\text{classes}\}} \prod_{i \in \{\text{variables}\}} P_C^i(x_{i,k_{\text{event}}})}{} \]
Naïve Bayesian Classifier
“often called: (projective) Likelihood”

How parameterise the 1-dim PDFs ??

- event counting (histogramming)
  - Automatic, unbiased, but suboptimal
- parametric (function) fitting
  - Difficult to automate for arbitrary PDFs
- nonparametric fitting (i.e. splines, kernel)
  - Easy to automate, can create artefacts/suppress information

example: original (underlying) distribution is Gaussian

- If the correlations between variables is really negligible, this classifier is “perfect” (simple, robust, performing)
- If not, you seriously lose performance 😞
- How can we “fix” this ?
What if there are correlations?

- Typically correlations are present: 
  \[ C_{ij} = \text{cov}(x_i, x_j) = E(x_i x_j) - E(x_i)E(x_j) \neq 0 \ (i \neq j) \]

\[ \rightarrow \text{pre-processing: choose set of linear transformed input variables for which } C_{ij} = 0 \ (i \neq j) \]
Decorrelation

- Determine *square-root* $C'$ of correlation matrix $C$, *i.e.*, $C = C'C$
  - compute $C'$ by diagonalising $C$: $D = S^T CS \Rightarrow C' = S\sqrt{D}S^T$
  - transformation from original $(x)$ in de-correlated variable space $(x')$ by: $x' = C'^{-1}x$

Attention. This is able to eliminate only linear correlations!!
Decorrelation

- Determine *square-root* $C'$ of correlation matrix $C$, i.e., $C = C'C$
  - compute $C'$ by diagonalising $C$: $D = S^TCS \Rightarrow C' = S\sqrt{D}S^T$
  - transformation from original $(x)$ in de-correlated variable space $(x')$ by: $x' = C' - 1x$

Attention. This is able to eliminate only linear correlations!!
Decorrelation: Principal Component Analysis

- PCA is typically used to:
  - reduce the dimensionality of a problem
  - find the most dominant features in your distribution by transforming

- The eigenvectors of the covariance matrix with the largest eigenvalues correspond to the dimensions that have the strongest correlation in the data set. Along these axis the variance is largest
  - sort the eigenvectors according to their eigenvalues

- Dataset is transformed in variable space along these eigenvectors
  - Along the “first” dimension the data show the largest “features”, the smallest features are found in the “last” dimension.

\[ x_k^{PC}(i_{\text{event}}) = \sum_{v \in \{\text{variables}\}} [x_v(i_{\text{event}}) - \bar{x}_v] \cdot v_v^{(k)}, \quad \forall k \in \{\text{variables}\} \]

- Matrix of eigenvectors V obey the relation: \( C \cdot V = D \cdot V \) → PCA eliminates correlations!

Principle Component (PC) of variable \( k \)

Sample means

Eigenvector

Correlation matrix

Diagonalised square root of C
How to Apply the Pre-Processing Transformation?

In general: the decorrelation for signal and background variables is different however: for a “test event” we don’t know beforehand if it is signal or background. ? What do we do?

→ for likelihood ratio, decorrelate signal and background independently

\[
y_L(i_{\text{event}}) = \frac{\prod_{k \in \{\text{variables}\}} p_k^S(x_k(i_{\text{event}}))}{\prod_{k \in \{\text{variables}\}} p_k^S(x_k(i_{\text{event}})) + \prod_{k \in \{\text{variables}\}} p_k^B(x_k(i_{\text{event}}))}
\]

signal transformation

\[
y_{L,\text{trans}}(i_{\text{event}}) = \frac{\prod_{k \in \{\text{variables}\}} p_k^S(\hat{x}_k(i_{\text{event}}))}{\prod_{k \in \{\text{variables}\}} p_k^S(\hat{x}_k(i_{\text{event}})) + \prod_{k \in \{\text{variables}\}} p_k^B(\hat{x}_k(i_{\text{event}}))}
\]

background transformation

→ for other estimators, one needs to decide on one of the two… (or decorrelate on a mixture of signal and background events)
Decorrelation at Work

- Example: linear correlated Gaussians → decorrelation works to 100%
  → 1-D Likelihood on decorrelated sample give best possible performance
  → compare also the effect on the MVA-output variable!

Correlated variables:

(Note the different scale on the y-axis... sorry)
Limitations of the Decorrelation

- in cases with non-Gaussian distributions and/or nonlinear correlations, the decorrelation needs to be treated with care

- How does linear decorrelation affect cases where correlations between signal and background differ?

![Original correlations](image)

Signal

Background
Limitations of the Decorrelation

- in cases with non-Gaussian distributions and/or nonlinear correlations, the decorrelation needs to be treated with care.

- How does linear decorrelation affect cases where correlations between signal and background differ?

SQRT decorrelation

![Signal](image1)

![Background](image2)
Limitations of the Decorrelation

- in cases with non-Gaussian distributions and/or nonlinear correlations, the decorrelation needs to be treated with care

- How does linear decorrelation affect strongly nonlinear cases?

![Original correlations](image)

- Signal
- Background
Limitations of the Decorrelation

- in cases with non-Gaussian distributions and/or nonlinear correlations, the decorrelation needs to be treated with care

- How does linear decorrelation affect strongly nonlinear cases?

Watch out before you used decorrelation “blindly”!!
Perhaps “decorrelate” only a subspace!
"Gaussian-isation"

- Improve decorrelation by pre-Gaussianisation of variables
  
  - First: transformation to achieve uniform (flat) distribution:
    \[
    x_k^{\text{flat}}(i_{\text{event}}) = \int_{-\infty}^{x_k(i_{\text{event}})} p_k(x'_k) dx'_k, \quad \forall k \in \{\text{variables}\}
    \]
    Rarity transform of variable \(k\)  
    Measured value  
    PDF of variable \(k\)

    The integral can be solved in an unbinned way by event counting, or by creating non-parametric PDFs (see later for likelihood section)

  - Second: make Gaussian via inverse error function:
    \[
    x_k^{\text{Gauss}}(i_{\text{event}}) = \sqrt{2} \cdot \text{erf}^{-1}\left(2x_k^{\text{flat}}(i_{\text{event}}) - 1\right), \quad \forall k \in \{\text{variables}\}
    \]

  - Third: decorrelate (and "iterate" this procedure)
“Gaussian-isation”

We cannot simultaneously “Gaussianise” both signal and background!
"Gaussian-isation"

We cannot simultaneously “Gaussianise” both signal and background!
We cannot simultaneously “Gaussianise” both signal and background!
If non-parametric methods like ‘k-Nearest-Neighbour’ or “Kernel Density Estimators” suffer from
- lack of training data → “curse of dimensionality”
- slow response time → need to evaluate the whole training data for every test event

→ use of parametric models $y(x)$ to fit to the training data

$$ y \ (x = \{x_1, ..., x_D\}) = \sum_{i=0}^{M} w_i h_i(x) $$

**Linear Discriminant:**

i.e. any **linear** function of the input variables:
giving rise to **linear** decision boundaries

$$ y \ (x = \{x_1, ..., x_D\}) = w_0 + \sum_{i=1}^{D} w_i x_i $$

How do we determine the “weights” $w$ that do “best”??

Watch out: these lines are NOT the linear function $y(x)$, but rather the decision boundary given by $y(x) = \text{const.}$
Fisher’s Linear Discriminant

\[ y (x = \{x_1, \ldots, x_D\}) = y (x, w) = w_0 + \sum_{i=1}^{D} w_i x_i \]

How do we determine the “weights” \( w \) that do “best”??

- Maximise the “separation” between the two classes S and B
- minimise overlap of the distribution \( y_S(x) \) and \( y_B(x) \)
  - maximise the distance between the two mean values of the classes
  - minimise the variance within each class

\[ \rightarrow \text{maximise} \quad J(\mathbf{w}) = \frac{(E(y_B) - E(y_S))^2}{\sigma_{y_B}^2 + \sigma_{y_S}^2} = \frac{\mathbf{w}^T \mathbf{Bw}}{\mathbf{w}^T \mathbf{Ww}} = \frac{\text{"in between" variance}}{\text{"within" variance}} \]

\[ \nabla_w J (\mathbf{w}) = 0 \Rightarrow \mathbf{w} \propto \mathbf{W}^{-1} \langle \bar{x}_S \rangle - \langle \bar{x}_B \rangle \quad \text{the Fisher coefficients} \]

note: these quantities can be calculated from the training data
Linear Discriminant and non linear correlations

assume the following non-linear correlated data:

- the Linear discriminant obviously doesn’t do a very good job here:

- Of course, these can easily be de-correlated:
  ➔ here: linear discriminator works perfectly on de-correlated data

$$\text{var}\ 0' = \sqrt{\text{var}\ 0^2 + \text{var}\ 1^2}$$

$$\text{var}\ 1' = \tan\left(\frac{\text{var}\ 0}{\text{var}\ 1}\right)$$
Linear Discriminant with Quadratic input:

- A simple extension of the linear decision boundary to "quadratic" decision boundary:
  
  while:
  - \( \text{var0} \)
  - \( \text{var1} \)
  - \( \text{var0} \times \text{var0} \)
  - \( \text{var1} \times \text{var1} \)
  - \( \text{var0} \times \text{var1} \)

  \( \rightarrow \) linear decision boundaries in \( \text{var0}, \text{var1} \)

  \( \rightarrow \) quadratic decision boundaries in \( \text{var0}, \text{var1} \)

Performance of Fisher Discriminant:

with quadratic input:

- Fisher
- Fisher with decorrelated variables
- Fisher with quadratic input
Linear Discriminant with Quadratic input:

- A simple extension of the linear decision boundary to “quadratic” decision boundary:
  - $\var{var0} \cdot \var{var0}$
  - $\var{var1} \cdot \var{var1}$
  - $\var{var0} \cdot \var{var1}$

- Of course, if one “finds out”/”knows” correlations they are best treated explicitly!
  - explicit decorrelation
  - or e.g:
    - Function discriminant analysis (FDA)
      - Fit any user-defined function of input variables requiring that signal events return $\rightarrow 1$ and background $\rightarrow 0$
        - Parameter fitting: Genetics Alg., MINUIT, MC and combinations
        - Easy reproduction of Fisher result, but can add nonlinearities
        - Very transparent discriminator
All classifiers seen so far (KNN, Likelihood, Fisher) could be calculated and didn’t require parameter fitting → the class that “estimate” the underlying PDF rather than a decision boundary.

Other classifiers provide a set of “basis” functions (or model) that need to be optimally adjusted to find the appropriate separating hyperplane/surface.

Let $x \in R^D$ be a random variable (i.e. our observables) and $y(x)$

- a real valued output variable with joint distribution $Pr(x,y)$ → regression
- $y$ value → 1 for signal, $y \to 0$ for background → classification

we are looking for a function $y(x)$ that predicts $y$ given certain input variables: $y(x): R^D \to R$:

Loss function: $L(y, y(x))$ penalizes errors made in the prediction:

- $EPE(y(x)) = E(y - y(x))^2$ squared error loss, typical “loss function” used in regression
- by conditioning on “x” we can write:
- $EPE(y(x)) = E(|y-y(x)|)$ misclassification error, typical “loss function” for classification problems
naturally, if we want to go to “arbitrary” non-linear decision boundaries, \( y(x) \) needs to be constructed in “any” non-linear fashion

\[
y(\vec{x}) = \sum_{i}^{M} \left( w_i h_i(\vec{x}) \right)
\]

- Think of \( h_i(x) \) as a set of “basis” functions
- If \( h(x) \) is sufficiently general (i.e. non linear), a linear combination of “enough” basis function should allow to describe any possible discriminating function \( y(x) \)

K. Weierstrass Theorem: proves just that previous statement.

Imagine you chose do the following:

\[
y(x) = \sum_{i}^{M} w_{0i} A \left( w_{i0} + \sum_{j=1}^{D} w_{ij} x_j \right)
\]

\( y(x) = \) a linear combination of non linear functions of linear combination of the input data

\[
A(x) = \frac{1}{1+e^{-x}}:
\]

the sigmoid function

Ready is the Neural Network
Now we “only” need to find the appropriate “weights” \( w \)
Neural Networks: Multilayer Perceptron MLP

But before talking about the weights, let's try to “interpret” the formula as a Neural Network:

- Nodes in hidden layer represent the “activation functions” whose arguments are linear combinations of input variables $\rightarrow$ non-linear response to the input
- The output is a linear combination of the output of the activation functions at the internal nodes
- Input to the layers from preceding nodes only $\rightarrow$ feed forward network (no backward loops)
- It is straightforward to extend this to “several” input layers

$$y(x) = \sum_{i}^{M} w_{0i} A \left( w_{i0} + \sum_{j=1}^{D} w_{ij} \cdot x_{j} \right)$$

"Activation" function e.g. sigmoid:

$$A(x) = \left(1 + e^{-x}\right)^{-1}$$

or tanh or …
Neural Networks: Multilayer Perceptron MLP

But before talking about the weights, let’s try to “interpret” the formula as a Neural Network:

\[
y(x) = \sum_{i}^{M} w_{0i} A \left( w_{i0} + \sum_{j=1}^{D} w_{ij} \cdot x_j \right)
\]

“Activation” function e.g. sigmoid:
\[A(x) = \left(1 + e^{-x}\right)^{-1}\]
or tanh or …

Nodes \(\rightarrow\) Neurons
Links (weights) \(\rightarrow\) Synapses

Neural network: try to simulate reactions of a brain to certain stimulus (input data)
Neural Network Training

idea: using the “training events” adjust the weights such, that

- $y(x) \rightarrow 0$ for background events
- $y(x) \rightarrow 1$ for signal events

how do we adjust?

- minimize Loss function:

$$L(w) = \sum_{i}^{\text{events}} (y(x_i) - y(C))^2$$

where

$$y(C) = \begin{cases} 
1 & \text{for } C = \text{signal} \\
0 & \text{for } C = \text{backgr.} 
\end{cases}$$

- $y(x)$ is a very “wiggly” function with many local minima. A global overall fit in the many parameters is possible but not the most efficient method to train neural networks

- back propagation (learn from experience, gradually adjust your perception to match reality)
- online learning (learn event by event -- not only at the end of your life from all experience)
Neural Network Training
back-propagation and online learning

- start with random weights
- adjust weights in each step a bit in the direction of the steepest descend of the “Loss”-function $L$

$$ w^{n+1} = w^n + \eta \cdot \nabla_w L(w) \quad \eta = \text{learning rate} \quad L(w) = (y(x_i) - y(C))^2 $$

- for weights connected to output nodes
  \[
  \frac{\partial L}{\partial w_{0i}} = (y(x) - y(C)) A \left( w_{i0} + \sum_{j=1}^{D} w_{ij} \cdot x_j \right)
  \]

- for weights connected to output nodes
  ... a bit more complicated formula

- note: all these gradients are easily calculated from the training event

- training is repeated $n$-times over the whole training data sample. how often ??

for online learning, the training events should be mixed randomly, otherwise you first steer in a wrong direction from which it is afterwards hard to get out again!
Bias ↔ Variance Trade off

- High bias, low variance
- Low bias, high variance

Model Complexity vs. Prediction Error

- Training sample
- Test sample

Overtraining!

- Save Monte Carlo training events using “Cross Validation” to optimize tuning parameters ($\alpha$) in the algorithm:
  - Divide the data sample into say 5 sub-sets
  - Train 5 times the same classifier (for a given tune parameter $\alpha$=e.g. #training cycles, size of kernel) → for each event in the training you have a statistically independent classifier
  - Repeat with varied $\alpha$ to find optimal value of $\alpha$ → train classifier on full sample using this $\alpha$

Too bad 😞, cross validation it is still NOT implemented in TMVA
Support Vector Machines

- If Neural Networks are complicated by finding the proper optimum “weights” for best separation power by “wiggly” functional behaviour of the piecewise defined separation hyperplane.

- If KNN (multidimensional likelihood) suffers disadvantage that calculating the MVA-output of each test event involves evaluation of ALL training events.

- If Boosted Decision Trees in theory are always weaker than a perfect Neural Network.

→ Try to get the best of all worlds…
Support Vector Machine

- There are methods to create linear decision boundaries using only measures of distances (= inner (scalar) products)
  - leads to quadratic optimisation problem

- The decision boundary in the end is defined only by training events that are closest to the boundary

- We’ve seen that variable transformations, i.e moving into a higher dimensional space (i.e. using \( \text{var1} \times \text{var1} \) in Fisher Discriminant) can allow you to separate with linear decision boundaries non linear problems

- Support Vector Machine
Support Vector Machines

- Find hyperplane that best separates signal from background
  - Linear decision boundary
  - Best separation: maximum distance (margin) between closest events (support) to hyperplane
  - If data non-separable add \textit{misclassification cost} parameter $C \cdot \sum_i \xi_i$ to minimisation function
- Solution of largest margin depends only on inner product of support vectors (distances)
  - \textit{quadratic minimisation problem}
Support Vector Machines

- Find hyperplane that best separates signal from background
  - Best separation: maximum distance (margin) between closest events (support) to hyperplane
  - Linear decision boundary
  - If data non-separable add misclassification cost parameter \( C \cdot \sum \xi_i \) to minimisation function
  - Solution depends only on inner product of support vectors \( \rightarrow \) quadratic minimisation problem

- Non-linear cases:
  - Transform variables into higher dimensional feature space where again a linear boundary (hyperplane) can separate the data
Support Vector Machines

- Find hyperplane that best separates signal from background
  - Best separation: maximum distance (margin) between closest events (support) to hyperplane
  - Linear decision boundary
  - If data non-separable add misclassification cost parameter $C \cdot \sum \xi_i$ to minimisation function
- Solution depends only on inner product of support vectors $\rightarrow$ quadratic minimisation problem
  - Non-linear cases:
    - Transform variables into higher dimensional feature space where again a linear boundary (hyperplane) can separate the data
    - Explicit transformation doesn’t need to be specified. Only need the “scalar product” (inner product) $x \cdot x \rightarrow \Phi(x) \cdot \Phi(x)$.
      - certain Kernel Functions can be interpreted as scalar products between transformed vectors in the higher dimensional feature space. e.g.: Gaussian, Polynomial, Sigmoid
    - Choose Kernel and fit the hyperplane using the linear techniques developed above
      - Kernel size parameter typically needs careful tuning! (Overtraining!)
Support Vector Machines

SVM: the Kernel size parameter: example: Gaussian Kernels

- Kernel size ($\sigma$ of the Gaussian) choosen too large: $\rightarrow$ not enough “flexibility” in the underlying transformation

- Kernel size ($\sigma$ of the Gaussian) choosen properly for the given problem

colour code:
Red $\rightarrow$ large signal probability:
Boosted Decision Trees

- **Decision Tree:** Sequential application of cuts splits the data into nodes, where the final nodes (leaves) classify an event as **signal** or **background**.
Boosted Decision Trees

- **Decision Tree**: Sequential application of cuts splits the data into nodes, where the final nodes (leafs) classify an event as **signal** or **background**

- Used since a long time in general “data-mining” applications, less known in (High Energy) Physics (although very similar to “simple Cuts”)
  - easy to interpret, visualised
  - independent of monotonous variable transformations, immune against outliers
  - weak variables are ignored (and don’t (much) deteriorate performance)

- Disadvantage → very sensitive to statistical fluctuations in training data

- **Boosted Decision Trees (1996)**: combine a whole forest of Decision Trees, derived from the same sample, e.g. using different event weights.
  - overcomes the stability problem
  - became popular in HEP since MiniBooNE, B.Roe et al., NIM 543(2005)
Growing a Decision Tree

- start with training sample at the root node
- split training sample at node into two, using a cut in the one variable that gives best separation gain
- continue splitting until:
  - minimal #events per node
  - maximum number of nodes
  - maximum depth specified
  - (a split doesn’t give a minimum separation gain) → not a good idea → see “chessboard”
- leaf-nodes classify S,B according to the majority of events or give a S/B probability
- Why no multiple branches (splits) per node?
  → Fragments data too quickly; also: multiple:
- What about multivariate splits?
  → time consuming
  → other methods more adapted for such correlations
Separation Gain

- What do we mean by “best separation gain”?

- define a measure on how mixed S and B in a node are:
  - Gini-index: (Corrado Gini 1912, typically used to measure income inequality)
    \[ p (1-p) : p=\text{purity} \]
  - Cross Entropy:
    \[ -(p \ln p + (1-p) \ln(1-p)) \]
  - Misidentification:
    \[ 1-\max(p,1-p) \]

- difference in the various indices are small, most commonly used: Gini-index

separation gain: e.g. \[ N_{\text{Parent}} \times Gini_{\text{Parent}} - N_{\text{left}} \times Gini_{\text{LeftNode}} - N_{\text{right}} \times Gini_{\text{RightNode}} \]

- Choose amongst all possible variables and cut values the one that maximised the this.
Decision Tree Pruning

- One can continue node splitting until all leaf nodes are basically pure (using the training sample)
  - obviously: that’s overtraining

- Two possibilities:
  - stop growing earlier
    - generally not a good idea, useless splits might open up subsequent useful splits
  - grow tree to the end and “cut back”, nodes that seem statistically dominated:
    - pruning

- e.g. Cost Complexity pruning:
  - assign to every sub-tree, $T \ C(T, \alpha)$:
  - find subtree $T$ with minimal $C(T, \alpha)$ for given $\alpha$
  - prune up to value of $\alpha$ that does not show overtraining in the test sample

\[
C(T, \alpha) = \sum_{\text{leaves events in leaf}} \sum_{\text{events in leaf}} |y(x) - y(C)| + \alpha N_{\text{leaf nodes}}
\]

Loss function

regularisation

cost parameter
“Real life” example of an optimally pruned Decision Tree:

- Pruning algorithms are developed and applied on individual trees
- Optimally pruned single trees are not necessarily optimal in a forest!
Boosting

Training Sample → re-weight → classifier $C^{(0)}(x)$

Weighted Sample → re-weight → classifier $C^{(1)}(x)$

Weighted Sample → re-weight → classifier $C^{(2)}(x)$

Weighted Sample → re-weight → classifier $C^{(3)}(x)$

Weighted Sample → classifier $C^{(m)}(x)$

$y(x) = \sum_{i=1}^{N_{\text{Classifier}}} w_i C^{(i)}(x)$
Adaptive Boosting (AdaBoost)

- AdaBoost re-weights events misclassified by previous classifier by:
  \[ \frac{1 - f_{\text{err}}}{f_{\text{err}}} \]
  with:
  \[ f_{\text{err}} = \frac{\text{misclassified events}}{\text{all events}} \]

- AdaBoost weights the classifiers also using the error rate of the individual classifier according to:
  \[
y(x) = \sum_{i}^{N_{\text{Classifier}}} \log \left( \frac{1 - f_{\text{err}}^{(i)}}{f_{\text{err}}^{(i)}} \right) C^{(i)}(x)
  \]
AdaBoost: A simple demonstration

The example: (somewhat artificial...but nice for demonstration):
- Data file with three “bumps”
- Weak classifier (i.e. one single simple “cut” ↔ decision tree stumps)

Two reasonable cuts:

a) \( \text{Var0} > 0.5 \Rightarrow \varepsilon_{\text{signal}} = 66\% \quad \varepsilon_{\text{bkg}} \approx 0\% \quad \text{misclassified events in total 16.5\%} \)

b) \( \text{Var0} < -0.5 \Rightarrow \varepsilon_{\text{signal}} = 33\% \quad \varepsilon_{\text{bkg}} \approx 0\% \quad \text{misclassified events in total 33\%} \)

the training of a single decision tree stump will find “cut a)”
AdaBoost: A simple demonstration

The first “tree”, choosing cut a) will give an error fraction: err = 0.165

→ before building the next “tree”: weight wrong classified training events by \((1 - \text{err}/\text{err})\) ≈ 5

→ the next “tree” sees essentially the following data sample:

The combined classifier: Tree1 + Tree2

the (weighted) average of the response to a test event from both trees is able to separate signal from background as good as one would expect from the most powerful classifier

.. and hence will chose: “cut b)”: Var0 < -0.5
AdaBoost: A simple demonstration

Only 1 tree “stump”

Only 2 tree “stumps” with AdaBoost
Gradient Boost

- Gradient Boost is a way to implement “boosting” with arbitrary “loss functions” by approximating “somehow” the gradient of the loss function

- AdaBoost: Exponential loss $\exp(-y_0y(\alpha, x))$ → theoretically sensitive to outliers

- Binomial log-likelihood loss $\ln(1 + \exp(-2y_0y(\alpha, x)))$ → more well behaved loss function, (the corresponding “GradientBoost” is implemented in TMVA)
Bagging and Randomised Trees

other classifier combinations:

- **Bagging:**
  - combine trees grown from “bootstrap” samples
  - (i.e re-sample training data with replacement)

- **Randomised Trees:** *(Random Forest: trademark L.Breiman, A.Cutler)*
  - combine trees grown with:
    - random bootstrap (or subsets) of the training data only
    - consider at each node only a random subsets of variables for the split
    - NO Pruning!

- These combined classifiers work surprisingly well, are very stable and almost perfect “out of the box” classifiers
Another plot showing how important it is to choose good “tuning parameters”
The Schachbrett Toy (chess board)

- Performance achieved without parameter adjustments: nearest Neighbour and BDTs are best “out of the box”
- After some parameter tuning, also SVM und ANN(MLP) perform

Event Distribution

- Events weighted by SVM response

![Event Distribution Diagram]
Generalised Classifier Boosting

- Principle (just as in BDT): multiple training cycles, each time wrongly classified events get a higher event weight

\[
y(x) = \sum_{i}^{N_{\text{Classifier}}} \log \left( \frac{1 - f_{\text{err}}^{(i)}}{f_{\text{err}}^{(i)}} \right) C^{(i)}(x)
\]

Response is weighted sum of each classifier response

Boosting will be interesting especially for Methods like Cuts, MLP, and SVM
AdaBoost On a linear Classifier (e.g. Fisher)

J. Sochman, J. Matas, cmp.felk.cvut.cz

Start with a problem for which a linear classifier is weak:

\[ t = 1 \]

\[ t = 3 \]
AdaBoost On a linear Classifier (e.g. Fisher)
Learning with Rule Ensembles

- Following RuleFit approach by Friedman-Popescu

- Model is linear combination of rules, where a rule is a sequence of cuts

\[
y_{RF}(\vec{x}) = a_0 + \sum_{m=1}^{M_R} a_m r_m(\vec{x}) + \sum_{k=1}^{n_R} b_k \hat{x}_k
\]

The problem to solve is

- Create rule ensemble: use forest of decision trees
- Fit coefficients \( a_m, b_k \): gradient direct regularization minimising Risk (Friedman et al.)

- Pruning removes topologically equal rules" (same variables in cut sequence)
See some Classifiers at Work
Toy Examples: Linear-, Cross-, Circular Correlations

- Illustrate the behaviour of linear and nonlinear classifiers

Linear correlations (same for signal and background)

Linear correlations (opposite for signal and background)

Circular correlations (same for signal and background)
How well do the classifier resolve the various correlation patterns?

- **Linear correlations** (same for signal and background)
- **Cross-linear correlations** (opposite for signal and background)
- **Circular correlations** (same for signal and background)
Final Classifier Performance

- Background rejection versus signal efficiency curve:
Stability with Respect to Irrelevant Variables

- Toy example with 2 discriminating and 4 non-discriminating variables?
Stability with Respect to Irrelevant Variables

- Toy example with 2 discriminating and 4 non-discriminating variables?

![Graph of Background rejection versus Signal efficiency]

- Use only two discriminant variables in classifiers.

MVA Methods:
- Fisher
- MLP
- Likelihood
- PDERS
- RuleFit
- BDT
Stability with Respect to Irrelevant Variables

- Toy example with 2 discriminating and 4 non-discriminating variables?

![Graph showing background rejection versus signal efficiency]

- Use all discriminant variables in classifiers

MVA Method:
- Fisher
- MLP
- Likelihood
- RuleFit
- BDT
- PDERS
More Visualisation
(following example taken from M. Schmelling)

A simple K-short selection:

- consider all opposite tracks with $450\text{GeV} < M_{\pi^{-}\pi^{+}} < 550\text{GeV}$
- reconstruct secondary vertex for those tracks
- require the secondary vertex to lie “downstream” $Z_{\text{svtx}} - Z_{\text{pvtx}} > 0$

![Invariant mass spectrum graph]

- $d_1$: impact parameter of the positive track at the primary vertex
- $d_2$: impact parameter of the negative track at the primary vertex
- $d$: impact parameter of the $K_s$ at the primary vertex
- $d_{12}$: distance of closest approach between the tracks
- $d_z$: distance between secondary and primary vertex

![Diagram with impact parameters and distances]
→ criteria

- informative: good separation between signal and background
- well behaved: no “singularities” in the PDFs
  → avoid finetuning in placement of cuts
- independent: every variable contributes new information

❖ variables used in the following

A measure for impact parameter reduction

\[ v_1 = \log_{10} \frac{d_1 d_2}{d} \]

Quality of the secondary vertex

\[ v_2 = \log_{10}(d_{12}) \]

Lifetime of the particle (correlated with \(d_1\) and \(d_2\), hence taken 3rd)

\[ v_3 = \log_{10}(d_3) \]
More Visualisation

(following example taken from M. Schmelling)

- some number of signal and background candidates in training
  ➡️ same statistical power to estimate PDFs
- imbalanced samples in testing
  ➡️ gauge performance under realistic conditions
Visualisation of Decision Boundary

Fisher: classifier contours

MLP: classifier contours

Likelihood: classifier contours

BDT: classifier contours
Visualisation in 3 Variables

- some number of signal and background candidates in training
  - same statistical power to estimate PDFs
- imbalanced samples in testing
  - gauge performance under realistic conditions
Visualisation of Decision Boundary

Fisher-3D: optimal cut contour

MLP-3D: optimal cut contour

Likelihood-3D: optimal cut contour

BDT-3D: optimal cut contour
General Advice for (MVA) Analyses

- There is no magic in MVA-Methods:
  - no need to be too afraid of “black boxes”
  - you typically still need to make careful tuning and do some “hard work”

- The most important thing at the start is finding good observables
  - good separation power between S and B
  - little correlations amongst each other
  - no correlation with the parameters you try to measure in your signal sample!

- Think also about possible combination of variables
  - this may allow you to eliminate correlations
    - rem.: all intelligence that you already put into the system is MUCH better than what the machine will do

- Always apply straightforward preselection cuts and let the MVA only do the rest.

- “Sharp features should be avoided” → numerical problems, loss of information when binning is applied
  - simple variable transformations (i.e. log(variable) ) can often smooth out these areas and allow signal and background differences to appear in a clearer way

- Treat regions in the detector that have different features “independent”
  - can introduce correlations where otherwise the variables would be uncorrelated!
“Categorising” Classifiers

- Multivariate training samples often have distinct sub-populations of data
  - A detector element may only exist in the barrel, but not in the endcaps
  - A variable may have different distributions in barrel, overlap, endcap regions

- Ignoring this dependence creates correlations between variables, which must be learned by the classifier
  - Classifiers such as the projective likelihood, which do not account for correlations, significantly loose performance if the sub-populations are not separated

- Categorisation means splitting the data sample into categories defining disjoint data samples with the following (idealised) properties:
  - Events belonging to the same category are statistically indistinguishable
  - Events belonging to different categories have different properties

- In TMVA: All categories are treated independently for training and application (transparent for user), but evaluation is done for the whole data sample
“Categorising” Classifiers

Let’s try our standard example of 4 Gaussian-distributed input variables:

- Now, “var4” depends on a new variable “eta” (which may not be used for classification)

→ for $|\eta| > 1.3$ the Signal and Background Gaussian means are shifted w.r.t. $|\eta| < 1.3$
“Categorising” Classifiers

- Let’s try our standard example of 4 Gaussian-distributed input variables:
  - Now, “var4” depends on a new variable “eta” (which may not be used for classification)
  - for $|\eta| > 1.3$ the Signal and Background Gaussian means are shifted w.r.t. $|\eta| < 1.3$

The category technique is heavily used in multivariate likelihood fits, eg, RooFit (RooSimultaneousPdf)
Multi-Class Classification

Binary classification: two classes, “signal” and “background”
Multi-Class Classification

Multi-class classification – natural extension for many classifiers
Systematic Errors/Uncertainties
Some Words about Systematic Errors

- Typical worries are:
  - What happens if the estimated “Probability Density” is wrong?
  - Can the Classifier, i.e. the discrimination function $y(x)$, introduce systematic uncertainties?
  - What happens if the training data do not match “reality”

$\Rightarrow$ Any wrong PDF leads to imperfect discrimination function

$$y(x) = \frac{P(x | S)}{P(x | B)}$$

$\Rightarrow$ Imperfect (calling it “wrong” isn’t “right”) $y(x) \Rightarrow$ loss of discrimination power

that’s all!

$\Rightarrow$ classical cuts face exactly the same problem, however:

- in addition to cutting on features that are not correct, now you can also “exploit” correlations that are in fact not correct

- Systematic error are only introduced once “Monte Carlo events” with imperfect modeling are used for
  - efficiency; purity
  - #expected events
  - same problem with classical “cut” analysis
  - use control samples to test MVA-output distribution ($y(x)$)

- Combined variable (MVA-output, $y(x)$) might “hide” problems in ONE individual variable more than if looked at alone $\Rightarrow$ train classifier with few variables only and compare with data
Systematic “Error” in Correlations

- Use as training sample events that have correlations
  - optimize CUTs
  - train an proper MVA (e.g. Likelihood, BDT)

- Assume in “real data” there are NO correlations → SEE what happens!!
Systematic “Error” in Correlations

- Compare “Data” (TestSample) and Monte-Carlo (both taken from the same underlying distribution)
Systematic “Error” in Correlations

- Compare “Data” (TestSample) and Monte-Carlo (both taken from the same underlying distributions that differ by the correlation!!!)

Differences are ONLY visible in the MVA-output plots (and if you’d look at cut sequences....)
Treatment of Systematic Uncertainties

- Is there a strategy however to become ‘less sensitive’ to possible systematic uncertainties
  - i.e. classically: variable that is prone to uncertainties → do not cut in the region of steepest gradient
  - classically one would not choose the most important cut on an uncertain variable

- Try to make classifier less sensitive to “uncertain variables”
  - i.e. re-weight events in training to decrease separation in variables with large systematic uncertainty

(certainly not yet a recipe that can strictly be followed, more an idea of what could perhaps be done)

“Calibration uncertainty” may shift the central value and hence worsen (or increase) the discrimination power of “var4”
Treatment of Systematic Uncertainties

1st Way

Classifier output distributions for signal only
Treatment of Systematic Uncertainties

2nd Way

Classifier output distributions for signal only

MVA_Fisher

MVA_BDT

MVA_Likelihood

MVA_LikelihoodD
What is TMVA

One framework for “all” MVA-techniques, available in ROOT

- Have a common platform/interface for all MVA classification and regression-methods:
- Have common data pre-processing capabilities
- Train and test all classifiers on same data sample and evaluate consistently
- Provide common analysis (ROOT scripts) and application framework
- Provide access with and without ROOT, through macros, C++ executables or python

TMVA is a sourceforge (SF) package for world-wide access

- Home page .................. http://tmva.sf.net/
- SF project page .............. http://sf.net/projects/tmva
- Mailing list .................. http://sf.net/mail/?group_id=152074
- Tutorial TWiki ............... https://twiki.cern.ch/twiki/bin/view/TMVA/WebHome

Integrated and distributed with ROOT since ROOT v5.11/03

- now fully developed within ROOT SVN  →  same release cycles
Currently implemented classifiers and regression methods:
- Rectangular cut optimisation
- Projective and multidimensional likelihood estimator (incl. regression)
- k-Nearest Neighbor algorithm (incl. regression)
- Fisher and H-Matrix discriminants
- Function discriminant
- Artificial neural networks (3 multilayer perceptron implementations) (incl. regression)
- Boosted/bagged decision trees (incl. regression)
- Rule Fitting
- Support Vector Machine (incl. regression)

Currently implemented data preprocessing stages:
- Decorrelation, Principal Value Decomposition, “Gaussianisation”

Examples for combination methods:
- Boosting, Categorisation, MVA Committees
A typical TMVA analysis consists of two main steps:

1. **Training phase**: training, testing and evaluation of classifiers using data samples with known signal and background composition

2. **Application phase**: using selected trained classifiers to classify unknown data samples

Illustration of these steps with toy data samples
Code Flow for *Training* and *Application* Phases

**User Training Script**
- create
- create `ROOT Target File`
- create `TMVA::Factory`
- execute
- execute `Add Variables`
- execute `API` to `Initialise Training and Test Trees`
- execute `API` to `Book MVA kType, Options`
- execute `API` to `Train MVAs` with `write weight file`
- execute `API` to `Test MVAs`
- execute `API` to `Evaluate MVAs`

**User Application Script**
- create `TMVA::Reader`
- execute
- execute `API` to `Add Variables`
- execute `API` to `Book MVA weight file to read`
- execute `begin event loop`
- update event
- execute
- execute `API` to `Compute MVA`
- execute
- execute `API` to `Compute MVA`
- execute `end event loop`
void TMVC Classification ( )
{
    TFile* outputFile = TFile::Open( "TMVA.root", "RECREATE" );

    TMVA::Factory *factory = new TMVA::Factory( "MVAnalysis", outputFile,"!V" );

    TFile *input = TFile::Open( "tmva_example.root" );

    factory->AddSignalTree ( (TTree*)input->Get("TreeS"), 1.0 );
    factory->AddBackgroundTree ( (TTree*)input->Get("TreeB"), 1.0 );

    factory->AddVariable("var1+var2", 'F');
    factory->AddVariable("var1-var2", 'F');
    factory->AddVariable("var3", 'F');
    factory->AddVariable("var4", 'F');

    factory->PrepareTrainingAndTestTree("", "NSigTrain=3000:NBkgTrain=3000:SplitMode=Random:!V" );

    factory->BookMethod( TMVA::Types::kLikelihood, "Likelihood",
                       "!V!:TransformOutput:Spline=2:NSmooth=5:NAvEvtPerBin=50" );
    factory->BookMethod( TMVA::Types::kMLP, "MLP", "!V!:NCycles=200:HiddenLayers=N+1,N:TestRate=5" );

    factory->TrainAllMethods();
    factory->TestAllMethods();
    factory->EvaluateAllMethods();

    outputFile->Close();
    delete factory;
}
A Simple Example for an Application

```cpp
void TMVClassificationApplication() {
  TMVA::Reader *reader = new TMVA::Reader("!Color");
  Float_t var1, var2, var3, var4;
  reader->AddVariable( "var1+var2", &var1 );
  reader->AddVariable( "var1-var2", &var2 );
  reader->AddVariable( "var3", &var3 );
  reader->AddVariable( "var4", &var4 );
  reader->BookMVA( "MLP classifier", "weights/MVAnalysis_MLP.weights.txt" );
  TFile *input = TFile::Open("tmva_example.root");
  TTree* theTree = (TTree*)input->Get("TreeS");

  // … set branch addresses for user TTree
  for (Long64_t ievt=3000; ievt<theTree->GetEntries(); ievt++) {
    theTree->GetEntry(ievt);
    var1 = userVar1 + userVar2;
    var2 = userVar1 - userVar2;
    var3 = userVar3;
    var4 = userVar4;
    Double_t out = reader->EvaluateMVA( "MLP classifier" );
    // do something with it …
  }
  delete reader;
}
```

create Reader

register the variables

book classifier(s)

prepare event loop

compute input variables

calculate classifier output
Data Preparation

- Data input format: ROOT TTree or ASCII
- Selection any subset or combination or function of available variables
- Apply pre-selection cuts (possibly independent for signal and bkg)
- Define global event weights for signal or background input files
- Define individual event weight (use of any input variable present in training data)
- Choose one out of various methods for splitting into training and test samples:
  - Block wise
  - Randomly
  - Periodically (i.e. periodically 3 testing ev., 2 training ev., 3 testing ev, 2 training ev. ….)
  - User defined training and test trees
- Choose preprocessing of input variables (e.g., decorrelation)
MVA Evaluation Framework

- TMVA is not only a collection of classifiers, but an MVA framework.
- After training, TMVA provides ROOT evaluation scripts (through GUI).

Plot all signal (S) and background (B) input variables with and without pre-processing.

Correlation scatters and linear coefficients for S & B.

Classifier outputs (S & B) for test and training samples (spot overtraining).

Classifier Rarity distribution.

Classifier significance with optimal cuts.

B rejection versus S efficiency.

Classifier-specific plots:
- Likelihood reference distributions
- Classifier PDFs (for probability output and Rarity)
- Network architecture, weights and convergence
- Rule Fitting analysis plots

- Visualise decision trees.
Example: Toy Monte Carlo

Data set with 4 linearly correlated Gaussian distributed variables:

<table>
<thead>
<tr>
<th>Rank</th>
<th>Variable</th>
<th>Separation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>var4</td>
<td>0.606</td>
</tr>
<tr>
<td>2</td>
<td>var1+var2</td>
<td>0.182</td>
</tr>
<tr>
<td>3</td>
<td>var3</td>
<td>0.173</td>
</tr>
<tr>
<td>4</td>
<td>var1-var2</td>
<td>0.014</td>
</tr>
</tbody>
</table>
Evaluating the Classifier Training (I)

- Projective likelihood PDFs, MLP training, BDTs, …

average no. of nodes before/after pruning: 4193 / 968
Testing the Classifiers

- Classifier output distributions for independent test sample:
Evaluating the Classifier Training

- Check for overtraining: classifier output for test and training samples ...

- Remark on overtraining
  - Occurs when classifier training has too few degrees of freedom because the classifier has too many adjustable parameters for too few training events
  - Sensitivity to overtraining depends on classifier: e.g., Fisher weak, BDT strong
  - Compare performance between training and test sample to detect overtraining
  - Actively counteract overtraining: e.g., smooth likelihood PDFs, prune decision trees, …
Evaluating the Classifier Training (IV)

- There is no unique way to express the performance of a classifier → several benchmark quantities computed by TMVA
  - Signal eff. at various background effs. (= 1 – rejection) when cutting on classifier output
  - The Separation: \[ \frac{1}{2} \int \frac{(\hat{y}_S(y) - \hat{y}_B(y))^2}{\hat{y}_S(y) + \hat{y}_B(y)} dy \]
  - “Rarity” implemented (background flat): \[ R(y) = \int_{-\infty}^{y} \hat{y}(y') dy' \]
Evaluating the Classifier Training

- Optimal cut for each classifiers …

Determine the optimal cut (working point) on a classifier output

For 1000 signal and 1000 background events the maximum $S/\sqrt{S+B}$ is 25.9327 when cutting at 0.4392
Evaluating the Classifiers Training

--- Fisher : Ranking result (top variable is best ranked)
--- Fisher : ---------------------------------------------
--- Fisher : Rank : Variable : Discr. power
--- Fisher : ---------------------------------------------
--- Fisher : 1 : var4 : 2.175e-01
--- Fisher : 2 : var3 : 1.718e-01
--- Fisher : 3 : var1 : 9.549e-02
--- Fisher : 4 : var2 : 2.841e-02
--- Fisher : ---------------------------------------------

Better variable

How discriminating is a variable?

Classifier correlation and overlap

--- Factory : Inter-MVA overlap matrix (signal):
--- Factory : ----------------------------------------
--- Factory : Likelihood   Fisher
--- Factory : Likelihood:  +1.000  +0.667
--- Factory : Fisher:      +0.667  +1.000
--- Factory : ----------------------------------------

Do classifiers select the same events as signal and background? If not, there is something to gain!
### Evaluating the Classifiers Training
(taken from TMVA output...)

#### Evaluation results ranked by best signal efficiency and purity (area)

<table>
<thead>
<tr>
<th>MVA Methods</th>
<th>Signal efficiency at bkg eff. (error): @B=0.01</th>
<th>@B=0.10</th>
<th>@B=0.30</th>
<th>Area</th>
<th>Separation:</th>
<th>Significance:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fisher</td>
<td>0.268(03)</td>
<td>0.653(03)</td>
<td>0.873(02)</td>
<td>0.882</td>
<td>0.444</td>
<td>1.189</td>
</tr>
<tr>
<td>MLP</td>
<td>0.266(03)</td>
<td>0.656(03)</td>
<td>0.873(02)</td>
<td>0.882</td>
<td>0.444</td>
<td>1.260</td>
</tr>
<tr>
<td>LikelihoodD</td>
<td>0.259(03)</td>
<td>0.649(03)</td>
<td>0.871(02)</td>
<td>0.880</td>
<td>0.441</td>
<td>1.251</td>
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<tr>
<td>PDERS</td>
<td>0.223(03)</td>
<td>0.628(03)</td>
<td>0.861(02)</td>
<td>0.870</td>
<td>0.417</td>
<td>1.192</td>
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<tr>
<td>RuleFit</td>
<td>0.196(03)</td>
<td>0.607(03)</td>
<td>0.845(02)</td>
<td>0.859</td>
<td>0.390</td>
<td>1.092</td>
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<tr>
<td>HMatrix</td>
<td>0.058(01)</td>
<td>0.622(03)</td>
<td>0.868(02)</td>
<td>0.855</td>
<td>0.410</td>
<td>1.093</td>
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<tr>
<td>BDT</td>
<td>0.154(02)</td>
<td>0.594(04)</td>
<td>0.838(03)</td>
<td>0.852</td>
<td>0.380</td>
<td>1.099</td>
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<tr>
<td>CutsGA</td>
<td>0.109(02)</td>
<td>1.000(00)</td>
<td>0.717(03)</td>
<td>0.784</td>
<td>0.000</td>
<td>0.000</td>
</tr>
<tr>
<td>Likelihood</td>
<td>0.086(02)</td>
<td>0.387(03)</td>
<td>0.677(03)</td>
<td>0.757</td>
<td>0.199</td>
<td>0.682</td>
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</table>

Better classifier check for overtraining.
Receiver Operating Characteristics (ROC) Curve

- Smooth background rejection versus signal efficiency curve:
  (from cut on classifier output)

![Graph showing Receiver Operating Characteristics (ROC) Curve with various MVA Methods including Fisher, FDA_MG, MLP, LikelihoodPCA, SVM_Gauss, RuleFit, HMatrix, BDT, PDERS, KNN, CutsGA, and Likelihood. The x-axis represents signal efficiency, and the y-axis represents background rejection. The graph illustrates the trade-off between sensitivity (probability to predict S if true S) and specificity (probability to predict B if true B).]
### Summary of Classifiers and their Properties

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Cuts</th>
<th>Likelihood</th>
<th>PDERS / k-NN</th>
<th>H-Matrix</th>
<th>Fisher</th>
<th>MLP</th>
<th>BDT</th>
<th>RuleFit</th>
<th>SVM</th>
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</tbody>
</table>

The properties of the Function discriminant (FDA) depend on the chosen function $A$. 

---

**Helge Voss**

TMVA Seminar, Lausanne, 12. April 2010 — Data Analysis With TMVA
Summary

- Seen the general idea of MVA’s for Classification and Regression:
- The most important classifiers implemented in TMVA:
  - reconstructing the PDF and use Likelihood Ratio:
    - Nearest Neighbour (Multidimensional Likelihood)
    - Naïve-Bayesian classifier (1dim (projective) Likelihood)
  - fitting directly the decision boundary:
    - Linear discriminant (Fisher)
    - Neuronal Network
    - Support Vector Machine
    - Boosted Decision Tress
- Seen some actual decision boundaries for simple 2D/3D problems
- General analysis advise (for MVAs)
- Systematic errors
  - be as careful as with “cuts” and check against data
- Introduction to TMVA
**Minimisation Techniques**

**Brute force methods: Random Monte Carlo or Grid search**
- Sample entire solution space, and chose solution providing minimum estimator
- Good global minimum finder, but poor accuracy

**Default solution in HEP: Minuit**
- Gradient-driven search, using variable metric, can use quadratic Newton-type solution
- Poor global minimum finder, gets quickly stuck in presence of local minima

**Biology-inspired**
- "Genetic" representation of points in the parameter space
- Uses mutation and "crossover"
- Finds approximately global minima

**Like heating up metal and slowly cooling it down ("annealing")**
Atoms in metal move towards the state of lowest energy while for sudden cooling atoms tend to freeze in intermediate higher energy states → slow "cooling" of system to avoid "freezing" in local solution
Minimisation Techniques

**Brute force methods:** Random Monte Carlo or Grid search

- Sample entire solution space, and chose solution providing minimum estimator
- Good global minimum finder, but poor accuracy

one can also chain minimisers

For example, one can use MC sampling to detect the vicinity of a global minimum, and then use Minuit to accurately converge to it.

**Like heating up metal and slowly cooling it down ("annealing")**

Atoms in metal move towards the state of lowest energy while for sudden cooling atoms tend to freeze in intermediate higher energy states $\rightarrow$ slow “cooling” of system to avoid “freezing” in local solution