

# Unfolding algorithms and RooUnfold

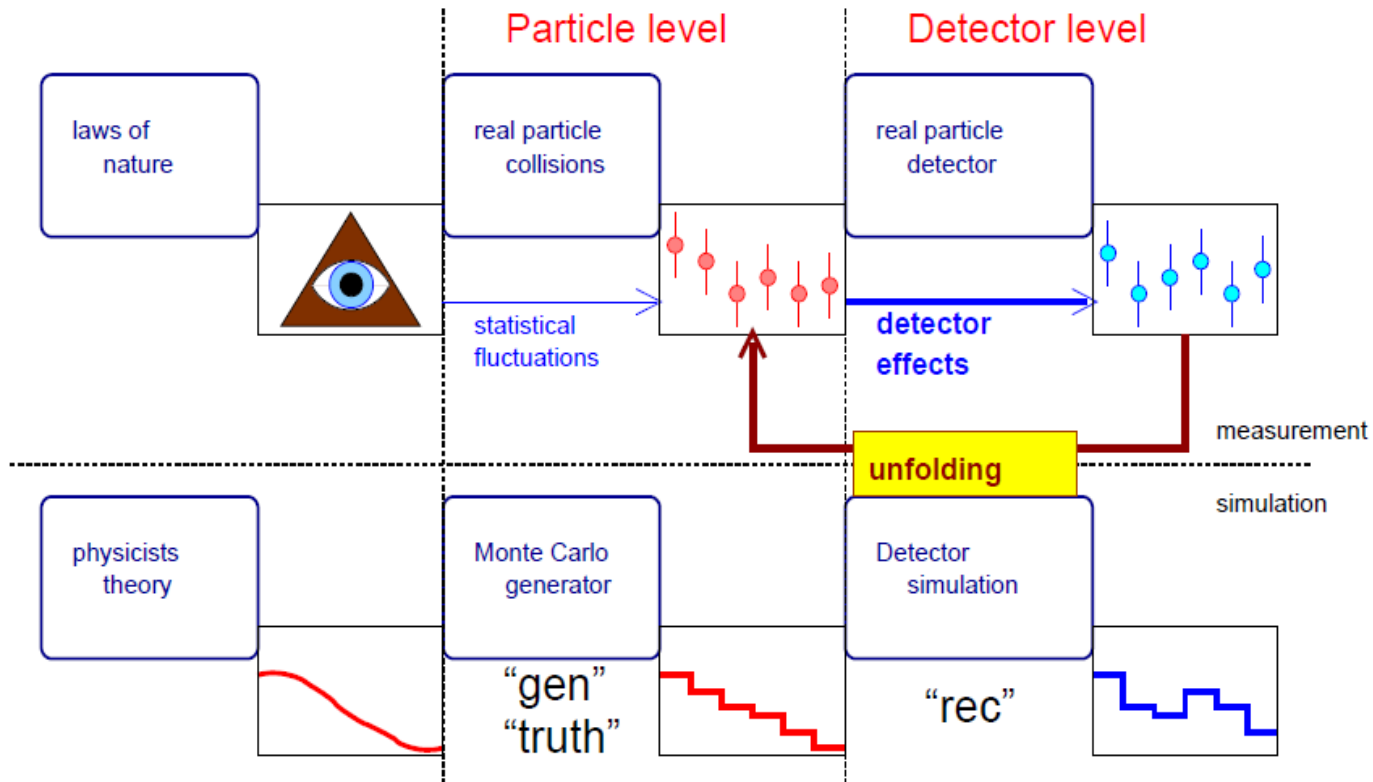
## □ Unfolding algorithms:

- **Extracted from slides by T. Adye at PHYSTAT 2016 and K. Reygers lectures at Heilderbeg Univ.**
- **S. Smitt and D. Britzger, DESY Stat School 2014**
- **S. Schmitt , QCDHS conference in 2016**

# Unfolding

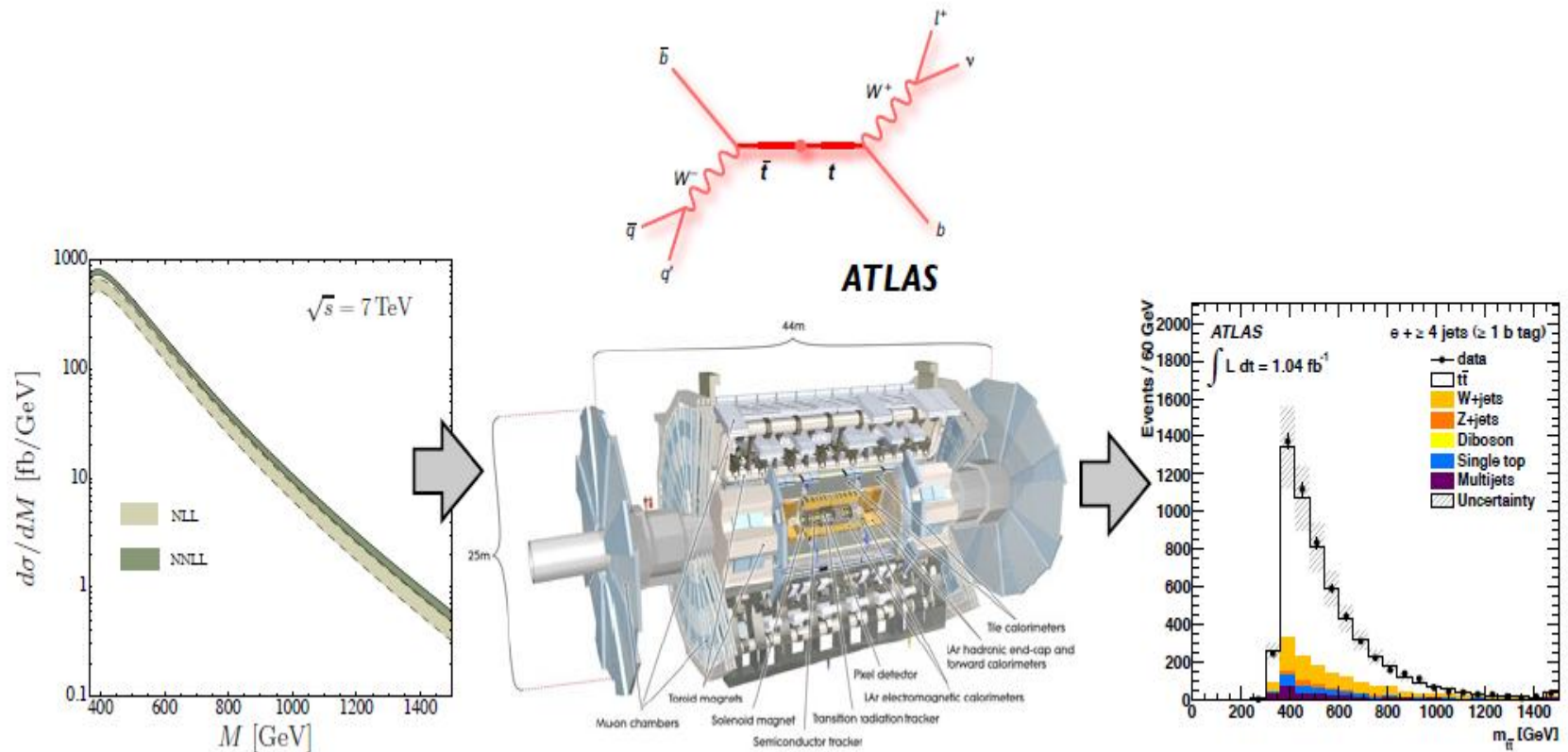
- Unfolding: estimate truth distribution from measurement, distorted by
  - detector effects
  - statistical fluctuations
- truth distribution: cross sections or similar quantities
- Unfolding is also referred to as “correction for detector effects”
- Integral equation of 1<sup>st</sup> kind
$$\int k(x, y)f(y)dy + \delta(x) = g(x)$$
given observations  $g(x)$   
the kernel  $k(x, y)$   
and fluctuations  $\delta(x)$   
estimate the truth  $f(y)$
- $k(x, y)$ : detector effects, background, etc
- $g(x)$  has uncertainties
- $k(x, y)$  has syst. uncertainties  
→ not covered in this talk

# What is unfolding



- Obtain measurements independent of detector effects, using the simulation
- Propagate statistical uncertainties back to particle level
- Require results to be independent of theory assumptions

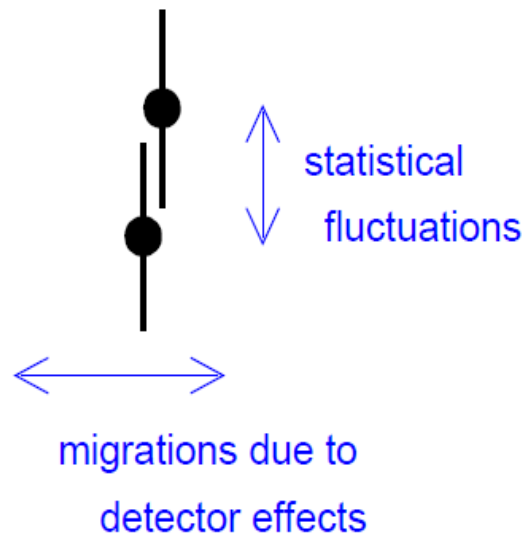
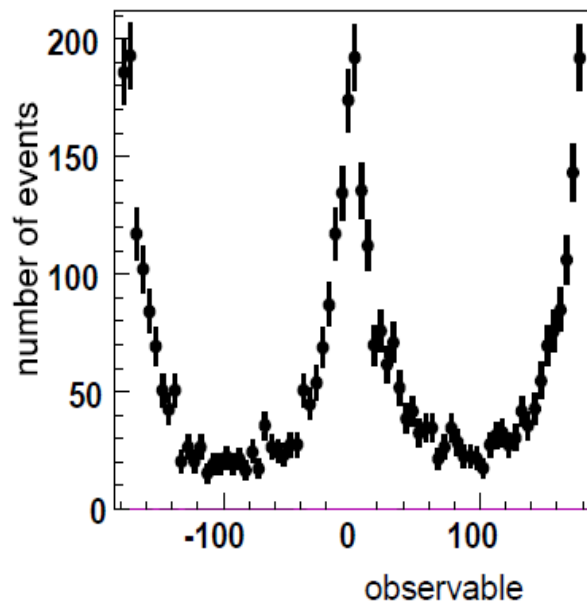
# ATLAS analysis example



G. Aad *et al.* [ATLAS Collaboration], Eur. Phys. J. C **72** (2012) 2039, arXiv:1203.4211 [hep-ex].

# Migrations and stat. fluctuations

Histogram of observed event counts is affected by statistical fluctuations (vertical axis) and detector effects (horizontal axis)



Unfolding: correct for migration effects in the presence of statistical fluctuations

Result: estimator of the "truth" and its covariance matrix (statistical uncertainties)

# Formal definitions

- measurements are given by event (jet, track, ...) counts, grouped in bins  $x_i^{\text{rec}}$
- Bins are defined by regions in phase-space (observables)
- Not covered: unbinned methods
- Detector effects: **detector response matrix A**

$A_{ij}$ : probability that event from truth bin  $j$  is found in rec bin  $i$

Expected number of events in bin  $i$ :  $\mu_i = \sum A_{ij} x_j^{\text{truth}}$

- Statistical fluctuations: **Poisson distribution or Gaussian approximation**

$$P(Y_i = y_i^{\text{rec}}) = \frac{\exp[-\mu_i] (\mu_i)^{y_i}}{y_i!}$$

# Unfolding methods

- Bin-by-bin "correction"
- Matrix methods:
  - Matrix inversion
  - "Bayesian" [D'Agostini 1995]
  - "Iterative" [D'Agostini 1995 iterated]
  - Fraction fit: TUnfold
  - Regularised unfolding: TUnfold

(truth+background) × detector × stat.fluctuations → measurement

Result: estimator of truth ← unfolding algorithm ← measurement

# „Bin-by-bin“

- Assumption: migration effects are “small” so they can be “corrected” using a multiplicative factor
- The factor is determined from MC
- Two variants:
  - Simple bin-by-bin
  - Bin-by-bin with background subtraction



# Bin-by-bin correction factors

- Very simple method:

$$x_i = (y_i - b_i) \frac{N_i^{\text{gen}}}{N_i^{\text{rec}}} \quad \text{Correction factor}$$

$y_i$  : observed in bin  $i$

$b_i$  : expected background in bin  $i$

$N_i^{\text{gen}}$  : MC truth in bin  $i$

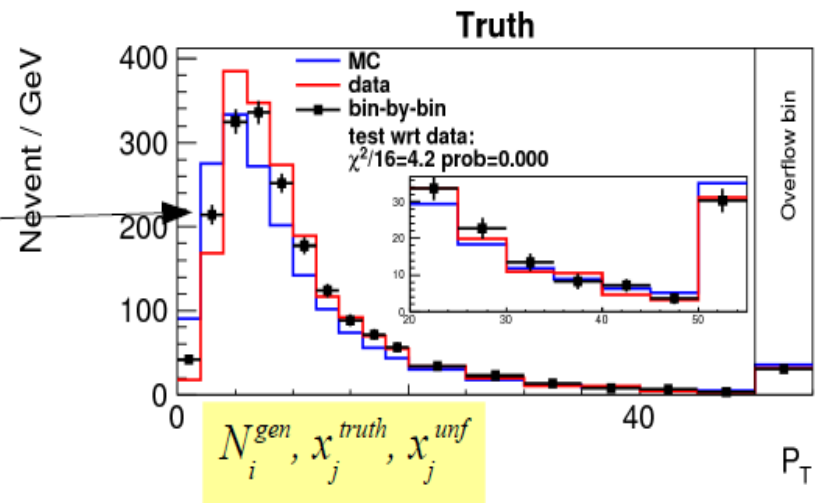
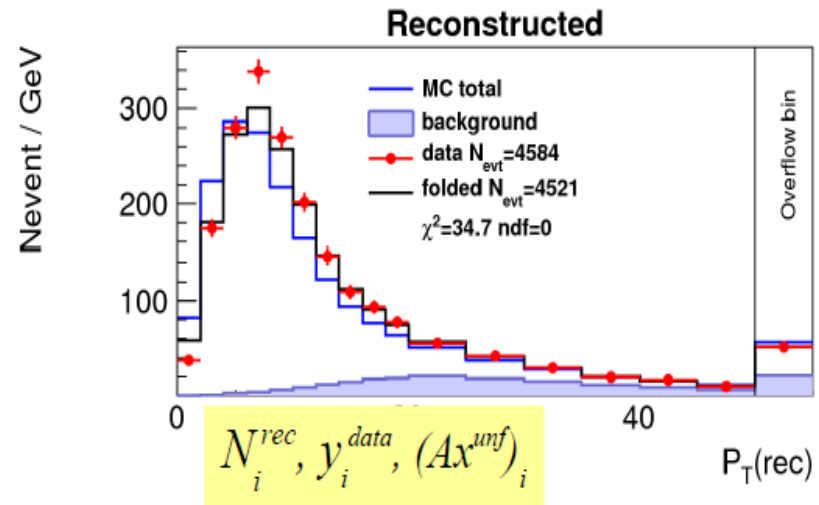
$N_i^{\text{rec}} = \sum_j A_{ij} N_j^{\text{gen}}$  : MC reconstructed in bin  $i$

Results “looks nice”

No statistical bin-to-bin correlations

but

Method is wrong, fails very basic tests



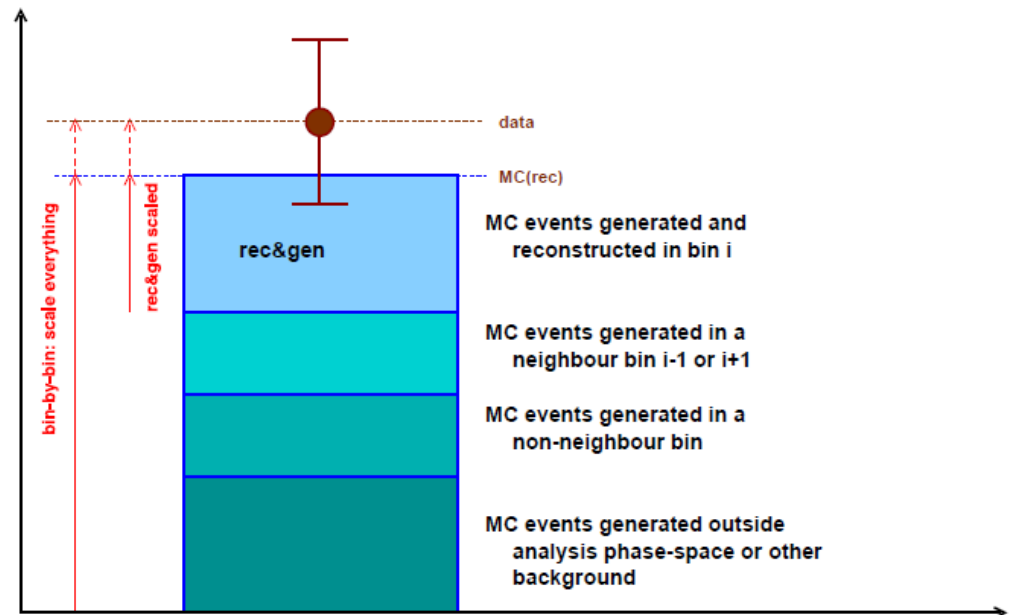
# Simple „Bin-by-bin”: why is it wrong?

- Migrations are additive, while BBB correction is multiplicative → wrong type of correction

$$x_i^{\text{BBB}} = x_i^{\text{gen}} \frac{y_i^{\text{data}}}{y_i^{\text{rec}}}$$

- It should be:

$$\begin{aligned} x_i^{\text{BBBSUB}} &= x_i^{\text{gen}} \frac{y_i^{\text{data}} - (y_i^{\text{rec}} - y_i^{\text{rec\&gen}})}{y_i^{\text{rec\&gen}}} \\ &= x_i^{\text{gen}} \frac{y_i^{\text{data}} - y_i^{\text{rec}} (1 - P_i)}{y_i^{\text{rec}} P_i} \end{aligned}$$



- Relevant quantity: purity

$$P_i = \frac{y_i^{\text{rec\&gen}}}{y_i^{\text{rec}}}$$

# Matrix methods

- All matrix methods are based on the matrix of probabilities:

Expected number of events in bin  $i$ :  $\mu_i = \sum A_{ij} x_j^{\text{truth}}$

- The  $A_{ij}$  are calculated from Monte Carlo

$$A_{ij} = \frac{y_{ij}^{\text{rec,gen}}}{y_j^{\text{gen}}} \text{ and the reconstruction efficiencies are } \varepsilon_j = \sum_i A_{ij}$$

- $A_{ij}$  is normalized to the generated number of events in bin  $j$ , so it is (largely) model independent, only depends on the detector response.

# Matrix inversion

- If the number of bins is equal on gen and rec level:  $A$  is a square matrix

→ invert it

folding equation:  $y = Ax + b$

invert matrix:  $x = A^{-1}(y - b)$

Covariance:  $V_{xx} = A^{-1}V_{yy}(A^{-1})^T$

correlation coefficients:  $\rho_{ij} = \frac{(V_{xx})_{ij}}{\sqrt{(V_{xx})_{ii}(V_{xx})_{jj}}}$

$y$  : measurements

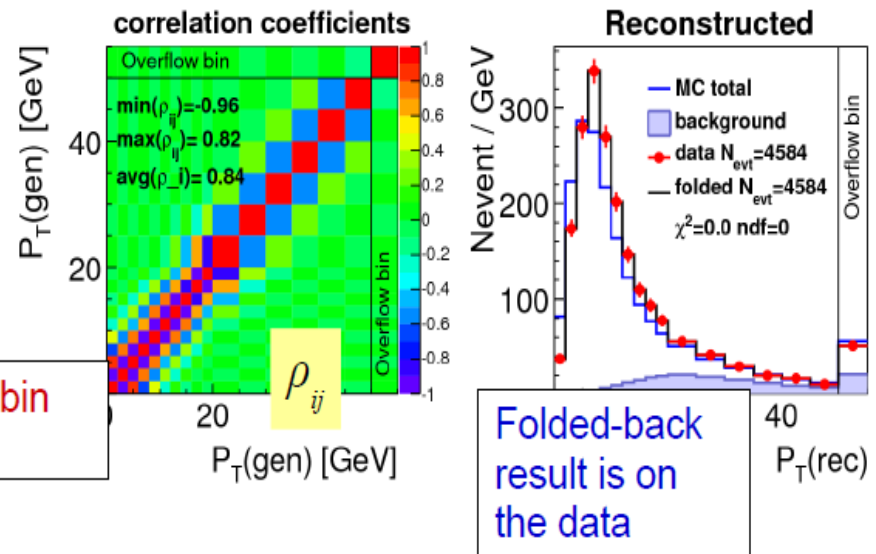
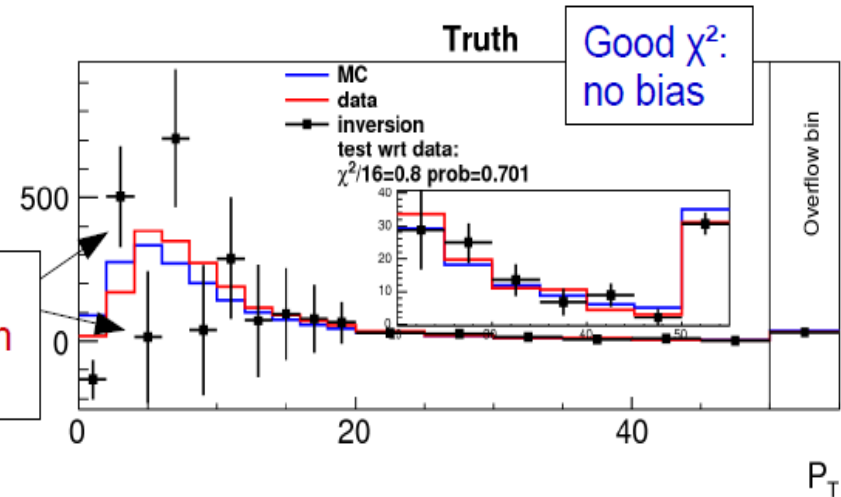
$V_{yy}$  : covariance matrix of measurements

$b$  : background

$A$  : matrix of migrations

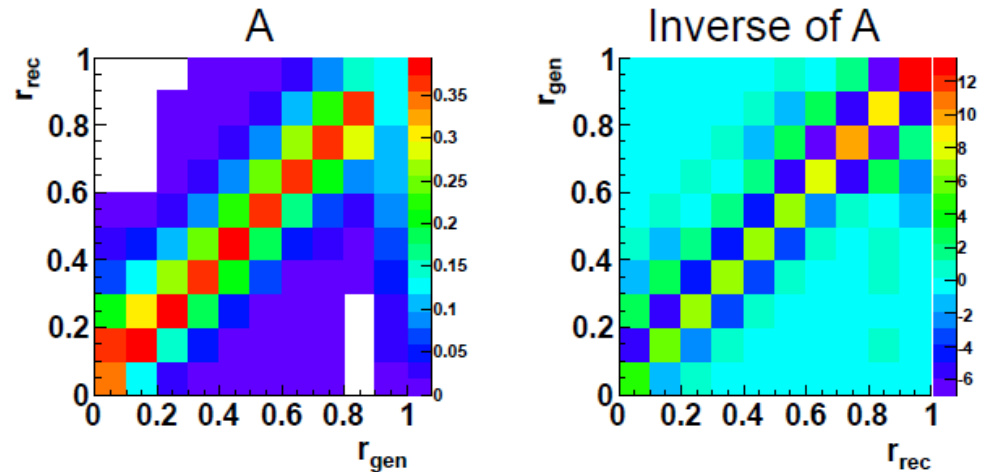
Unfolded result exhibits bin-to-bin oscillations

Large bin-to-bin correlations



# Cause of large fluctuations

- Matrix inversion: creates large negative off-diagonals  
→ statistical fluctuations of the data are amplified
- Possible improvements
  - Avoid matrix inversion “Bayesian” or “Iterative”
  - Use more reconstructed bins → TFractionFitter, TUnfold
  - Regularisation:  
TSVDUnfold, TUnfold



# Template fit

- Choose larger number of reconstructed bins than truth bins → least-square fit
- Idea: use more information → obtain better result?

$$\chi^2 = (y - b - Ax)^T V_{yy}^{-1} (y - b - Ax)$$

$y$  : measurements

$V_{yy}$  : covariance matrix of measurements

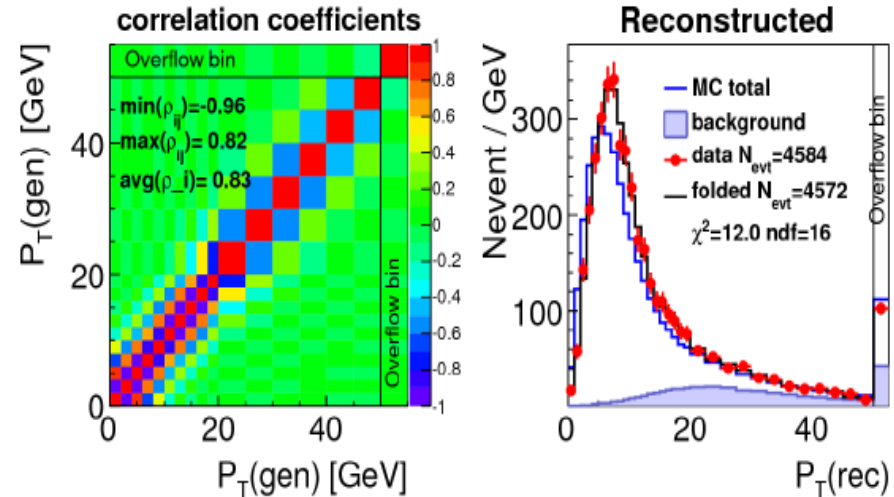
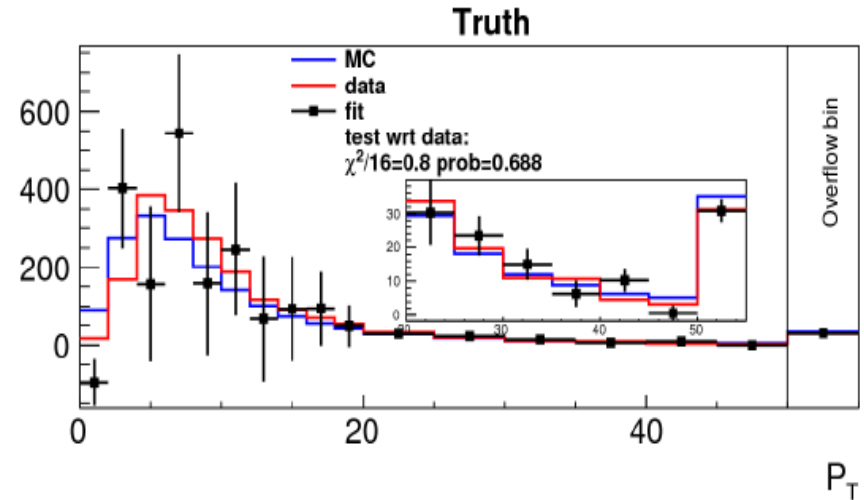
$b$  : background

$A$  : matrix of migrations

$A_{ij}$  : MC template for truth bin  $j$

$$x = (A^T V_{yy}^{-1} A)^{-1} A^T V_{yy}^{-1} (y - b)$$

covariance of  $x$  :  $V_{xx} = (A^T V_{yy}^{-1} A)^{-1}$



# Template fit with area constraint

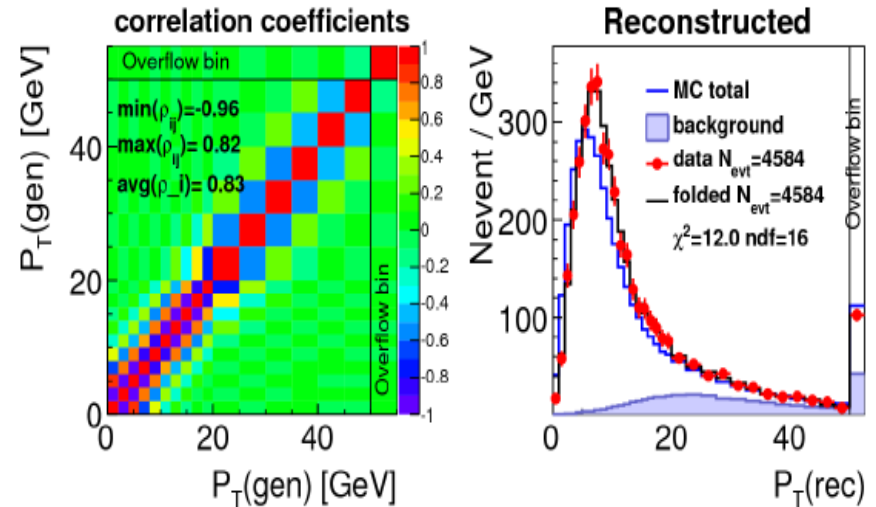
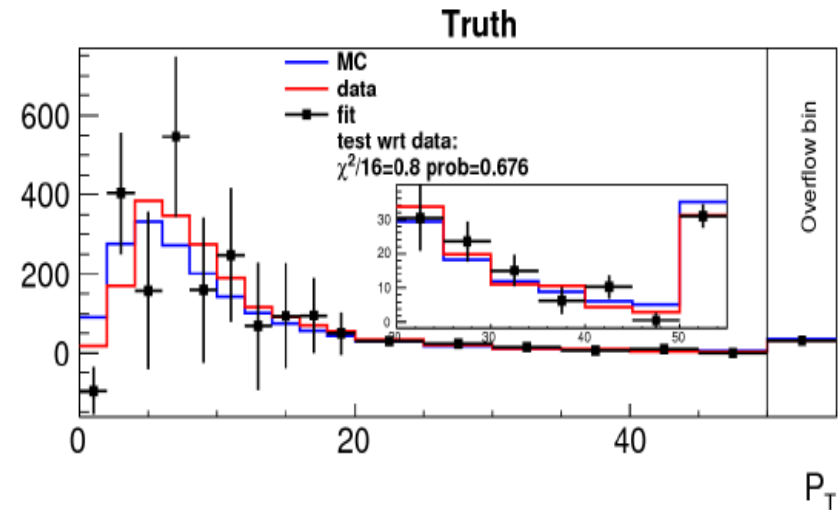
- Template with with constraint on the total number of events
- Basic idea: preserve normalisation for the folded-back result by adding the constraint

$$\sum (y_i - b_i) = \sum_{i,j} A_{ij} x_j$$

- Technical implementation: see TUnfold documentation

→ Result does not change much over unconstrained template fit, but normalisation is recovered

$$[N_{\text{data}} = N_{\text{fold}} = 4584]$$



# Tikhonov regularisation

- Basic idea: add terms to the likelihood which damp oscillations in the result.

$$\chi^2 = (y - b - Ax)^T V_{yy}^{-1} (y - b - Ax) + \tau^2 (L(x - x_B))^T L(x - x_B)$$

$y$  : measurements

$V_{yy}$  : covariance matrix of measurements

$b$  : background

$A$  : matrix of migrations

$x_B$  : regularisation bias

$L$  : regularisation conditions

$\tau$  : regularisation strength

In addition, apply area constraint to preserve normalisation

- Regularisation bias  $x_B$  : set to zero or to MC truth
- Regularisation conditions  $L$  : set to unity matrix [or mimic second derivatives, “curvature”]
- Regularisation strength  $\tau$  : “small” number

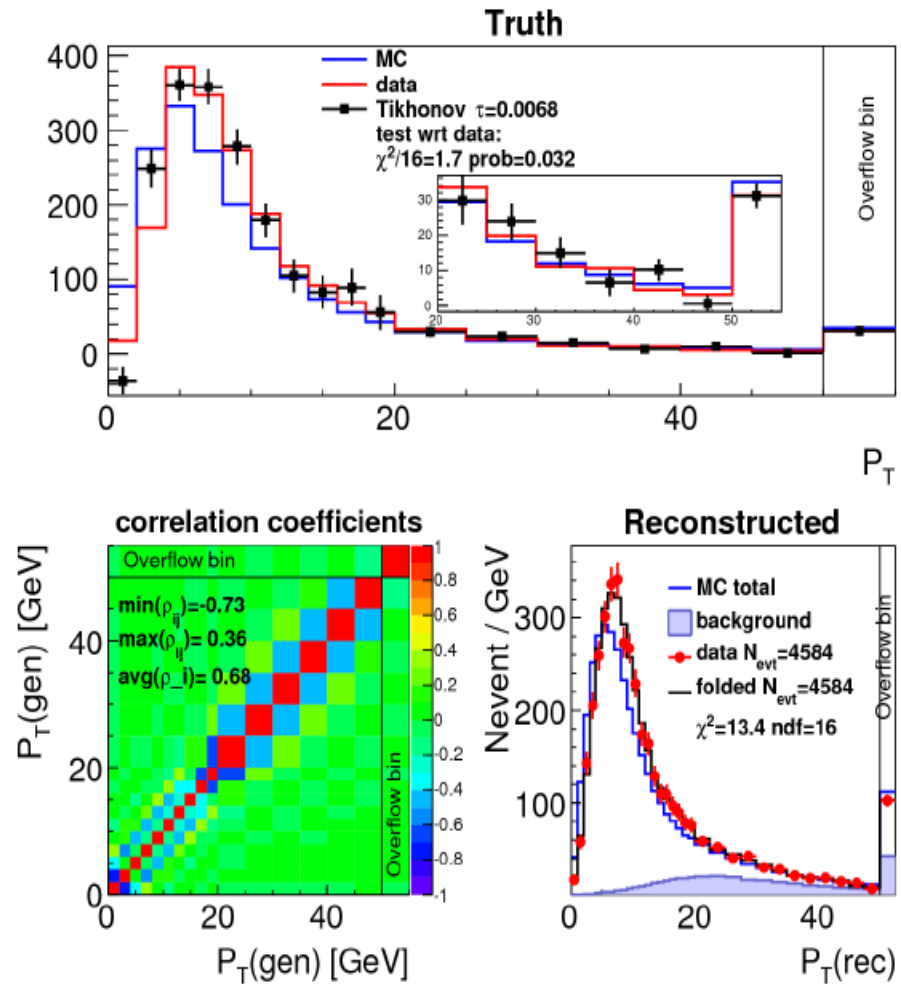
$$\tau \ll 1/\sigma$$

where  $\sigma$  ~ uncertainty after unfolding



# Tikhonov regularisation (eg. TUnfold)

- Basic idea: add terms to the likelihood which damp oscillations in the result.
- This is working well: no oscillations, moderate correlations and uncertainties
- Basic tests look reasonable
- Question: objective to choose  $\tau$



# Iterative method

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i}{\sum_k A_{ik} x_k^{(N)}}$$

Ratio data to folded  
→ iterate until ~1

efficiency:  $\epsilon_j = \sum_i A_{ij}$

start values:  $x_j^{(-1)}$  [e.g. MC truth]

iterate until  $N$  is sufficiently large

- Original works by Shepp/Vardi 1982, Kondor 1983, Mülthei/Schorr 1987
- Re-invented by D'Agostini 1995 as “Iterative Bayesian unfolding”

Note: efficiency is absorbed in a redefinition of  $A$ ,  $x$  in the original works:  $x' = \epsilon x$  and  $A' = A/\epsilon$

- Mathematical properties (Shepp/Vardi 1982 and Mülthei/Schorr 1987)
  - Ultimately converges to a maximum of the (Poisson) Likelihood  
→ like matrix inversion but with all  $x \geq 0$
  - Convergence is very slow
- Use in HEP:
  - Stop after  $N$  iterations → result will be “smooth” [regularized] but is biased to the start value

Regularisation strength:

Tikhonov:  $\tau \leftrightarrow$  Iterative:  $N_{\text{iter}}$

# Iterative method with background

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i - b_i}{\sum_k A_{ik} x_k^{(N)}}$$

efficiency:  $\epsilon_j = \sum_i A_{ij}$

start values:  $x_j^{(-1)}$  [e.g. MC truth]

OR

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i}{\sum_k A_{ik} x_k^{(N)} + b_i}$$

efficiency:  $\epsilon_j = \sum_i A_{ij}$

start values:  $x_j^{(-1)}$  [e.g. MC truth]

- Background could be subtracted from the data
- Or: background could be added to the folded MC in the denominator. This guarantees the desired property  $x \geq 0$
- D'Agostini suggests to include the background normalisation as extra bin  $x_{n+1}$ . This also guarantees  $x \geq 0$  but results in an extra parameter  $\rightarrow$  make sure to then include a background control bin in the set of measurement bins

# Evaluation of the covariance matrix

- Matrix inversion methods (with or without Tikhonov regularisation): covariance matrix is calculated analytically
- Iterative methods: non-linear, covariance matrix calculation in general has to be done by other means
- Replica method
  - Apply statistical fluctuations on the data histogram
    - N replicas of the data
  - Repeat the unfolding for each replica
  - Covariance is estimated from RMS of the results
- Bootstrap method:
  - similar idea, but based on events
  - test complete analysis chain

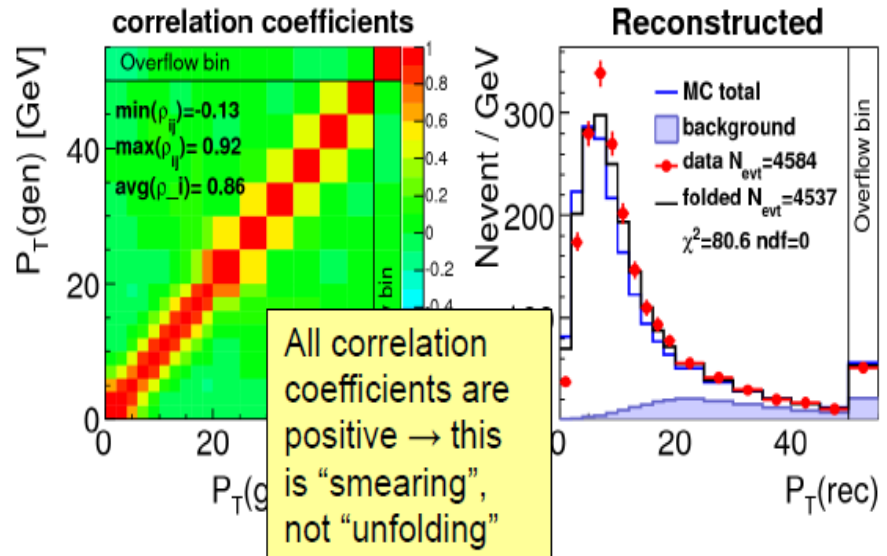
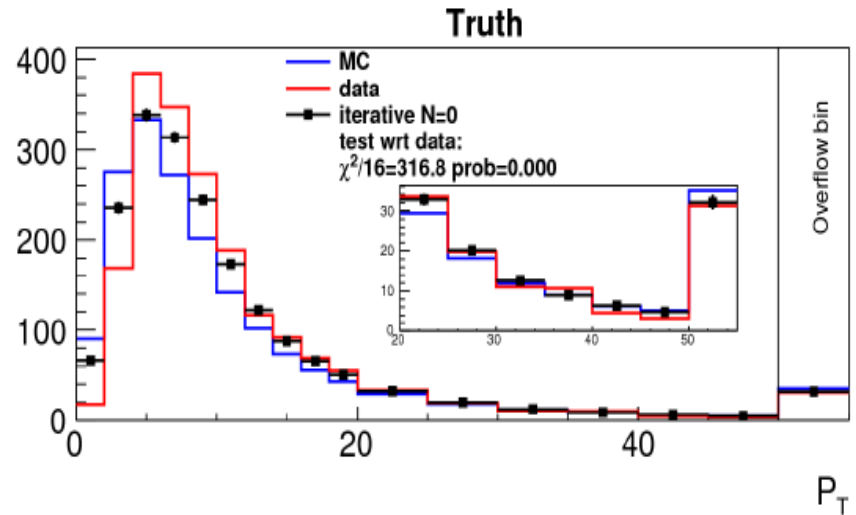
# Iterative method: 0<sup>th</sup> iteration

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i}{\sum_k A_{ik} x_k^{(N)} + b_i}$$

$$\text{efficiency: } \epsilon_j = \sum_i A_{ij}$$

start values  $x_j^{(-1)}$  set to MC truth

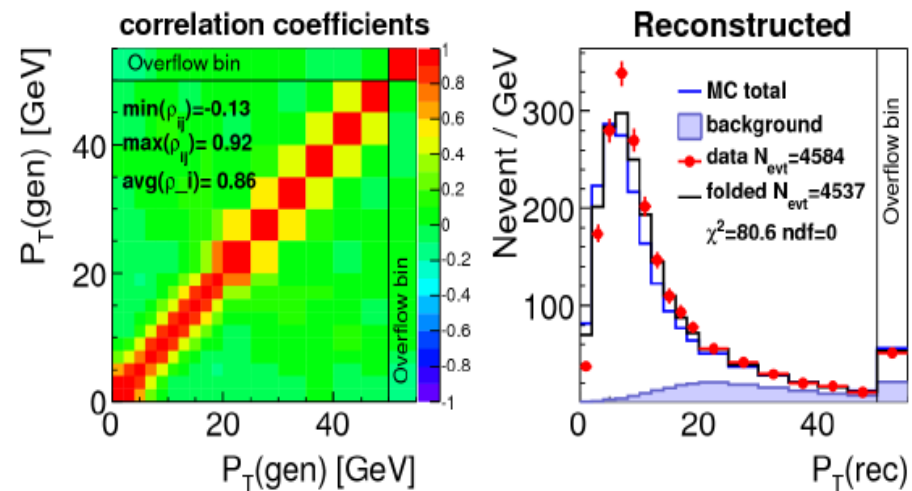
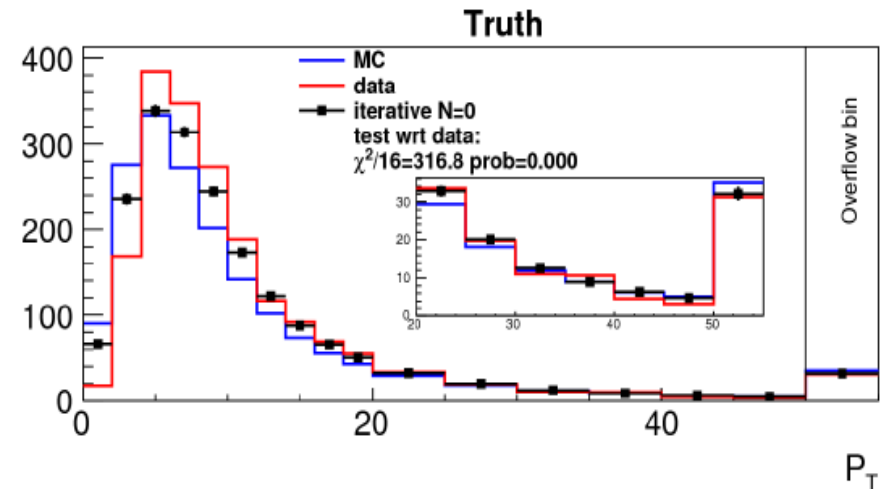
- 0<sup>th</sup> iteration: “Bayesian unfolding” from 1995 D'Agostini paper
- Result “looks nice”, very small uncertainties, but fails all tests  
→ the method has to be iterated



# Iterative method: 1<sup>st</sup> iteration

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i}{\sum_k A_{ik} x_k^{(N)} + b_i}$$

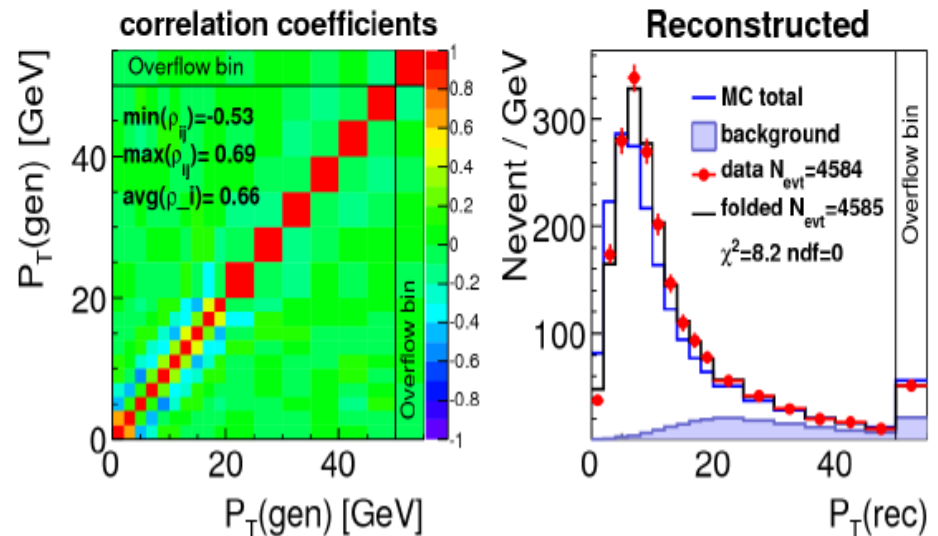
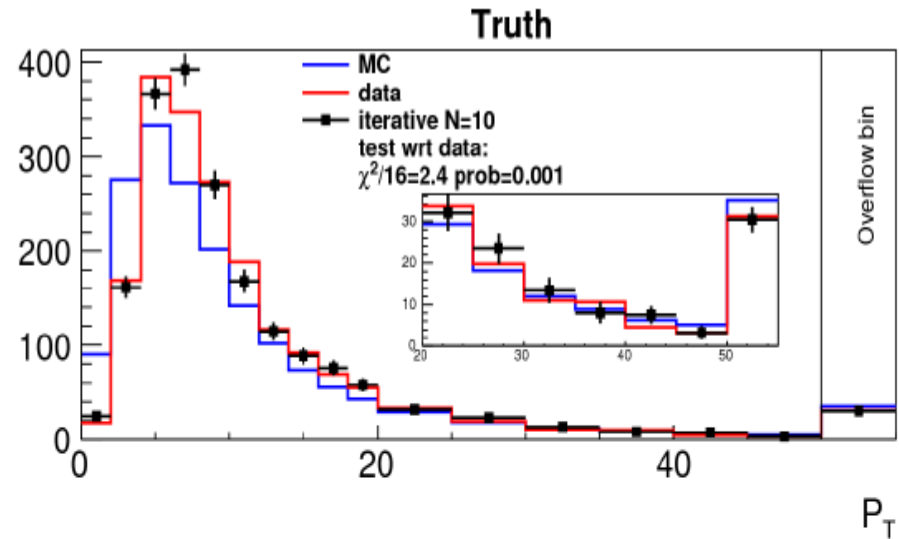
- Convergence rate is expected to grow quadratically with the number of bins [Mülthei/Schorr 1987]
- Look at 1<sup>st</sup> iteration
  - Neighboring bins have positive correlation (expect: negative)
  - Shape not described
  - Folded-back different from data  
→ have to iterate further



# Iterative method: 10<sup>th</sup> iteration

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i}{\sum_k A_{ik} x_k^{(N)} + b_i}$$

- Convergence rate is expected to grow quadratically with the number of bins [Mülthei/Schorr 1987]
- Look at 10<sup>th</sup> iteration
  - Similar to Tikhonov with strong regularisation

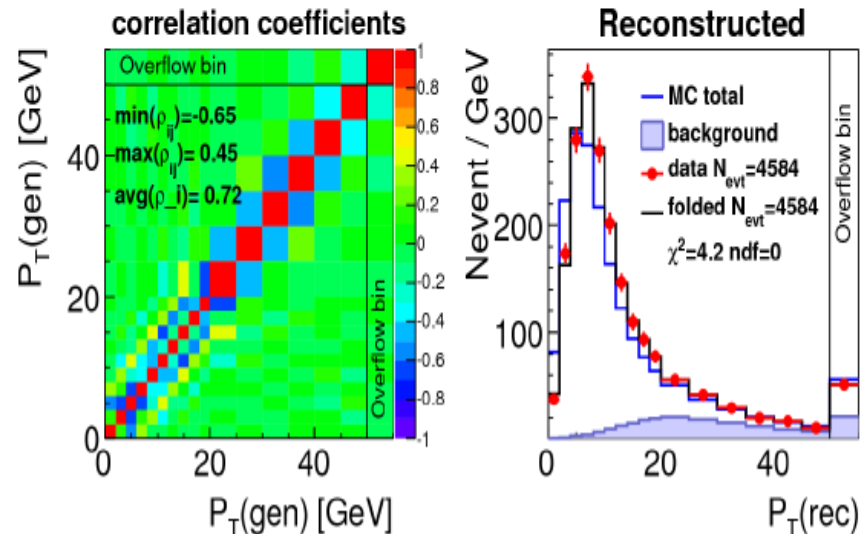
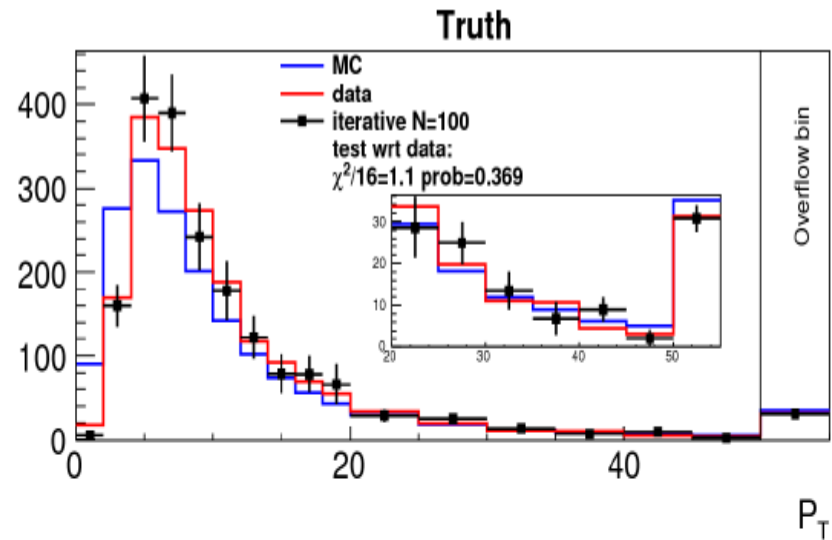




# Iterative method: 100<sup>th</sup> iteration

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i}{\sum_k A_{ik} x_k^{(N)} + b_i}$$

- Convergence rate is expected to grow quadratically with the number of bins [Mülthei/Schorr 1987]
- Look at 100<sup>th</sup> iteration
  - Similar to Tikhonov with weak regularisation

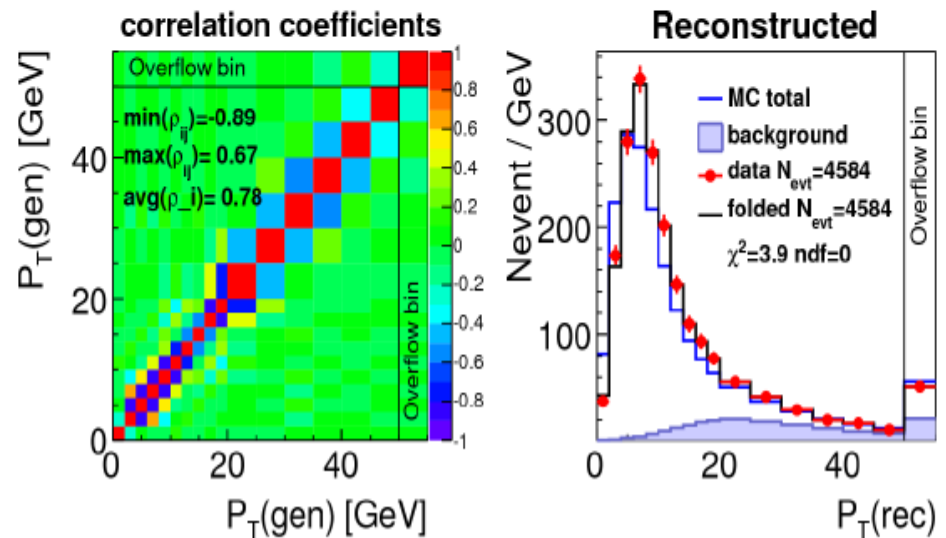
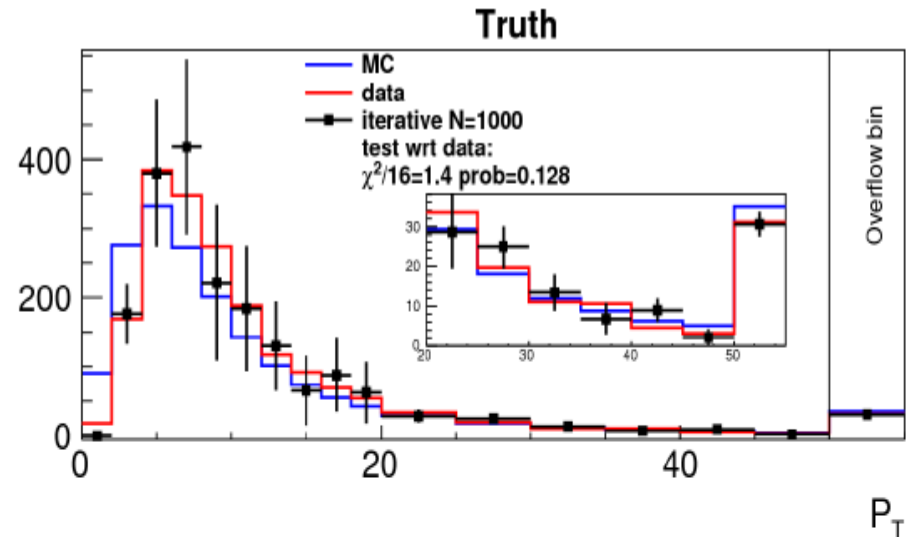




# Iterative method: 1000<sup>th</sup> iteration

$$x_j^{(N+1)} = x_j^{(N)} \sum_i \frac{A_{ij}}{\epsilon_j} \frac{y_i}{\sum_k A_{ik} x_k^{(N)} + b_i}$$

- Convergence rate is expected to grow quadratically with the number of bins [Mülthei/Schorr 1987]
- Look at 1000<sup>th</sup> iteration
  - Similar to matrix inversion, but all guaranteed to be  $x \geq 0$
  - Objective to choose number of iterations? Scan of correlation?

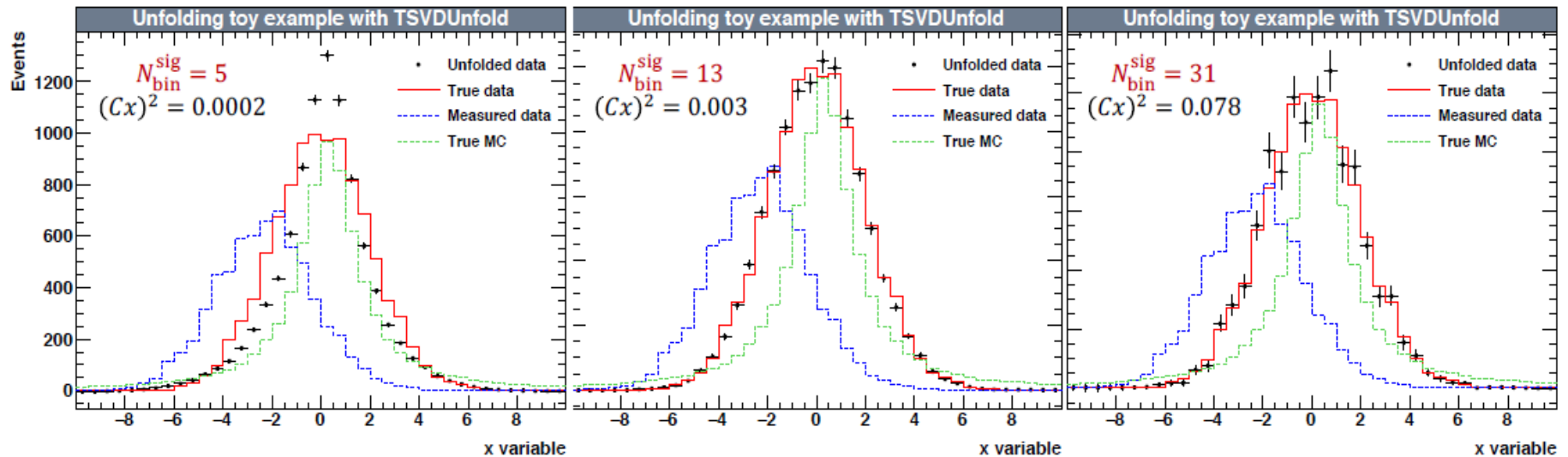


# Over- and Under-Regularised Unfolding

Over-regularised

Best regularisation choice

Under-regularised



The parameter determines the strength of the regularization

- ▶  $\tau$  too small  $\rightarrow$  oscillations
- ▶  $\tau$  too large  $\rightarrow$  unfolded spectrum biased towards MC

# Selection of other unfolding methods

- SVD [Hoecker et al, 1995]
  - Equivalent to matrix inversion with Tikhonov regularisation, parameter  $\tau$  from Eigenvalue analysis
- Shape-constrained unfolding [Kuusela, Panaretos 2015]
- Improved D'Agostini [2010]
- Fully Bayesian [Choudalakis 2012]

# RooUnfold package

- Provide a framework for different algorithms
  - Can compare performance directly, with common user code
    - RooUnfold takes care of different binning, normalisation, efficiency conventions
  - Can use common RooUnfold utilities
    - Write once, use for all algorithms
  - Currently implement or interface to iterative Bayes, SVD, TUnfold, unregularised matrix inversion, and bin-by-bin correction factors algorithms
- Simple OO design
  - “response matrix” object can be filled separately from training sample
    - in a different routine, or a different program (ROOT I/O support)
- Simple interface for the user
  - From program, ROOT/CINT script, or interactive ROOT prompt
  - Fill with histograms, vectors/matrices, ... or direct methods:
    - `response->Fill( $x_{\text{measured}}, x_{\text{true}}$ )` and `Miss( $x_{\text{true}}$ )` methods takes care of normalisation
  - Results as a histogram with errors, or vector and covariance matrix

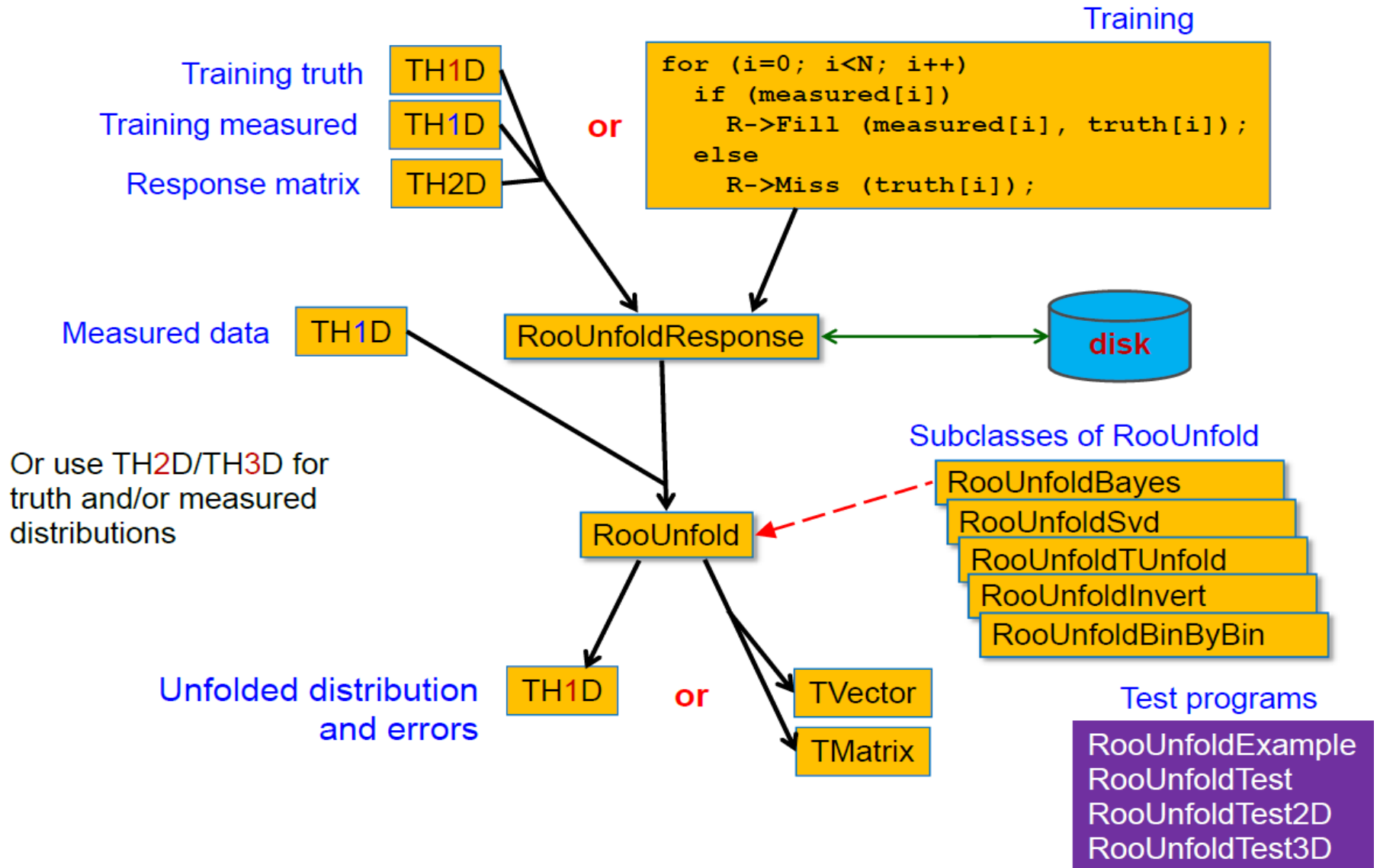
# RooUnfold features

- Supports different binning scenarios
  - multi-dimensional distributions (1D, 2D, and 3D)
  - Different binning (or even dimensionality) for measured and truth
  - Option to include or exclude histogram under/overflow bins in the unfolding
- Supports different methods for error computation (simple switch). In order of increasing CPU time:
  - No error calculation (uses  $\sqrt{N}$ )
  - bin-by-bin errors (no correlations)
  - full covariance matrix from the propagation of measurement errors in the unfolding, or
  - covariance matrix from MC toys
    - useful to test error propagation and when it is inaccurate
- These details are handled by the framework, so don't need to be implemented for each algorithm

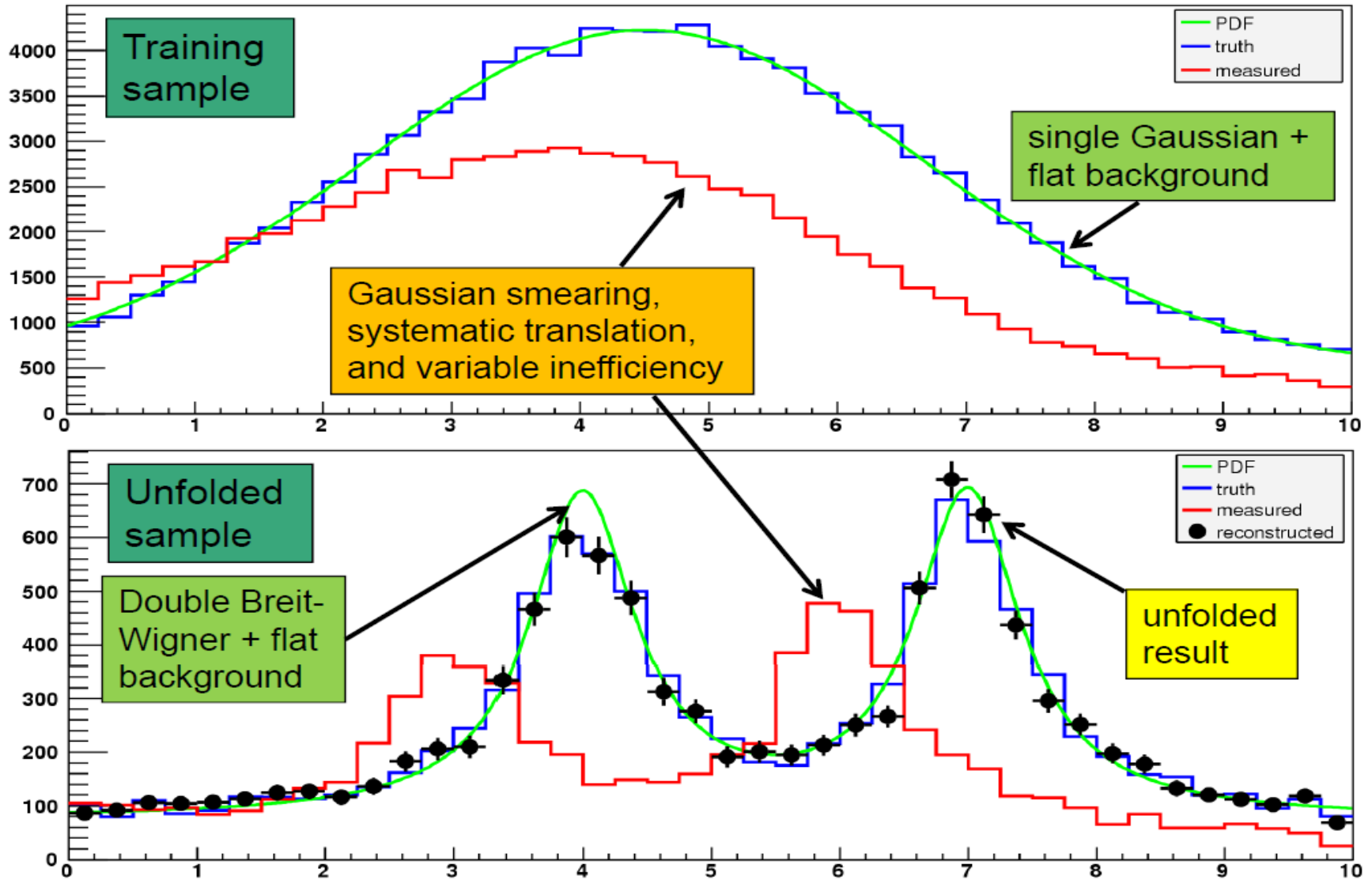
# RooUnfold testing

- Calculates **resolutions**, **pulls**, and  $\chi^2$
- Includes a **toy MC test framework**, allowing selection of different
  - PDFs and PDF parameters
  - binning
  - 1D, 2D, 3D tests
  - unfolding methods and parameters
  - Test procedures for the regularisation parameter and errorsand plotting results from a single command

# Roofold classes

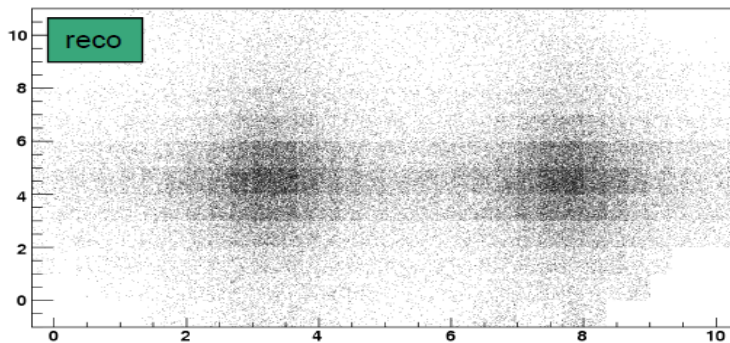
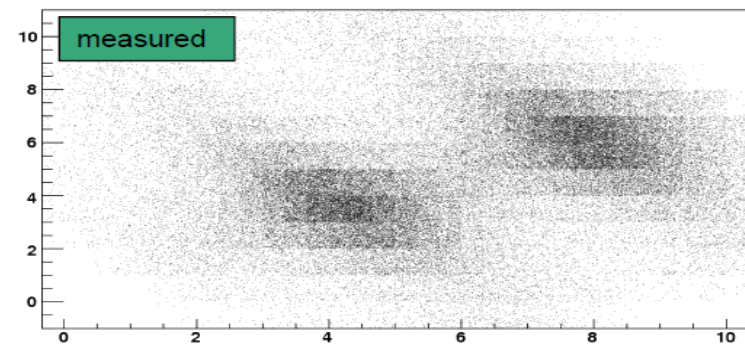
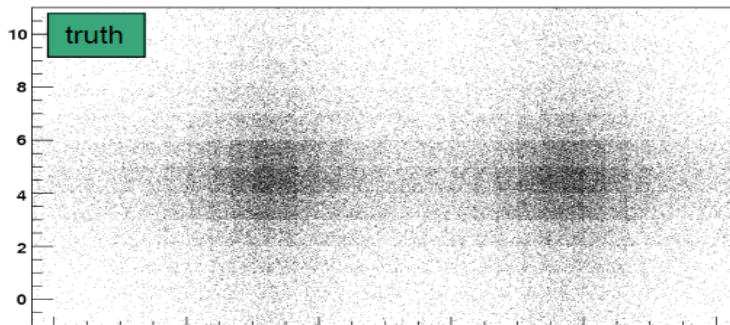
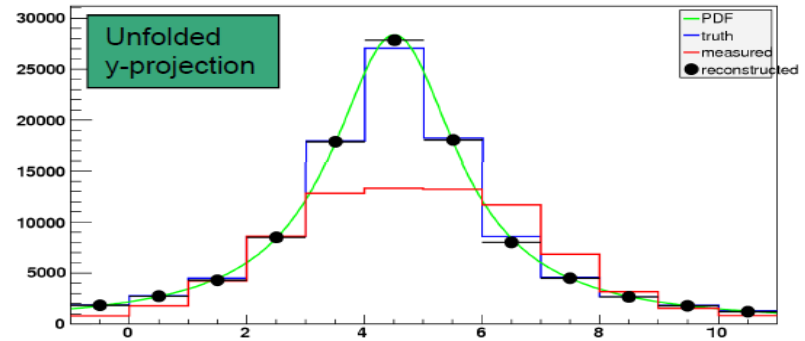
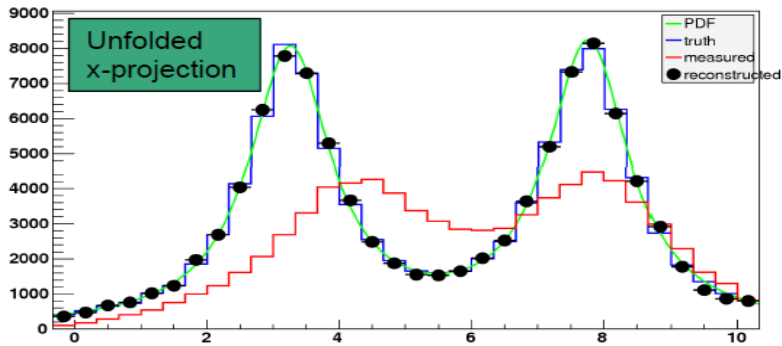


# RooUnfold example (Bayes)





# RoofUnfold example (Bayes)



2D unfolding

2D Smearing, bias, variable efficiency, and variable rotation

# RooUnfold algorithms: Iterative Bayes

- Uses the method of **Giulio D'Agostini** (1995), implemented by **Fergus Wilson** and Tim Adye
  - Uses repeated application of **Bayes' theorem** to invert the response matrix
  - Regularisation by stopping iterations before reaching “true” (but wildly fluctuating) inverse
    - Regularisation parameters is the **number of iterations**, which in principle has to be tuned according to the statistics, number of bins, etc. In practice, the results are fairly **insensitive** to the precise setting.
- Implementation details:
  - Initial **prior** is taken from training truth, rather than a flat distribution
    - Does not bias result once we have iterated, but perhaps reach optimum faster
  - Takes account of multinomial errors on the data sample but not, by default, uncertainties in the response matrix (finite MC statistics), which is very slow
  - Does not normally do **smoothing** (can be enabled with an option)

# RooUnfold algorithms: SVD

- Uses the method of **Andreas Höcker** and **Vato Kartvelishvili**
- Obtains inverse of response matrix using singular value decomposition
  - Use number-of-events matrix to keep track of MC uncertainties
- Regularisation with a smooth cut-off on small singular value contributions (these correspond to high-frequency fluctuations)
  - Replace  $s_i^2 \rightarrow s_i^2 / (s_i^2 + s_k^2)$
  - $k$  determines the relative contributions of MC truth and data
    - $k$  too small  $\rightarrow$  result dominated by **MC truth**
    - $k$  too large  $\rightarrow$  result dominated by **statistical fluctuations**
  - $k$  needs to be tuned for the particular type of distribution, number of bins, and approximate sample size
- Unfolded error matrix includes effect of finite MC training statistics (usually small)

# RooUnfold algorithms: TUnfold

- Uses the TUnfold method implemented by **Stefan Schmitt** and included in ROOT
  - RooUnfold includes an interface to this class
- Performs a **matrix inversion** with 0-, 1-, or 2-order polynomial **regularisation** of neighbouring bins
  - RooUnfold automatically takes care of packing 2D and 3D distributions and creating the appropriate regularisation matrix required by TUnfold
- TUnfold can determine an **optimal regularisation parameter** ( $\tau$ ) by scanning the “L-curve” of  $\log_{10}(\chi^2)$  vs  $\log_{10}(\tau)$ .

# RooUnfold algorithms: Unregularised

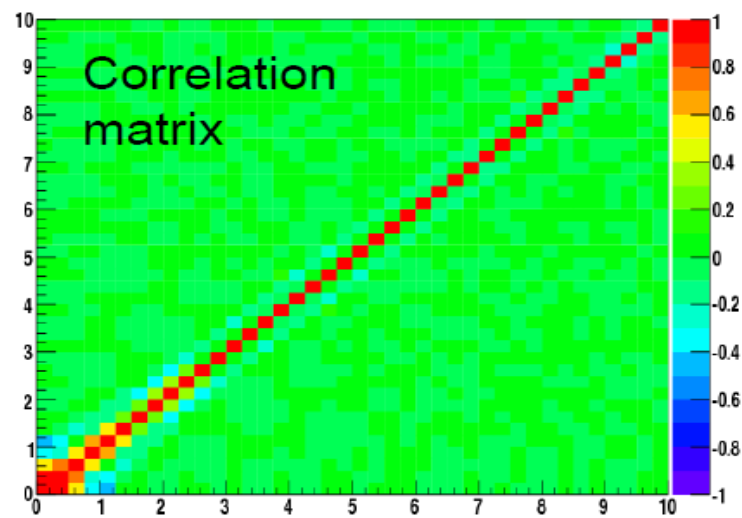
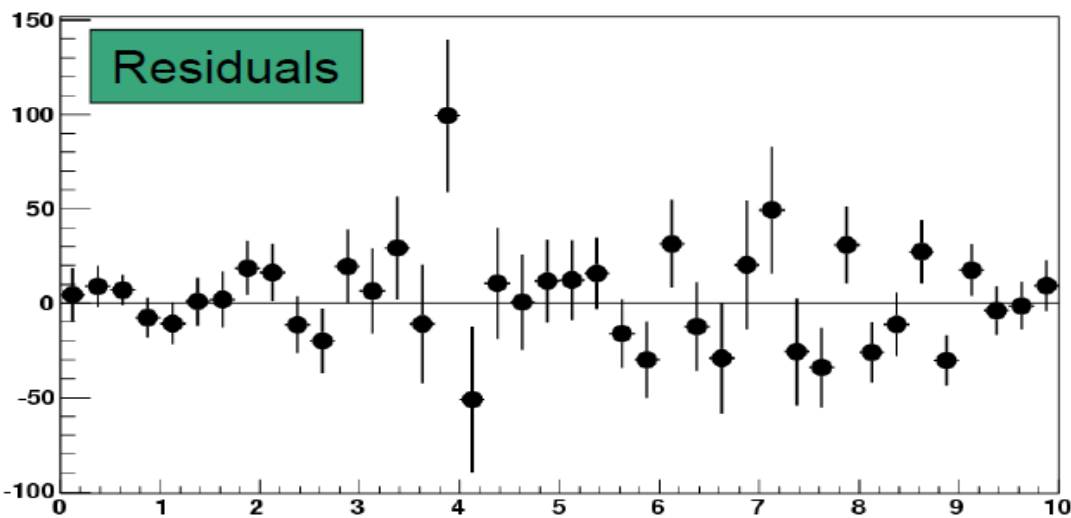
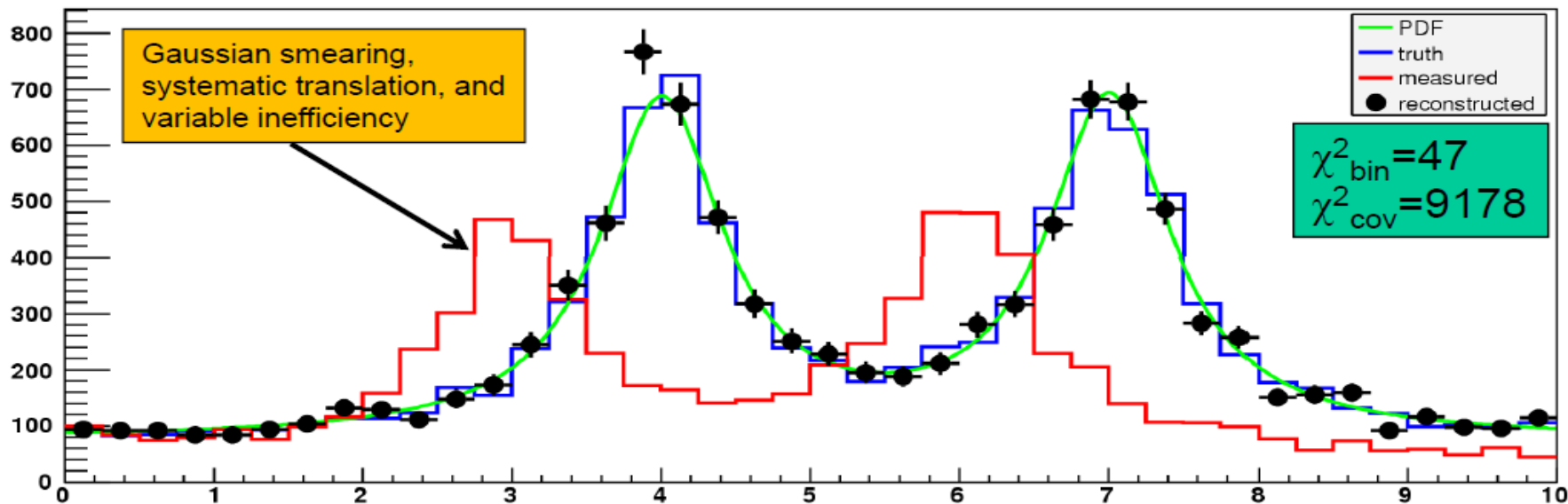
- Very simple algorithms
  - using bin-by-bin correction factors, with no inter-bin migration
  - using unregularised matrix inversion with singular value removal (TDecompSVD)

are included for comparison – and to demonstrate why they should not be used in most cases!

# RooUnfold algorithms: comparison

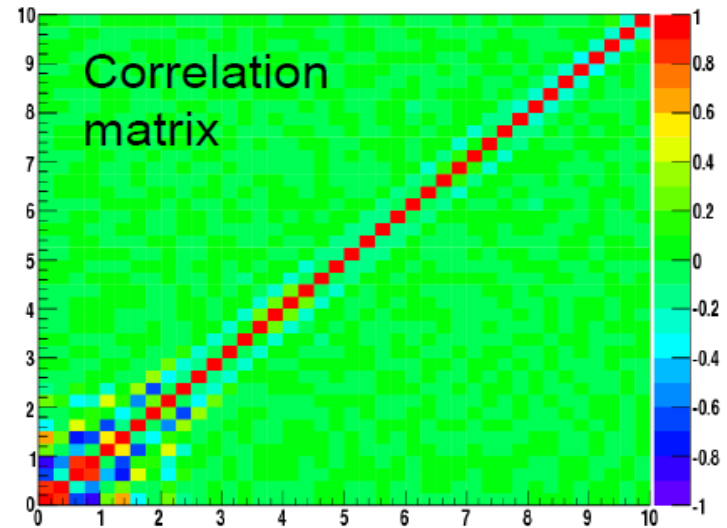
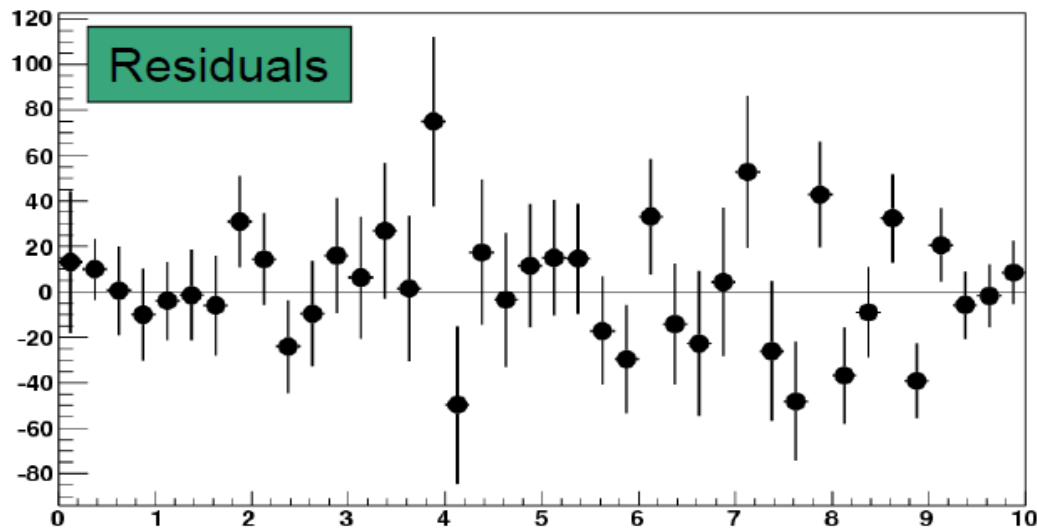
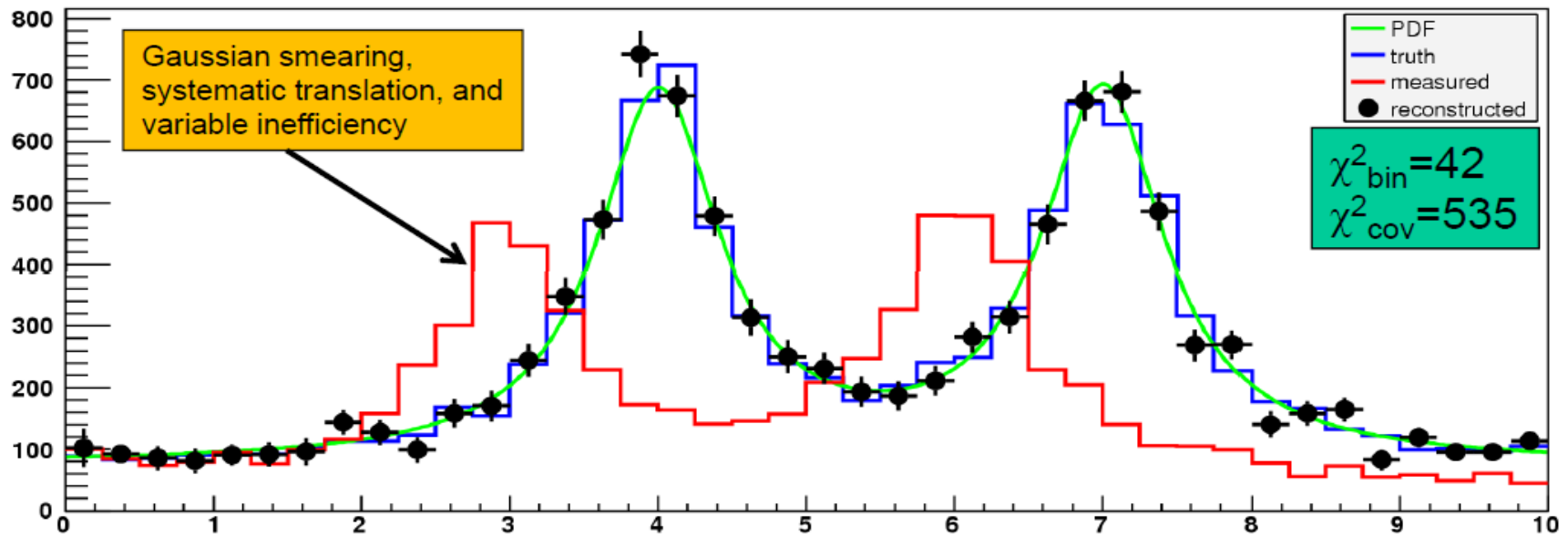
- TUnfold and unregularised matrix inversion require the number of bins,  $N_{\text{measured}} \geq N_{\text{true}}$ 
  - TUnfold claims best results if  $N_{\text{measured}} > N_{\text{true}}$ , eg.  $N_{\text{measured}} = 2N_{\text{true}}$ 
    - This is a common general recommendation from unfolding experts, but perhaps is most relevant to these types of algorithms with explicit regularisation
    - This is an implicit additional regularisation, since we are “smoothing” two bins into one
- SVD implementation and bin-by-bin methods only support  $N_{\text{measured}} = N_{\text{true}}$ 
  - SVD implementation also only works well for 1D distributions
- The choice of the SVD regularisation parameter has to be done by the user
  - TUnfold can often do this automatically
    - Can we do something similar for the SVD method?
  - The performance of the Bayes method is relatively insensitive to the regularisation parameter (number of iterations)

# RoofUnfold with Bayes algorithm (3 iterations)



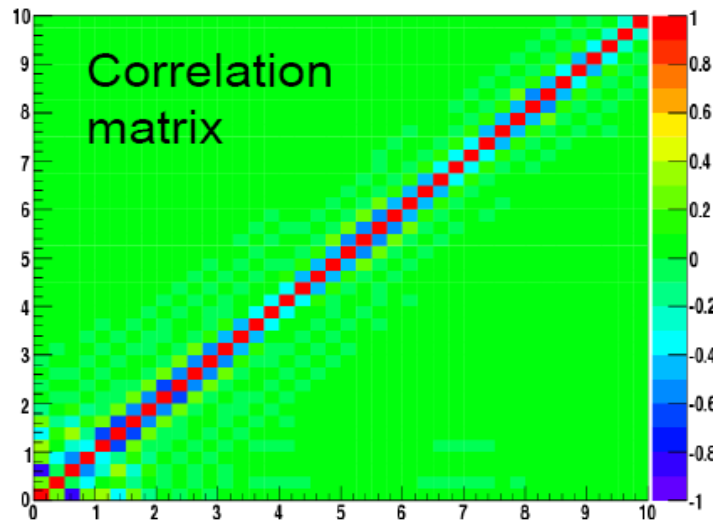
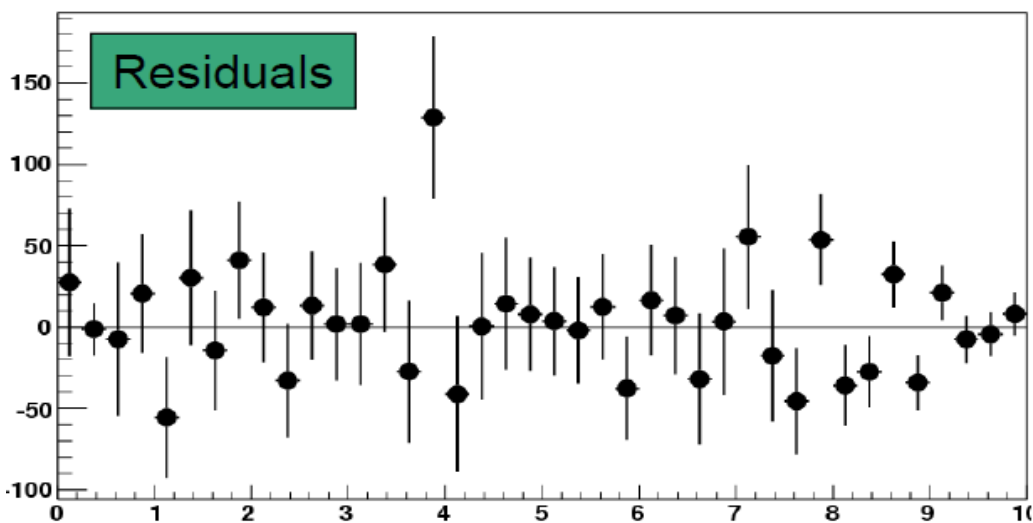
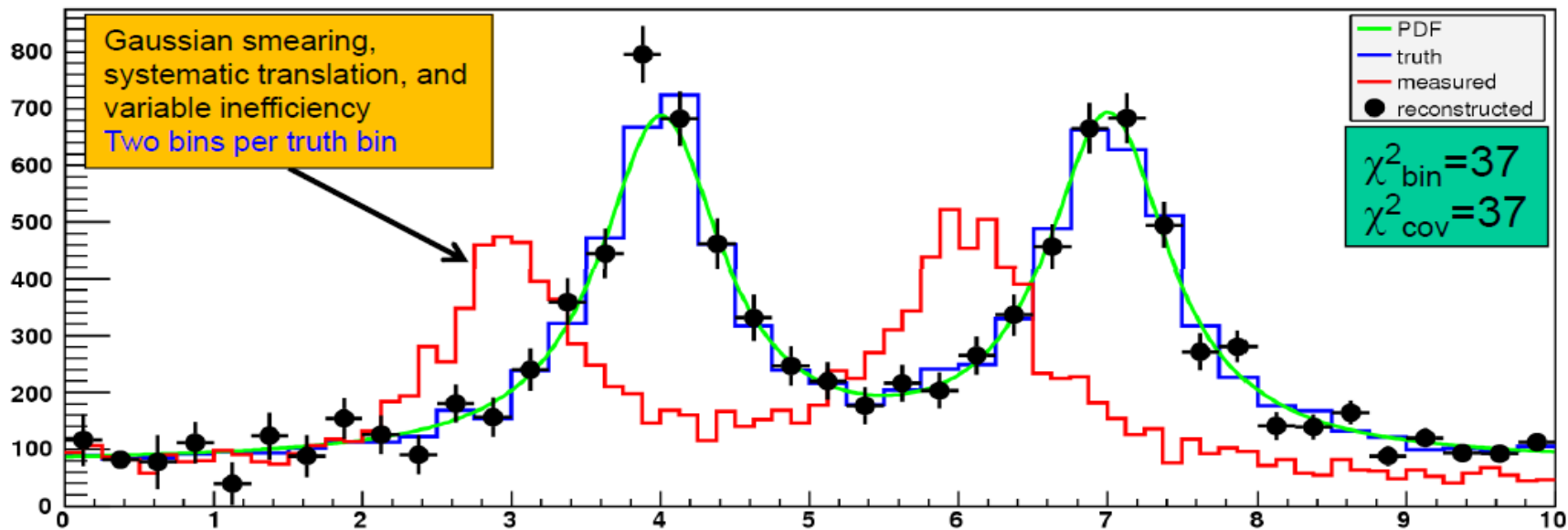


# RooUnfold with SVD algorithm ( k=30 )

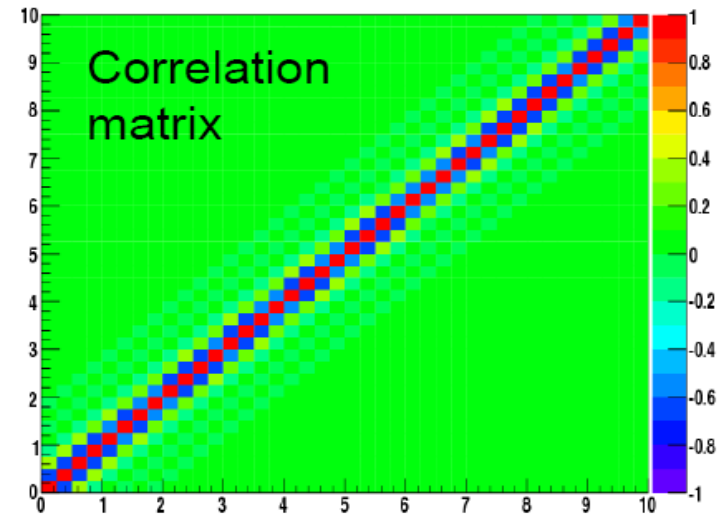
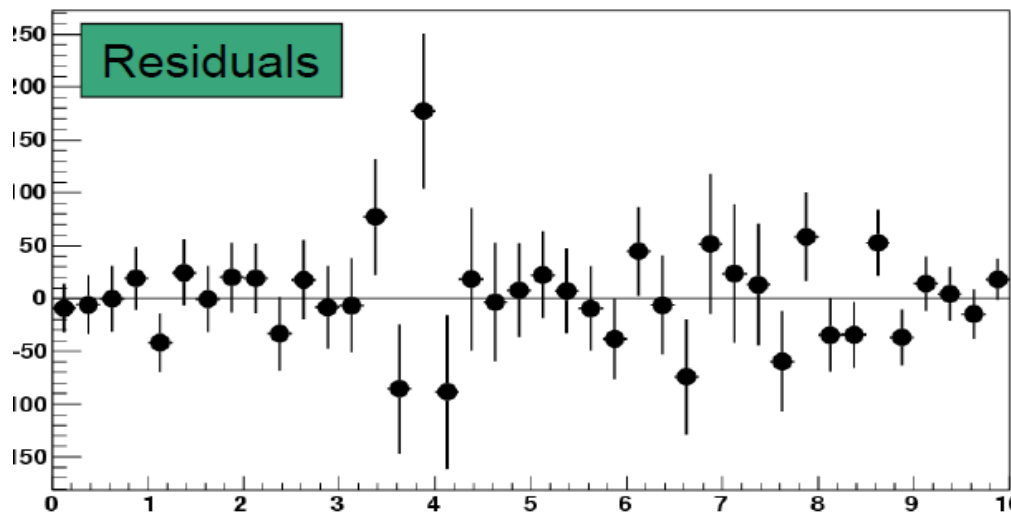
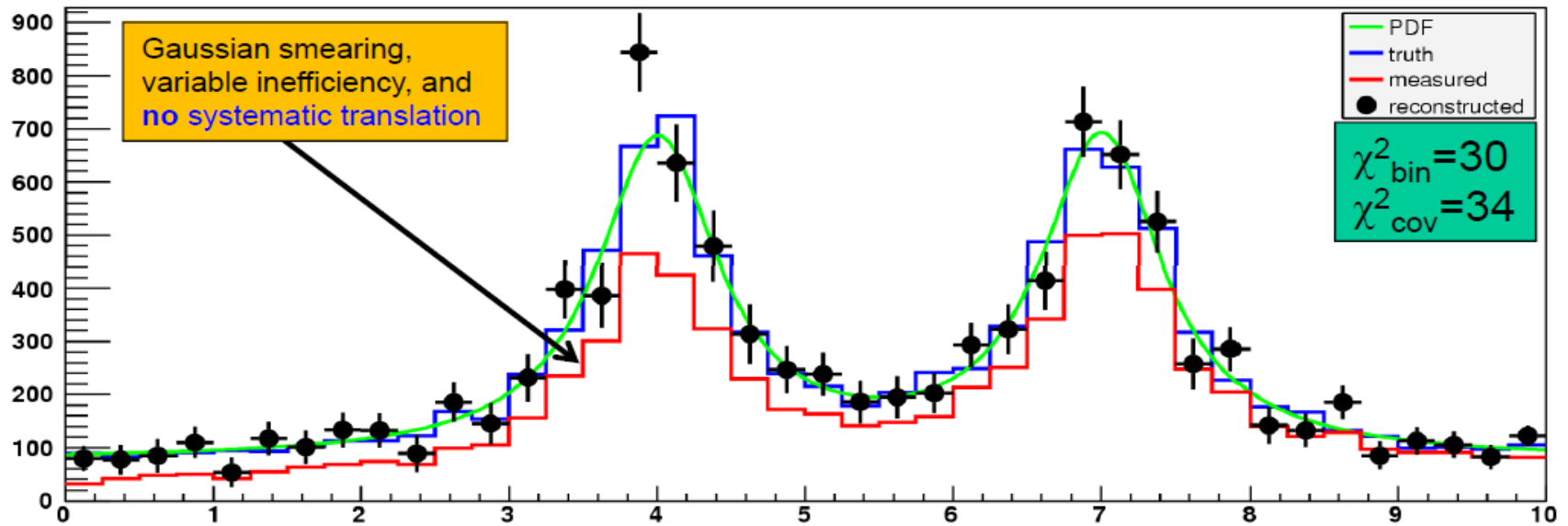




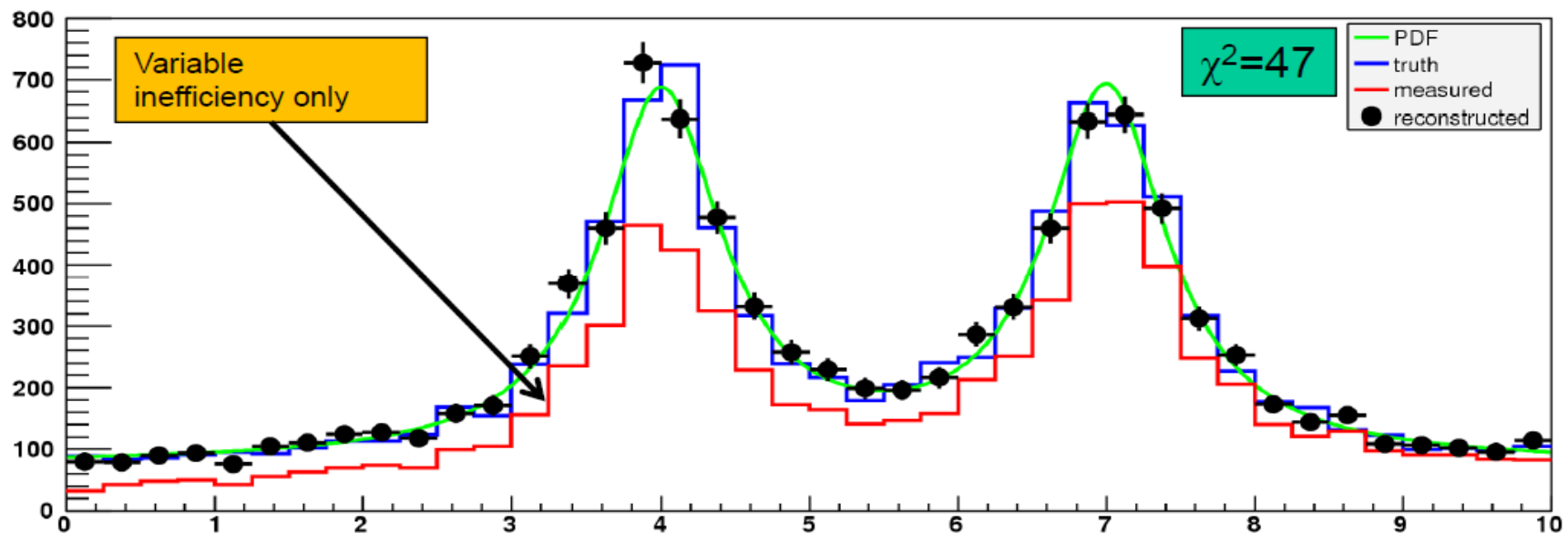
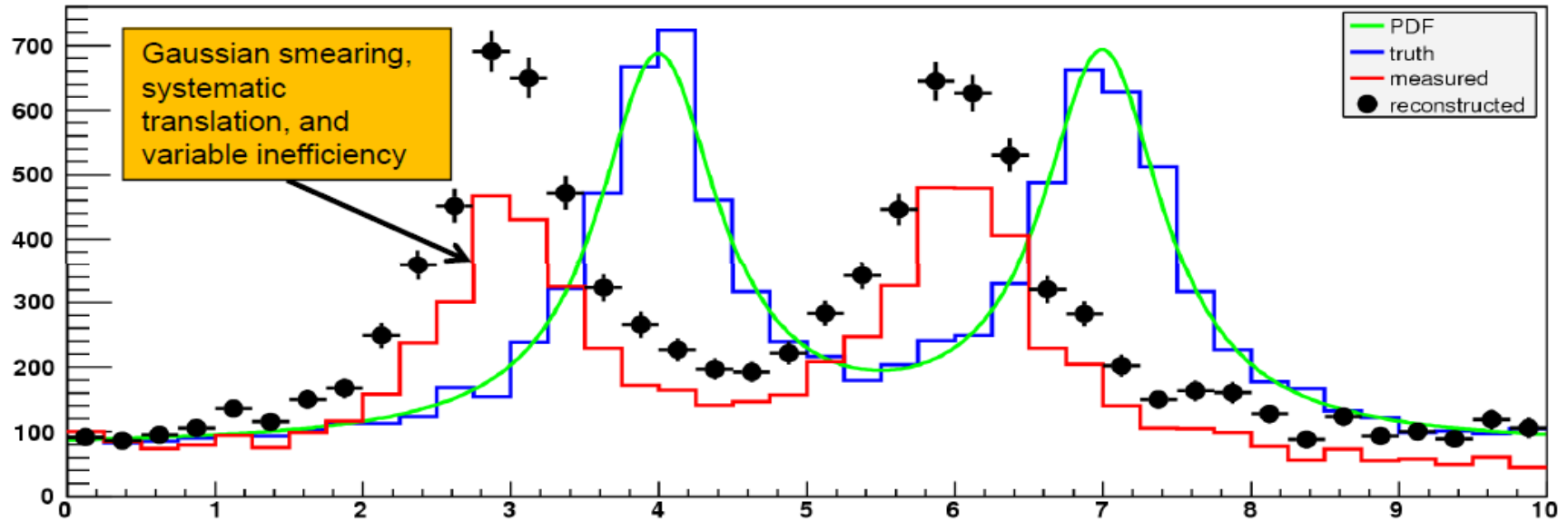
# RooUnfold with TUnfold algorithm ( $\tau=0.004$ )



# Unregularised matrix inversion

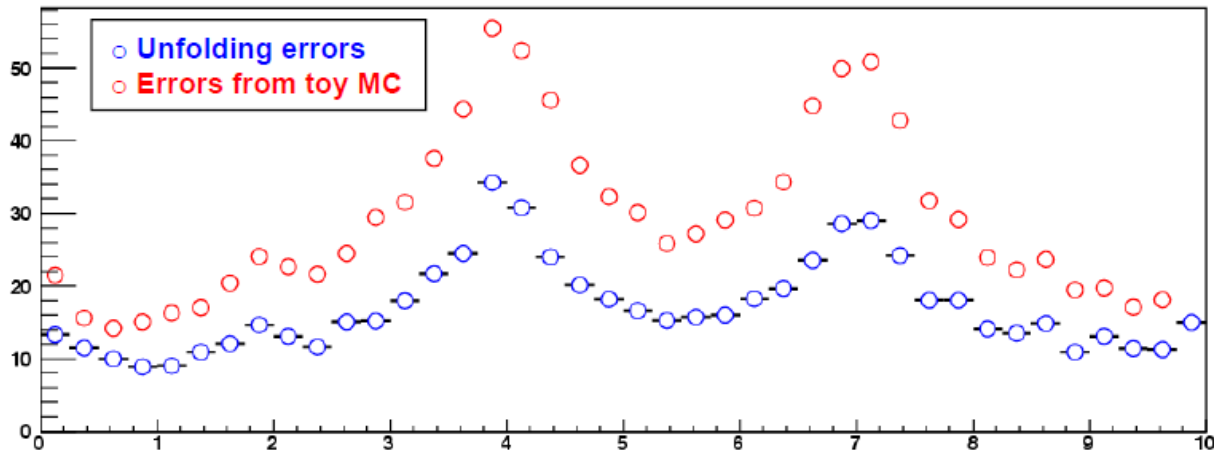


# Simple correction factors



# Unfolding errors

- All methods return a full covariance matrix of the errors on the unfolded histogram due to uncertainties on the measured distribution.
  - This is often calculated by propagation of errors
    - but not always possible if there are non-linearities or other problems, eg. the iterations in the Bayes method are not handled in D'Agostini's formalism:



- RooUnfold allows the covariance matrix to be calculated from toy MC instead
  - provides a cross-check of the error propagation or replace it if there are problems

# Bin-to-bin correlations

- Regularisation introduces inevitable correlations between bins in the unfolded distribution
  - To calculate a correct  $\chi^2$ , one has to invert the covariance matrix:  
$$\chi^2 = (\mathbf{x}_m - \mathbf{x}_t)^T \mathbf{V}^{-1} (\mathbf{x}_m - \mathbf{x}_t)$$
- However, in many cases, the covariance matrix is poorly conditioned, which makes calculating the inverse problematic
  - Inverting a poorly conditioned matrix involves subtracting large, but very similar numbers, leading to significant effects due to the machine precision
- In any case,  $\chi^2$  may not be the best figure of merit
  - could improve  $\chi^2$  by relaxing regularisation  $\rightarrow$  larger errors, but also larger residuals
  - Is there a better figure of merit?

# Which Method To Choose?

There is no "best" method. Depends on the analysis.

Main questions:

How to choose regularization parameters?

After how many iterations to stop in the iterative Bayesian unfolding?

Danger: Regularization and early stopping in iterative unfolding introduce a bias

Don't forget:

in some cases it is most useful to publish folding matrix with the result

# Summary

- Unfolding: get measurements independent of the detector response
- Alternative: publish folding matrix with the result
- Many methods exist, only a few have been compared in this talk
- Big unfolding families investigated in this talk:
  - Matrix inversion + Tikhonov regularisation (parameter  $\tau$ )
  - Iterative methods + truncation after  $N_{\text{iter}}$  steps
- Main question: how to choose the regularisation strength.
- Danger to obtain biased results if regularisation is too strong

# References

- Bayesian:
  - Nucl.Instrum.Meth. A362 (1995) 487-498
  - arXiv:1010.0632
- TSVDUnfold
  - Nucl.Instrum.Meth.A372 (1996) 469-481
- TUnfold
  - JINST 7 (2012) T10003 [arXiv:1205.6201]





# Deconvolution

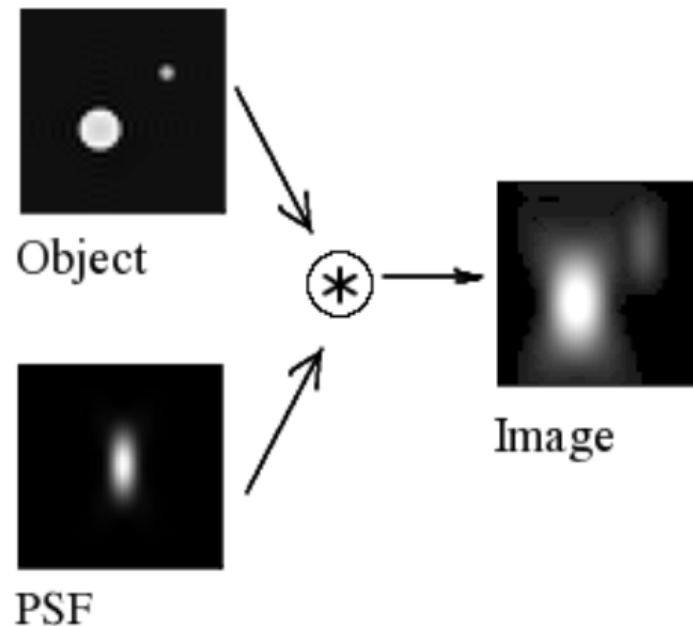
Finite resolution of the detector  
smears the quantities we're  
interested in.

Goal:  
smeared information  
→ original information

This is called *deconvolution* or  
*unfolding*

"Inverse problem"

Problem can be ill-posed in the  
sense that unfolded result can be  
very sensitive to small perturbations  
in the data



Example:

Smearing of a telescope image

[https://en.wikipedia.org/wiki/Point\\_spread\\_function](https://en.wikipedia.org/wiki/Point_spread_function)

# To Unfold or Not?

It's a lot of work, and often produces biased or otherwise unsatisfactory results. Moreover it's often unnecessary.

"Forward fitting" is much easier

- ▶ Take theory prediction
- ▶ Convolve it with the response of the detector
- ▶ Compare smeared theory directly with the data

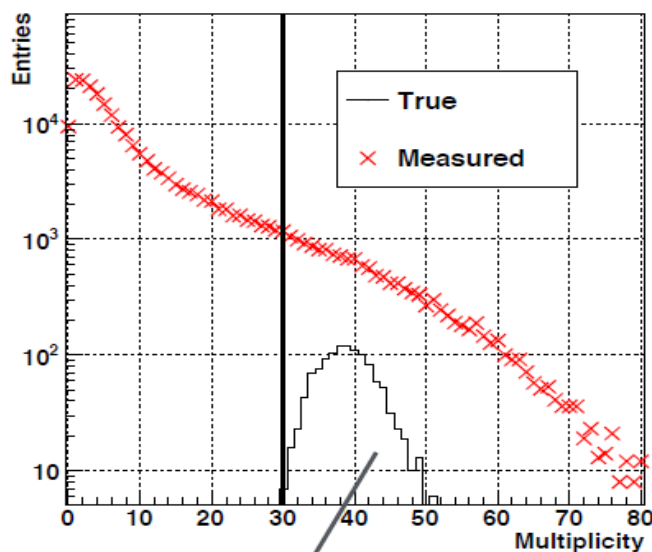
# When Unfolding Make Sense

1. Results from experiment A and B with different response function are to be compared
2. It is too complicated to publish the response function of the detector along with the data
  - ▶ Detector response might be very complex, e.g., time dependent
  - ▶ Sometimes computer code reflecting the response would have to be published
  - ▶ Danger that future users don't use the filter correctly

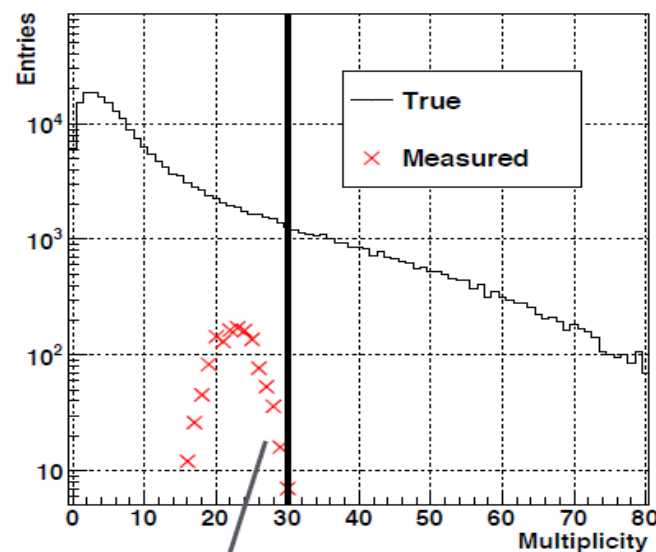
# When Unfolding Make Sense

- Multiplicity distributions  $P(N_{\text{ch}})$ 
  - ▶ Measured multiplicity differs from true charged particle multiplicity due to detector effects (efficiency, fake hits, ...)
- $p_T$  spectra, e.g.,  $\pi^0$  spectrum measured with a calorimeter
  - ▶ finite energy resolution and shower overlaps in a calorimeter affect the  $p_T$  of the reconstructed shower

Example: multiplicity distributions in pp collisions



true  $N$ 's contributing to  
measured  $N = 30$



measured  $N$ 's for a true  $N = 30$

arXiv:0912.0023

# Response Matrix (I)

Suppose that we deal with continuous variables (e.g., transverse momentum)

$f_t(x_t)$  : distribution of true values (normalized to unity)

$f_m(x_m)$  : distribution of measured values (normalized to unity)

$f_b(x_m)$  : distribution of background (normalized to unity)

Response function  $R$ :

$$R(x_m|x_t) = \underbrace{r(x_m|x_t)}_{\text{"smearing"}} \times \underbrace{\varepsilon(x_t)}_{\text{"efficiency"}} \quad \text{probability (density) to observe } x_m \text{ given } x_t$$

By construction, one has

$$\int_{\Omega_m} r(x_m|x_t) dx_m = 1$$

# Response Matrix (II)

Further definitions:

$m_{\text{tot}}$  : number of true events

$n_{\text{tot}}$  : number of measured events

$b_{\text{tot}}$  : number of background events

$$\mu_{\text{tot}} = E[m_{\text{tot}}], \quad \nu_{\text{tot}} = E[n_{\text{tot}}], \quad \beta_{\text{tot}} = E[b_{\text{tot}}]$$

It is practical to work with discrete bins. E.g., probability to find true in bin  $j$ :

$$p_j = \int_{\text{bin } j} dx_t f_t(x_t), \quad \mu_j = \mu_{\text{tot}} \times p_j$$

Ignoring backgrounds, the measured number of entries in bin  $i$  is:

$$\begin{aligned} \nu_i &= \mu_{\text{tot}} \int_{\Omega_t} dx_t \text{Prob}(x_m \text{ in } i | \text{true } x_t, \text{ detected}) \\ &\quad \times \text{Prob}(\text{detect } x_t) \times \text{Prob}(\text{produce } x_t) \\ &= \mu_{\text{tot}} \int_{\text{bin } i} dx_m \int_{\Omega_t} dx_t r(x_m | x_t) \varepsilon(x_t) f_t(x_t) \end{aligned}$$

# Response Matrix (III)

Further definitions:

$$\begin{aligned}\nu_i &= \mu_{\text{tot}} \int_{\text{bin } i} dx_m \sum_{j=1}^M \int_{\text{bin } j} dx_t r(x_m|x_t) \varepsilon(x_t) f_t(x_t) \\ &= \sum_{j=1}^M \int_{\text{bin } i} dx_m \int_{\text{bin } j} dx_t \frac{r(x_m|x_t) \varepsilon(x_t) f_t(x_t)}{\mu_j / \mu_{\text{tot}}} \mu_j\end{aligned}$$

This may be written as

$$\nu_i = \sum_{j=1}^M R_{ij} \mu_j$$

with the components of the response matrix

$$R_{ij} = \frac{\int_{\text{bin } i} dx_m \int_{\text{bin } j} dx_t r(x_m|x_t) \varepsilon(x_t) f_t(x_t)}{\int_{\text{bin } j} dx_t f_t(x_t)}$$



# Response Matrix (IV)

In other words:

$$R_{ij} = \text{Prob}(\text{observed in bin } i | \text{true in bin } j)$$

Obviously, summing the response matrix over  $i$  gives the efficiency:

$$\sum_{i=1}^N R_{ij} = \varepsilon_j$$

In compact matrix form (including background):

$$\nu_i = \sum_{j=1}^M R_{ij} \mu_j + \beta_i \quad \vec{\nu} = R \vec{\mu} + \vec{\beta}$$

Response matrix depends on  $f_t(x_t)$  which we want to know. However, if we make the bins small enough  $f_t(x_t) \approx \text{const.}$  within a bin and drops from the ratio:

$$R_{ij} = \frac{\int_{\text{bin } i} dx_m \int_{\text{bin } j} dx_t r(x_m | x_t) \varepsilon(x_t) f_t(x_t)}{\int_{\text{bin } j} dx_t f(x_t)} \approx \frac{1}{\Delta x_{t,j}} \int_{\text{bin } i} dx_m \int_{\text{bin } j} dx_t r(x_m | x_t) \varepsilon(x_t)$$

# Unfolding by Inverting Response Matrix (I)

We have

$$\vec{v} = R\vec{\mu} + \vec{\beta}$$

Replace  $\vec{v}$  by  $\vec{n}$  to obtain an obvious estimator for the true distribution:

$$\hat{\vec{\mu}} = R^{-1}(\vec{n} - \vec{\beta})$$

This solution minimizes

$$\chi^2(\vec{\mu}) = (\vec{v}(\vec{\mu}) - \vec{n})^T V^{-1}(\vec{v}(\vec{\mu}) - \vec{n}) \quad \text{where} \quad V_{i,j} = \text{cov}[n_i, n_j]$$

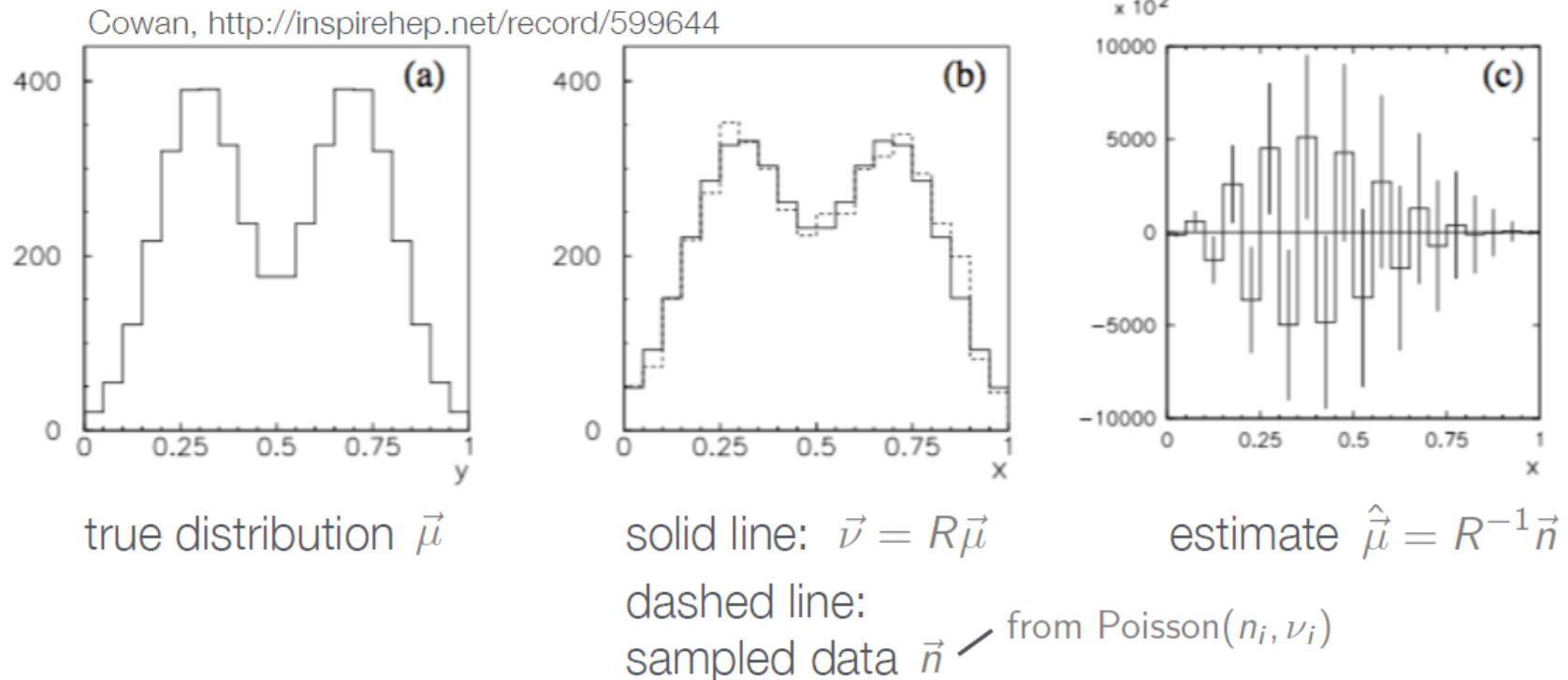
It can be shown that the covariance matrix of the solution is given by

$$U = R^{-1}V(R^{-1})^T$$

# Unfolding by Inverting Response Matrix (II)

It can also be shown that matrix inversion is unbiased and has minimal variance.

This sounds good ... let's try it.

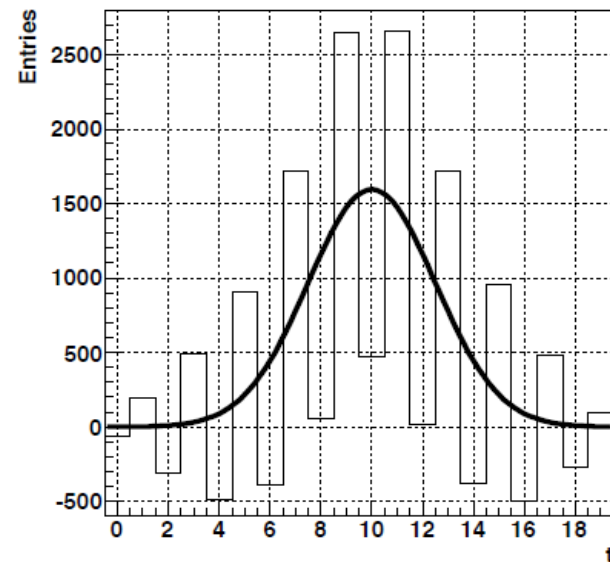
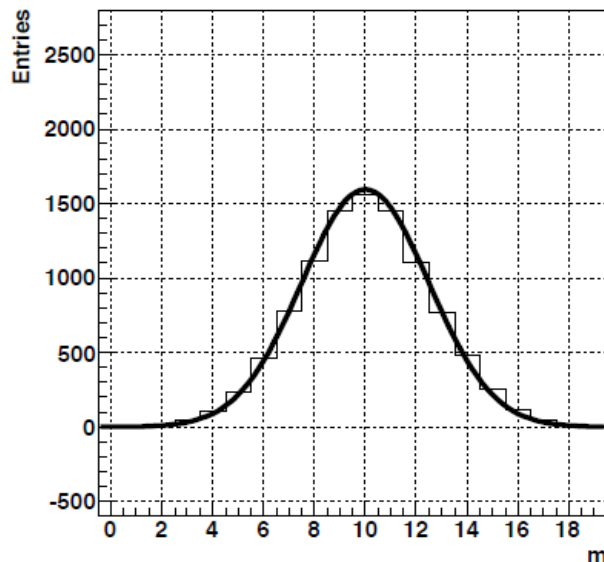


This looks like a disaster ... unfolded distribution very different from the true one

# Unfolding by Inverting Response Matrix (III)

Another example:

$$R = \begin{pmatrix} 0.75 & 0.25 & 0 & \dots \\ 0.25 & 0.50 & 0.25 & 0 \\ 0 & 0.25 & 0.50 & 0.25 \\ & & 0 & 0.25 & 0.50 \\ \vdots & & & & \ddots \end{pmatrix}.$$



Same conclusion: we don't get the desired (smooth) answer

# What's Wrong with the Matrix Inversion Method?

Unbiased, minimum variance, actually also a ML estimator ... all very nice!

The result is not wrong, it is just not desirable

- ▶ Does not really look like the original distribution
- ▶ Large correlation between bins

"Applying the response matrix  $R$  smears out fine structure  
→ applying  $R^{-1}$  creates (usually unwanted) structure"

More desirable solution by adding (smoothness) constraints.  
However, this will produce a bias.

The art of unfolding is to find an acceptable balance between bias and smoothness.

# Bin-by-Bin Method (I)

Used very often, but has issues ...

Assume shape of true spectrum and determine correction factor for each bin (usually determined from Monte Carlo simulation):

$$\mu_i = C_i(n_i - \beta_i) \qquad C_i = \frac{\mu_i^{\text{MC}}}{\nu_i^{\text{MC}}}$$

Works if smearing (bin-to-bin sharing) is negligible, only loss due to finite efficiency:

$$R_{ij} \approx \delta_{ij}\epsilon_j$$

Obviously works, too, if MC = nature.

Expectation value for corrected data:

$$E[\hat{\mu}_i] = C_i E[n_i - \beta_i] = C_i(\nu_i - \beta_i) \equiv C_i \nu_i^{\text{sig}}$$

# Bin-by-Bin Method (II)

Inserting the  $C_i$ 's one can determine the bias:

$$E[\hat{\mu}_i] = \frac{\mu_i^{\text{MC}}}{\nu_i^{\text{MC}}} \nu_i^{\text{sig}} = \underbrace{\left( \frac{\mu_i^{\text{MC}}}{\nu_i^{\text{MC}}} - \frac{\mu_i}{\nu_i^{\text{sig}}} \right)}_{\text{bias}} \nu_i^{\text{sig}} + \mu_i$$

no bias only if  
MC = nature

Covariance matrix of the corrected data (smearing fluctuations independent between bins)

$$U_{ij} = \text{cov}[\hat{\mu}_i, \hat{\mu}_j] = C_i C_j \underbrace{\text{cov}[n_i^{\text{sig}}, n_j^{\text{sig}}]}_{0 \text{ for } i \neq j} = C_i^2 \text{Var}[n_i^{\text{sig}}] \delta_{ij}$$

Iterative bin-by-by method

- ▶ Start with plausible guess of true spectrum
- ▶ Apply correction to measurement
- ▶ Generate new correction factors from corrected spectrum of previous iteration
- ▶ And so on ... usually a few iterations sufficient

# Regularized Unfolding

Matrix inversion is the maximum likelihood solution:

Independent Poisson fluctuations:

$$\ln L(\vec{\mu}) = \sum_{i=1}^M (n_i \ln \nu_i - \nu_i)$$

ML estimator:

$$\begin{aligned} \hat{\nu} &= \vec{n} \\ \rightarrow \hat{\mu} &= R^{-1}(\vec{n} - \vec{\beta}) \end{aligned}$$

Idea: accept solutions that are close to maximum likelihood estimate:

$$\ln L(\vec{\mu}) \geq \ln L(\vec{\mu}_{\max}) - \Delta \ln L(\vec{\mu})$$

Define a smoothness function  $S$  that gets bigger when the unfolded solution becomes smoother.

The task then is to maximize

$$\Phi(\vec{\mu}) = \alpha \ln L(\vec{\mu}) + S(\mu)$$

$\alpha$  depends on  $\Delta \ln \vec{\mu}$ ,  
 $\alpha \rightarrow \infty$  give ML solution

smoothness function



# Tikhonov Regularization

Measure of smoothness = mean square of  $k$ -th derivative of deconvoluted function  $f$ :

$$S[f] = - \int dx \left( \frac{d^k f}{d^k x} \right)^2 \quad k = 1, 2, 3, \dots$$

Minus sign makes  $S$  big when derivative is small

Tikhonov for  $k = 2$  with  $\log L = -\chi^2/2$ :

$$S(\vec{\mu}) = - \sum_{i=1}^{M-2} (-\mu_i + 2\mu_{i+1} - \mu_{i+2})^2$$

Implementation by A. Höcker, V. Kartvelishvili: *Singular Value Decomposition*  
(NIM A372 (1996) 469, hep-ph/9509307, TSVDUnfold in ROOT)

Minimizes  $\chi^2(\vec{\mu}) + \tau \sum_i [(\mu_{i+1} - \mu_i) - (\mu_i - \mu_{i-1})]^2$

Advice on how to choose  $\tau$  in the paper