Presentation and discussion about the energy measurement in the ECAL

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Problem: existence of the guard-rings



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Selection of the electrons

Obtained from the shape of the showers $(\sigma_x, \sigma_y \text{ and } \sigma_z)$



Effect of the guard-rings (drawings of the profiles)





Correction in Y

Fit : Plateau (1 parameter) – Gaussian (3 parameters) $[3]-[0] \cdot e^{-\frac{(Y-[1])^2}{2 \cdot [2]^2}}$ (where parameter [2] = RMS)

> Fonction of correction: (only 3 parameters)

$$\frac{[3]}{[3]-[0]\cdot e^{-\frac{(Y-[1])^2}{2.[2]^2}}} = \frac{1}{1-[0']\cdot e^{-\frac{(Y-[1])^2}{2.[2]^2}}}$$

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Layer per layer correction



Y Immi

Remarks:

In the previous histograms : Y = Y barycenter calculated in the studied layer Energy = Energy deposited in the studied layer

It is not always possible to
fit the response of each
layer, for instance : layer # 0.0.14In that case, for these layers,
there will be no correction in
y.0.12Y.0.08



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Fitting parameters (45 GeV)



With only 2 param. (\Rightarrow 1 param. for the corrective function)





Comparisons: 45 GeV, 30 GeV and 15 GeV (layer/layer)

20

30 Layer#

Layer #













Results: 45 GeV, 30 GeV and 15 GeV (selected region in X)





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Layer per layer correction in X (4 parameters)



Remarks:



Layer per layer correction in X (2 parameters)









Comparison with simulations

Data/Simulation, 15 GeV

Data/Simulation, 15 GeV

🗕 Data

(4 param.)

Simulation

(4 param.)

20 25

Layer #

📥 Data

Data

25

(%

15

10

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¥210

Layer #

Layer #



Other methods: 1. virtual cells

o In order to estimate the energy lost in the inter-wafer gaps virtual cells are added o The virtual cell energy is calculated from the interpolation of the deposited energies in the 2 closest cells: $E_{virtual} = K \cdot (E_{i-1} + E_{i+1})$

o This method is quite simple, it leads to an energy resolution improvement but do not really or sufficiently correct the gap problem

Virtual cell method

o Actually the value of the factor K depends on the shower barycenter in each layer and therefore this method looses one of its big advantage

$$E_{virtual} = K(X_b, Y_b) \cdot \left(E_{i-1} + E_{i+1}\right)$$



But what happens if the pad size changes : i.e. $1 \text{ cm} \times 1 \text{ cm} \rightarrow 5 \text{ mm} \times 5 \text{ mm}$?

2. Spline interpolation method

o A spline is an interpolation defined piecewise by polynomials o A spline (actually, third spline polynomial term) has been used to make interpolation with each row and each column of cells





Interpolation of a gaussian distribution by a spline

o Again the energy resolution is improved, but the tail at low energy is not well corrected
o It gives results close to what was obtained with the virtual cell method

Conclusion

Effect of the guard rings can be corrected thanks to a correction method using the barycenter calculated layer per layer. But this technique requires lots of parameters and is a little « heavy »

Other simpler methods have been tested, but for the time being, they does not lead to a good improvement for 1 cm x 1 cm pads

MOKKA with 5 mm x 5mm pads is needed in order to test these methods in a more realistic case