
Subject: Loss of efficiency for electrons at $\theta \sim 22^\circ$, due to association failure in EMC

Posted by [Ermias](#) on Fri, 20 Feb 2015 16:10:26 GMT

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Dear all,

While doing simulations on electrons, I noticed a localized efficiency loss for electrons at around $\theta \sim 22^\circ$. After digging around a bit, I was able to pinpoint that it was due to electrons in this location not being associated to *any* cluster, even though there is a valid reconstructed cluster sitting near the electron's projection. I first started to notice this problem in oct.14 release. Even though the efficiency drop with oct.14 was localized in a relatively smaller zone ($\sim 1^\circ$ window), the effect on the signal I was simulating was significant ($\sim 10\%$) because the electrons for this signal peak around 20° in θ . However with the current development version (26841) the loss in efficiency is striking (see attached figure, left panel, count of all electrons vs electrons with eid vs. θ). The efficiency loss is there for positrons too.

I looked at the change in the EMC association code and the only significant change that happened between apr.13 release and current trunk is the addition of the following conditions before starting the cluster association:

```
if ( (emcModule<3) && (helix->GetZ())>150.) ) continue; // not consider tracks after emc barrel for BARREL
if ( (emcModule==3) && (helix->GetZ())<165.) ) continue; // consider tracks only from last gem plane for FWD
if ( (emcModule==4) && (helix->GetZ())>-30.) ) continue; // consider tracks only ending at the back of STT for BKW
```

at L47 of PndPidEmcInfo.cxx. I assume these lines are there for a reason (would appreciate to hear from EMC experts why...), but I was able to recover most of the loss in efficiency by commenting them out (right panel). Could it be that the actual cut values are not correctly set?

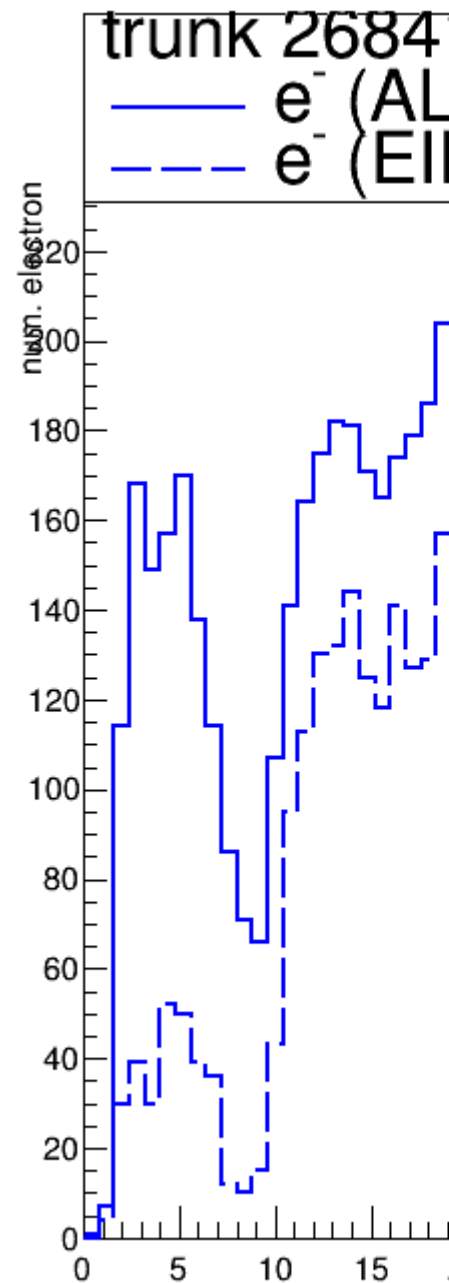
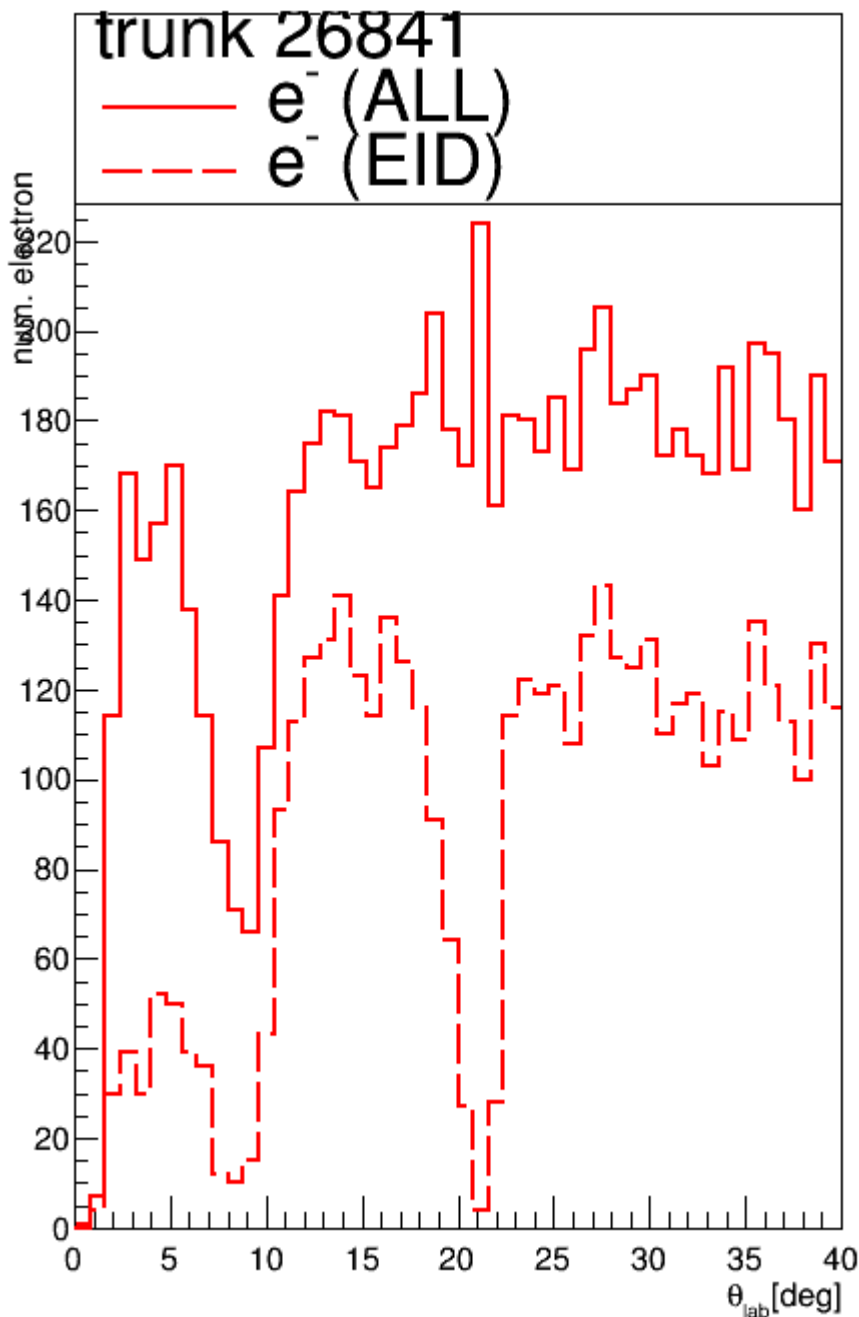
What fix do EMC experts suggest? Maybe its a known issue and people are working on it, but for "mass" simulation, would it be advisable to just go back and patch oct.14 version? or wait until a new release that includes fixes? What would be the approximate time scale for the next release, if it is okay to ask?

Thanks in advance!

Ermias.

File Attachments

1) [tc.png](#), downloaded 943 times



Subject: Re: Loss of efficiency for electrons at $\theta \sim 22^\circ$, due to association failure in EMC

Posted by [Stefano Spataro](#) on Fri, 20 Feb 2015 20:58:20 GMT

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Nobody of EMC has ever worked on the track-EMC correlation, you are asking to the wrong

guys.

```
if ( (emcModule<3) && (helix->GetZ())>150.) ) continue; // not consider tracks after emc barrel
for BARREL
if ( (emcModule==3) && (helix->GetZ())<165.) ) continue; // consider tracks only from last gem
plane for FWD
if ( (emcModule==4) && (helix->GetZ())>-30.) ) continue; // consider tracks only ending at the
back of STT for BKW
```

The lines are well commented, and they explain why they were put there. helix is the track parameters at the last point of the track. Since these are geometrical selections, in theory they should work. Which is the EMC module which is suffering from that drop? A check with MC id could help (but you need to use recent trunk since in oct14 the MC for EMC was bugged).

Subject: Re: Loss of efficiency for electrons at $\theta \sim 22^\circ$, due to association failure in EMC

Posted by [Ermias](#) on Fri, 20 Feb 2015 23:04:30 GMT

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Hi Stefano,

Sorry, I made the wrong assumption about who's working on that part of the code. I didn't mean to offend anyone and I should have known better to check...

For tracks that fail to get associated (emcIndex<0 after the loop over emcHits), I printed out the track MC index, its energy calculated from tracking using pion hypothesis together with the module number and energy of the emcHit with the closest energy to the track.

I only printed out a few hundred events, but It seems like module 3 is contributing to all of the unintended misses in the events I checked. Please let me know if I can provide any other useful feedback...

ps: Do you advise against using oct.14 for any simulation that uses the EMC?

Cheers,
Ermias.

pidCandMcIndex= 0	:	trackEnergy= 0.849534	Module= 3	emcHitEnergy= 0.740053
pidCandMcIndex= 0	:	trackEnergy= 4.03095	Module= 3	emcHitEnergy= 3.51169
pidCandMcIndex= 0	:	trackEnergy= 4.97663	Module= 3	emcHitEnergy= 4.8357
pidCandMcIndex= 0	:	trackEnergy= 3.74218	Module= 3	emcHitEnergy= 3.74057
pidCandMcIndex= 0	:	trackEnergy= 2.5047	Module= 3	emcHitEnergy= 2.40757
pidCandMcIndex= 0	:	trackEnergy= 0.405984	Module= 3	emcHitEnergy= 0.347898
pidCandMcIndex= 0	:	trackEnergy= 1.16359	Module= 3	emcHitEnergy= 1.08486
pidCandMcIndex= 0	:	trackEnergy= 2.81498	Module= 3	emcHitEnergy= 2.69472
pidCandMcIndex= 1084	:	trackEnergy= 0.22849	Module= 3	emcHitEnergy= 0.36478
pidCandMcIndex= 1083	:	trackEnergy= 0.428764	Module= 3	emcHitEnergy= 0.36478
pidCandMcIndex= 0	:	trackEnergy= 2.87692	Module= 3	emcHitEnergy= 2.68255

pidCandMcIndex= 0	:	trackEnergy= 1.8808	Module= 3	emcHitEnergy= 1.69046
pidCandMcIndex= 0	:	trackEnergy= 1.2923	Module= 3	emcHitEnergy= 1.2199
pidCandMcIndex= 0	:	trackEnergy= 3.45425	Module= 3	emcHitEnergy= 3.62943
pidCandMcIndex= 0	:	trackEnergy= 4.53307	Module= 3	emcHitEnergy= 3.92069
pidCandMcIndex= 0	:	trackEnergy= 3.95271	Module= 3	emcHitEnergy= 3.83431
pidCandMcIndex= 0	:	trackEnergy= 2.07854	Module= 3	emcHitEnergy= 3.70188
pidCandMcIndex= 0	:	trackEnergy= 0.840579	Module= 3	emcHitEnergy= 0.816676
pidCandMcIndex= 0	:	trackEnergy= 3.44526	Module= 3	emcHitEnergy= 3.43316
pidCandMcIndex= 0	:	trackEnergy= 4.48627	Module= 3	emcHitEnergy= 4.15238
pidCandMcIndex= 0	:	trackEnergy= 3.05255	Module= 3	emcHitEnergy= 3.01602
pidCandMcIndex= 0	:	trackEnergy= 1.46736	Module= 3	emcHitEnergy= 0.845704
pidCandMcIndex= 0	:	trackEnergy= 1.70518	Module= 3	emcHitEnergy= 1.62284
pidCandMcIndex= 0	:	trackEnergy= 1.37598	Module= 3	emcHitEnergy= 1.33291
pidCandMcIndex= 0	:	trackEnergy= 2.54198	Module= 3	emcHitEnergy= 3.89186
pidCandMcIndex= 0	:	trackEnergy= 4.27216	Module= 3	emcHitEnergy= 4.15942
pidCandMcIndex= 0	:	trackEnergy= 1.54658	Module= 3	emcHitEnergy= 1.48835
pidCandMcIndex= 0	:	trackEnergy= 3.80585	Module= 3	emcHitEnergy= 3.40713
pidCandMcIndex= 0	:	trackEnergy= 3.73259	Module= 3	emcHitEnergy= 3.56458
pidCandMcIndex= 0	:	trackEnergy= 0.898616	Module= 3	emcHitEnergy= 0.949504
pidCandMcIndex= 0	:	trackEnergy= 1.25923	Module= 3	emcHitEnergy= 0.920801
pidCandMcIndex= 0	:	trackEnergy= 0.463938	Module= 3	emcHitEnergy= 0.0957954
pidCandMcIndex= 0	:	trackEnergy= 2.92428	Module= 3	emcHitEnergy= 3.71379
pidCandMcIndex= 0	:	trackEnergy= 0.611837	Module= 3	emcHitEnergy= 0.546316
pidCandMcIndex= 0	:	trackEnergy= 4.05194	Module= 3	emcHitEnergy= 4.24035
pidCandMcIndex= 0	:	trackEnergy= 0.40836	Module= 3	emcHitEnergy= 0.320374
pidCandMcIndex= 349	:	trackEnergy= 0.262772	Module= 3	emcHitEnergy= 0.130423
pidCandMcIndex= 0	:	trackEnergy= 4.78844	Module= 3	emcHitEnergy= 4.82905
pidCandMcIndex= 0	:	trackEnergy= 0.384974	Module= 3	emcHitEnergy= 0.215389
pidCandMcIndex= 0	:	trackEnergy= 1.66274	Module= 3	emcHitEnergy= 1.65131
pidCandMcIndex= 1	:	trackEnergy= 0.212189	Module= 3	emcHitEnergy= 0.0214177

Subject: Re: Loss of efficiency for electrons at theta~22^deg, due to association failure in EMC

Posted by [StefanoSpataro](#) on Fri, 27 Feb 2015 11:04:55 GMT

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Oct14 suffers for a problem of MC truth for the neutrals, but apart from this the release is fine. All the lines you write come from geometrical considerations:

```
if ( (emcModule<3) && (helix->GetZ())>150.) ) continue; // not consider tracks after emc barrel for BARREL
```

If the position of the last hit is in the GEMS then most probably the will not hit the barrel, then skip this correlation

```
if ( (emcModule==3) && (helix->GetZ())<165.) ) continue; // consider tracks only from last gem plane for FWD
```

Consider only the tracks with are using the last GEM plane for the propagation to the forward endcap.

```
if ( (emcModule==4) && (helix->GetZ())>-30.) ) continue; // consider tracks only ending at the back of STT for BKW
```

If the last hit is not in the negative Z then it will not go to the backward endcap.

In theory, all these conditions make sense. BUT, maybe, if you suffer from a lack of counts for module 3, the 2nd command is somehow wrong, maybe not all the tracks hit the last plane of the GEM (problems in tracking). It could make sense to check the geometry of not GEMs and EMC to see how far we are in this "edge" region of 22°.
