

MEM Calculators

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1 Namespace Index

1.1 Namespace List

Here is a list of all documented namespaces with brief descriptions:

MEMNames **1**

2 Class Index

2.1 Class List

Here are the classes, structs, unions and interfaces with brief descriptions:

MEMs **2**

3 Namespace Documentation

3.1 MEMNames Namespace Reference

Enumerations

- enum [Processes](#) {
kSMHiggs, k0hplus, k0minus, k1plus,
k1minus, k2mplus_gg, k2mplus_qqbar, kqqZZ,
kggZZ, NUM_PROCESSES }

Enum type for supported processes in MELA and MEKD packages.

- enum `MEMCalcs` {
`kAnalytical`, `kMEKD`, `kJHUGen`, `kMCFM`,
`kMELA_HCP`, `NUM_MEMCALCS` }

Enum type for supported MEM calculators from MELA and MEKD packages.

3.1.1 Detailed Description

`MEMNames` namespace provides enum types for names of processes and names of tools/calculators supported by MELA and MEKD packages.

4 Class Documentation

4.1 MEMs Class Reference

Public Member Functions

- `MEMs` (double *collisionEnergy*=8, string *PDFName*="", bool *debug_*=false)
 - int `computeME` (`Processes` process, `MEMCalcs` calculator, vector< `TLorentzVector` > *partP*, vector< int > *partId*, double &*me2process*)
 - int `computeKD` (`Processes` processA, `Processes` processB, `MEMCalcs` calculator, vector< `TLorentzVector` > *partP*, vector< int > *partId*, double &*kd*, double &*me2processA*, double &*me2processB*)
 - int `computeMEs` (vector< `TLorentzVector` > *partP*, vector< int > *partId*)
 - int `retrieveME` (`Processes` process, `MEMCalcs` calculator, double &*me2process*)
 - int `computeKD` (`Processes` processA, `MEMCalcs` calculatorA, `Processes` processB, `MEMCalcs` calculatorB, double(MEMs::**funcKD*)(double, double), double &*kd*, double &*me2processA*, double &*me2processB*)
 - double `logRatio` (double *me2processA*, double *me2processB*)
 - double `probRatio` (double *me2processA*, double *me2processB*)
- Supproted KD functions, $kd = f_{KD}(me2processA, me2processB)$.*

Static Public Attributes

- static const bool `isProcSupported` [`NUM_PROCESSES`][`NUM_MEMCALCS`]
- Matrix of supproted processes.*

4.1.1 Constructor & Destructor Documentation

4.1.1.1 MEMs::MEMs (double *collisionEnergy* = 8, string *PDFName* = " ", bool *debug_* = false)

Constructor. Can specify the PDF to be use (ony CTEQ6L available at the moment).

Parameters

<i>collisionEnergy</i>	the sqrt(s) value in TeV (DEFAULT = 8).
<i>PDFName</i>	the name of the parton density functions to be used (DEFAULT = "", Optional: "CTEQ6L").

4.1.2 Member Function Documentation

4.1.2.1 int MEMs::computeME (`Processes` process, `MEMCalcs` calculator, vector< `TLorentzVector` > *partP*, vector< int > *partId*, double & *me2process*)

Compute individual ME for the specified process.

Parameters

in	<i>process</i>	names of the process for which the ME should be retrieved (REQUIRED).
in	<i>calculator</i>	name of the calculator tool to be used (REQUIRED).
in	<i>partP</i>	the input vector with TLorentzVectors for 4 leptons and 1 photon (REQUIRED).
in	<i>partId</i>	the input vecor with IDs (PDG) for 4 leptons and 1 photon (REQUIRED).
out	<i>me2process</i>	retrieved $ ME ^2$ for the specified process and calculator.

Returns

error code of the computation: 0 = NO_ERR, 1 = ERR_PROCESS, 2 = ERR_COMPUTE

4.1.2.2 `int MEMs::computeKD (Processes processA, Processes processB, MEMCalcs calculator, vector< TLorentzVector > partP, vector< int > partId, double & kd, double & me2processA, double & me2processB)`

Compute individual KD and MEs for process A and process B, obtained with the specified calculator tool.

Parameters

in	<i>processA,processB</i>	names of the processes A and B for which the KDs and MEs are computed (REQUIRED).
in	<i>calculator</i>	name of the calculator tool to be used (REQUIRED).
in	<i>partP</i>	the input vector with TLorentzVectors for 4 leptons and 1 photon (REQUIRED).
in	<i>partId</i>	the input vecor with IDs (PDG) for 4 leptons and 1 photon (REQUIRED).
out	<i>kd</i>	computed KD value for discrimination of processes A and B.
out	<i>me2processA</i>	computed $ ME ^2$ for process A.
out	<i>me2processB</i>	computed $ ME ^2$ for process B.

Returns

error code of the computation: 0 = NO_ERR, 1 = ERR_PROCESS, 2 = ERR_COMPUTE

4.1.2.3 `int MEMs::computeMEs (vector< TLorentzVector > partP, vector< int > partId)`

Compute MEs for all supported processes.

Individual MEs and KDs can be retrieved using [retrieveME\(Processes,MEMCalcs,double&\)](#) and [computeKD\(Processes,MEMCalcs,Processes,MEMCalcs,double*\)\(double,double\),double&,double&,double&\)](#).

Parameters

in	<i>partP</i>	the input vector with TLorentzVectors for 4 leptons and 1 photon (REQUIRED).
in	<i>partId</i>	the input vecor with IDs (PDG) for 4 leptons and 1 photon (REQUIRED).

Returns

error code of the computation: 0 = NO_ERR, 1 = ERR_PROCESS, 2 = ERR_COMPUTE

4.1.2.4 `int MEMs::retrieveME (Processes process, MEMCalcs calculator, double & me2process)`

Retrieve ME for specified process and specified calculator tool.

Method should be called only after running [computeMEs\(vector<TLorentzVector> partP,vector<int> partId\)](#).

Parameters

in	<i>process</i>	names of the process for which the ME should be retrieved (REQUIRED).
in	<i>calculator</i>	name of the calculator tool to be used (REQUIRED).
out	<i>me2process</i>	retrieved $ ME ^2$ for the specified process and calculator.

Returns

error codes: 0 = NO_ERR, 1 = ERR_PROCESS

4.1.2.5 `int MEMs::computeKD (Processes processA, MEMCalcs calculatorA, Processes processB, MEMCalcs calculatorB, double(MEMs::*)(double, double) funcKD, double & kd, double & me2processA, double & me2processB)`

Compute KD and retrieve MEs for process A and process B, obtained with the specified calculator tool. The KD is computed using KD function specified by the user as `kd = funcKD(me2processA, me2processB)`.

Method should be called only after running `computeMEs(vector<TLorentzVector> partP,vector<int> partId)`.

Parameters

in	<i>process-A,processB</i>	names of the processes for which the KD and MEs are computed (REQUIRED).
in	<i>calculator-A,calculatorB</i>	names of the calculator tools to be used (REQUIRED).
in	<i>funcKD</i>	name of the function to be used for KD computation (REQUIRED).
out	<i>kd</i>	computed KD value for discrimination of processes A and B.
out	<i>me2processA</i>	computed $ ME ^2$ for process A.
out	<i>me2processB</i>	computed $ ME ^2$ for process B.

Returns

error code of the computation: 0 = NO_ERR, 1 = ERR_PROCESS

4.1.3 Member Data Documentation

4.1.3.1 `const bool MEMs::isProcSupported` `[static]`

Initial value:

```
= {
    {1,      1,      1,      1,      1},
    {1,      1,      1,      0,      0},
    {1,      1,      1,      0,      0},
    {1,      0,      1,      0,      0},
    {1,      0,      1,      0,      0},
    {1,      1,      1,      0,      0},
    {1,      0,      1,      0,      0},
    {1,      1,      0,      1,      1},
    {0,      0,      0,      1,      0}}
```

Matrix of supported processes.

Matrix of supported processes - initialisation (to be updated)

The documentation for this class was generated from the following file:

- MEMCalculators.h

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