The particle diagnostic in SMILEI

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The particle diagnostic collects data from the macro-particles and processes them during runtime. It does <u>not</u> provide information on individual particles: instead, it produces <u>averaged quantities</u> like the particle density, currents, etc.

The data may be collected from one or several particle species.

The data is discretized inside a "grid" chosen by the user. This grid may be of any dimension. Examples:

- 1-dimensional grid along the position x (gives density variation along x)
- 2-dimensional grid along positions x and y (gives density map)
- 1-dimensional grid along the velocity v_x (gives the velocity distribution)
- 2-dimensional grid along position x and momentum p_x (gives the phase-space)
- 1-dimensional grid along the kinetic energy $E_{\rm kin}$ (gives the energy distribution)
- 3-dimensional grid along x, y and E_{kin} (gives the density map for several energies)
- 1-dimensional grid along the charge Z^* (gives the charge distribution)

Each dimension of the grid is called "axis".

The user may choose to average the data over several time-steps.

1 How to add a particle diagnostic in the input file

In the input file, you can add a particle diagnostic with the following syntax.

```
diagnostic particles
    output = output
    every = every
    time_average = time_average
    species = species1 species2 ...
    axis = type min max nsteps [logscale] [edge_inclusive]
    axis = type min max nsteps [logscale] [edge_inclusive]
    ...
end
```

- output must be set to density, charge_density, current_density_x, current_density_y, current_density_z. It determines the data that is summed in each cell of the grid. In the case of density, the weights are summed. In the case of charge_density, the weights×charge are summed. In the case of current_density_x, the weights×charge×v_x are summed (etc.)
- every must be a positive integer. It is the number of time-steps between each output.
- time_average must be a positive integer. It is the number of time-steps during which the data is averaged before output.
- species1 species2 ... must be one or several species. Species are recognized by their parameter species_type that is included in each group species ... end
- axis is an argument that describes one axis of the grid.

 Syntax: axis = type min max nsteps [logscale] [edge_inclusive]

 type can be x, y, z, px, py, pz, p, gamma, ekin, vx, vy, vz, v or charge.

 The axis is discretized for type from min to max in nsteps bins.

 The optional keyword logscale sets the axis scale to logarithmic instead of linear.

 The optional keyword edge_inclusive includes the particles outside the range [min, max] into the extrema bins.

There may be as many axis arguments as wanted in one diagnostic particles block.

There may be as many diagnostic particles blocks as wanted in the input file.

In the input file, you may add the following block as a reminder.

```
# DIAGNOSTICS ON PARTICLES - project the particles on a N-D arbitrary grid
# -----
# output = density, charge_density or current_density_[xyz]
              => parameter that describes what quantity is obtained
              => integer > 0 : number of time-steps between each output
# every
# time_average => integer > 0 : number of time-steps to average
# species
             => list of one or several species whose data will be used
       = _type_ _min_ _max_ _nsteps_ [logscale] [edge_inclusive]
# axis
              => _type_ can be one of the following:
                  x, y, z, px, py, pz, p, gamma, ekin, vx, vy, vz, v or charge
#
#
              => the data is discretized for _type_ between _min_ and
#
                  _max_, in _nsteps_ bins
#
              => the optional [logscale] sets the scale to logarithmic
#
              => the optional [edge_inclusive] forces the particles
#
                  outside (_min_,_max_) to be counted in the extrema bins
#
   example : axis = x \cdot 0 \cdot 1 \cdot 30
   example : axis = px -1 1 100
# >>>> MANY AXES CAN BE ADDED IN A SINGLE DIAGNOSTIC <><<
```

2 Examples

Variation of the density of species electron from x = 0 to 1, every 5 time-steps, without time-averaging.

Density map from x = 0 to 1, y = 0 to 1.

Velocity distribution from $v_x = -0.1$ to 0.1.

```
diagnostic particles
    output = density
    every = 5
    time_average = 1
    species = electron1
    axis = vx -0.1 0.1 100
end
```

Phase space from x = 0 to 1 and from px = -1 to 1.

Energy distribution from 0.01 to 1 MeV in logarithmic scale. Note that the input units are $m_e c^2 \sim 0.5$ MeV.

```
diagnostic particles
    output = density
    every = 5
    time_average = 1
    species = electron1
    axis = ekin  0.02  2  100  logscale
end
```

x-y density maps for three bands of energy: [0,1], [1,2], $[2,\infty]$. Note the use of edge_inclusive to reach energies up to ∞ .

Charge distribution from $Z^* = 0$ to 10.

```
diagnostic particles
    output = density
    every = 5
    time_average = 1
    species = electron1
    axis = charge    -0.5    10.5    11
end
```

3 How to view and post-process a particle diagnostic

Each diagnostic produces one file like ParticleDiagnostic0.h5, ParticleDiagnostic1.h5, etc. They are numbered the same way as the order of appearance in the input file.

A python script *ParticleDiagnostic.py* is provided to view or extract data from these files. To run this script, you will need *python2.7* with the following packages: numpy, matplotlib, pylab, h5py.

First, run python and include the script:

```
python -i ParticleDiagnostic.py
```

(Alternately, you can include this file into your own script using, for example, execfile("scripts/ParticleDiagnostic.py"))

In python, you can run ParticleDiagnostic(...) to extract the data and optionally plot it.

Syntax of the function *ParticleDiagnostic*:

```
ParticleDiagnostic(results_path, diagNumber=None, timesteps=None, slice=None, units="code", data_log=False, data_min=None, data_max=None, xmin=None, xmax=None, ymin=None, ymax=None, figure=None)
```

- results_path = _string_
 Path to the directory where the outputs are stored.

 (Also, this has to contain one and only one input file *.in)
- diagNumber = _int_ (optional)

 Number of the diagnostic. The first diagnostic has number 0.

 If not given, then a list of available diagnostics is printed.
- timesteps = _int_ (optional)

 timesteps = [_int_, _int_] (optional)

 If omitted, all timesteps are used.

 If one number given, the nearest timestep available is used.

 If two numbers given, all the timesteps in between are used.
- slice = { axis : "all", ... } (optional)
 slice = { axis : _double_, ... } (optional)
 slice = { axis : [_double_, _double_], ... } (optional)
 This parameter is used to reduce the number of dimensions of the array.
 axis must be "x", "y", "z", "px", "py", "pz", "p", "gamma", "ekin", "vx", "vy",
 "vz", "v" or "charge".

Any axis of the same name will be removed with the following technique:

- If the value is "all", then a sum is performed over all the axis.
- If the value is **_double_**, then only the bin closest to the value is kept.
- If the value is [_double_,_double_], then a sum is performed between the two values.

Example: $\{"x":[4,5]\}$ will sum all the data for x in the range [4,5].

• units = "nice" (optional)

If "nice" is chosen, then units are converted into usual units.

Distances in microns, density in cm⁻³, energy in MeV.

- data_log = True or (False) (optional)

 If True, then log 10 is applied to the output array before plotting.
- data_min = _double_ (optional)
 data_max = _double_ (optional)
 If present, output is rescaled before plotting.

• xmin = _double_ (optional)
xmax = _double_ (optional)
ymin = _double_ (optional)

ymax = _double_ (optional)

If present, axes are rescaled before plotting.

• figure = None (default)
figure = _int_ (optional)

Choses the figure number that is passed to matplotlib.

If absent or None, returns the first data without plotting.

Output of the function ParticleDiagnostic

If **figure** is a number, then the function does not return anything. Instead, it displays the results in a graph.

If figure is not provided, or figure=None, then the function returns the results as a python dictionary. The dictionary is of the form {"data": data_array, axis1:axis1_array, axis2:axis2_array, ...} where:

- data_array is the final array containing the data.
- axis1 and axis2 are the names of the axes (for example "x" or "px").
- axis1_array and axis2_array are the locations of the axes bins.

Examples of usage from the python prompt:

```
>>> ParticleDiagnostic("path/to/my/results", diagNumber=1, slice={"y":"all"}, units="nice",data_min=0, data_max=3e14, figure=1)
```

This will take the diagnostic #1, and sum the grid over all the y axis. Then it will plot the resulting array in figure 1 from 0 to 3e14.

```
>>> result = ParticleDiagnostic("path/to/my/results", diagNumber=3,
    slice={"ekin":[1,10]}, timesteps=1000)
```

This will take the diagnostic #3 for the timestep nearest to 1000, and sum for all energies between 1 and 10. The results are stored in the variable result.

The data can be accessed with result["data"].

If one of the axes is x, you can access the locations of the bins with result ["x"].