## DOS extraction

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## Index

## 1.1 Introduction

This program allows to extract the Density of States, assessed by mean capacitance-voltage measurements, in an organic semiconductor device. Simulated values are fitted to experimental data.

Source and header files are written in C++11 language.

The software is intended to be used on a Unix-like operating system.

## 1.2 Dependancies

The program requires the following libraries to be installed on your system:

- Eigen, to handle with matrices, vectors and linear algebra;
- Gnuplot, a graphical utility to generate plots;
- Boost, a C++ library used by the Gnuplot interface to C++;
- OpenMP, for parallel computing (recommended but not compulsory).

It also uses the following packages, provided in the include/folder:

- GetPot, to parse command-line and configuration files;
- gnuplot-iostream, the C++ interface for Gnuplot.

## 1.3 Compile

In order to compile a test executable, simply execute one of these commands in a terminal pointing to the root directory of this package:

```
$ make
```

or, if you want the compiler to produce also debugging informations:

```
$ make debug
```

You can specify the name of the test to be compiled (without extension, for example: *simulate\_dos*) by passing the variable **NAME** through command-line:

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```
$ make NAME=test_filename
```

The compiler will generate the test filename executable under the bin/directory.

Repeat these instructions for each test you want to compile.

## 1.4 Set up the configurations

Note

The default configuration directory is config/.

Before you can run an executable, you have to set up the configuration file (default: *config.pot*). Within it you can find a list of parameters, each of which is commented out to explain what modifying it will entail.

Particularly, the variables *input\_params* and *input\_experim* can be set, i.e. the filenames where to find input fitting parameters and experimental data respectively. It's recommended (but not compulsory) to put these files in the same directory as the configuration file (otherwise you can specify a relative or absolute path to them).

You can create multiple configuration files, each with different parameter values: the one you aim to use can be specified in the command-line before running.

## 1.5 Run!

Executables are placed under the bin/directory.

To run by using the default configuration filename (config.pot) simply move to the bin/directory and execute:

```
$ ./test_filename
```

To specify a different configuration file previously saved in the configuration directory:

```
$ ./test_filename -f configuration_filename

Or:
$ ./test_filename --file configuration_filename
```

The variable *configuration\_filename* should **not** contain the path.

## Warning

Furthermore, if you run the program from a different folder than *bin*/ or if you chose a different configuration directory, you have to manually specify the **full** path to the configuration directory (either absolute or relative to the current directory) by using:

```
$ ./test_filename -d configuration_directory

Or:
$ ./test_filename --directory configuration_directory
```

Once complete, you can find the results of the simulation(s) in the output directory specified in the configuration file (default: *output*/) under *bin*/.

# Namespace Index

## 2.1 Namespace List

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

constant	
	Numerical constants
numerics	<b>;</b>
	Namespace for generic numeric algorithms
utility	
	Namespace for utilities and auxiliary functions

Namespace Index

# **Hierarchical Index**

## 3.1 Class Hierarchy

This inheritance list is sorted roughly, but not completely, alphabetically:

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SsvParser	
OosModel	25
IonLinearPoisson1D	
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# **Class Index**

## 4.1 Class List

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

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Charge		
	Abstract class providing methods to calculate total electric charge (the rhs in the Poisson equa-	
	tion)	19
CsvPars	er	
	Class providing methods to read <b>numeric</b> content from a .csv file and to store it in Eigen matrices	
	or vectors	21
DosMod	el	
	Class providing methods to process a simulation to extract the Density of States starting from a	
	parameter list	25
GaussHe	ermiteRule	
	Class derived from QuadratureRule providing the Gauss-Hermite quadrature rule	28
Gaussiai	nCharge	
	Class derived from Charge, under the hypothesis that Density of States is a combination of	
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NonLine	arPoisson1D	
	Provide a solver for a non-linear Poisson equation	32
ParamLis	st	
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# File Index

## 5.1 File List

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

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# **Namespace Documentation**

## 6.1 constants Namespace Reference

Numerical constants.

### **Variables**

```
    const Real Q = 1.60217653000000e-19

     Electron charge [C].
• const Real Q2 = Q * Q
     Electron charge squared [C^2].

    const Real K B = 1.38065050000000e-23

     Boltzmann's constant [J \cdot K^{-1}].
• const Real EPS0 = 8.854187817e-12
      Vacuum electrical permittivity [C \cdot V^{-1} \cdot m^{-1}].
• const Real T = 300
     Reference temperature [K].
• const Real V_TH = K_B * T / Q
      Treshold voltage [V].
• const unsigned PARAMS_NO = 22
     Number of parameters required in input file.
• const Real PI = M_PI
const Real SQRT_PI = std::sqrt(PI)

    const Real PI_M4 = 0.7511255444649425

const Real SQRT_2 = std::sqrt(2)
     \sqrt{2}.
```

## 6.1.1 Detailed Description

Numerical constants.

## 6.2 numerics Namespace Reference

Namespace for generic numeric algorithms.

### **Functions**

template<typename ScalarType >

VectorX< ScalarType > sort (const VectorX< ScalarType > &vector)

Function to sort Eigen vectors.

template<typename ScalarType >

VectorXpair< ScalarType > sort\_pair (const VectorX< ScalarType > &vector)

Function to sort Eigen vectors, keeping track of indexes.

• Real trapz (const VectorXr &x, const VectorXr &y)

Function to compute approximate integral of y with spacing increment specified by x, using trapezoidal rule.

Real trapz (const VectorXr &y)

Compute the approximate integral of y with unit spacing, using trapezoidal rule.

VectorXr deriv (const VectorXr &, const VectorXr &)

Compute the numeric derivative:  $\frac{dy}{dx}$ .

• Real interp1 (const VectorXr &, const VectorXr &, const Real &)

Linear 1D interpolation. Interpolate y, defined at points x, at the point xNew.

VectorXr interp1 (const VectorXr &, const VectorXr &, const VectorXr &)

Linear 1D interpolation. Interpolate y, defined at points x, at the points xNew.

• Real error\_L2 (const VectorXr &, const VectorXr &, const VectorXr &, const Real &)

Compute the  $L^2$ -norm error between simulated and interpolated values, using trapz.

## 6.2.1 Detailed Description

Namespace for generic numeric algorithms.

### 6.2.2 Function Documentation

6.2.2.1 VectorX < ScalarType > sort ( const VectorX < ScalarType > & vector )

Function to sort Eigen vectors.

**Template Parameters** 

ScalarType	: the scalar type.
------------	--------------------

## **Parameters**

in	vector	: the vector to be sorted.
----	--------	----------------------------

## Returns

the sorted vector.

Definition at line 98 of file numerics.h.

6.2.2.2 VectorXpair < ScalarType > sort\_pair ( const VectorX < ScalarType > & vector )

Function to sort Eigen vectors, keeping track of indexes.

**Template Parameters** 

	ScalarType	: the scalar type.
Parameters		
in	vocto	r the vector to be corted

#### Returns

an Eigen vector of pairs: (sorted value, corresponding index in the unsorted vector).

Definition at line 107 of file numerics.h.

## 6.2.2.3 Real trapz (const VectorXr & x, const VectorXr & y)

Function to compute approximate integral of y with spacing increment specified by x, using trapezoidal rule.

#### **Parameters**

in	X	: the vector of the discrete domain;
in	у	: the vector of values to integrate.

#### Returns

the approximate integral value.

Definition at line 3 of file numerics.c++.

## 6.2.2.4 Real trapz (const VectorXr & y)

Compute the approximate integral of *y* with unit spacing, using trapezoidal rule.

## **Parameters**

in	У	: the vector of values to integrate.

## Returns

the approximate integral value.

Definition at line 18 of file numerics.c++.

## 6.2.2.5 VectorXr deriv (const VectorXr & y, const VectorXr & x)

Compute the numeric derivative:  $\frac{dy}{dx}$ .

## **Parameters**

in	у	: the vector of values to differentiate;
in	X	: the vector of the discrete domain.

## Returns

a vector of the same length as y containing the approximate derivative.

Definition at line 23 of file numerics.c++.

## 6.2.2.6 Real interp1 ( const VectorXr & x, const VectorXr & y, const Real & xNew )

Linear 1D interpolation. Interpolate *y*, defined at points *x*, at the point *xNew*.

#### **Parameters**

in	У	: the vector of values to interpolate;
in	X	: the vector of the discrete domain;
in	xNew	: the point to interpolate at.

## Returns

a scalar containing the interpolated value.

Definition at line 42 of file numerics.c++.

## 6.2.2.7 VectorXr interp1 (const VectorXr & x, const VectorXr & y, const VectorXr & xNew)

Linear 1D interpolation. Interpolate *y*, defined at points *x*, at the points *xNew*.

#### **Parameters**

in	у	: the vector of values to interpolate;
in	X	: the vector of the discrete domain;
in	xNew	: the vector of points to interpolate at.

### Returns

a vector of the same length as xNew containing the interpolated values.

Definition at line 61 of file numerics.c++.

## 6.2.2.8 Real error\_L2 ( const VectorXr & interp, const VectorXr & simulated, const VectorXr & V, const Real & V\_shift )

Compute the  $L^2$ -norm error between simulated and interpolated values, using *trapz*.

## **Parameters**

in	interp	: the interpolated values;
in	simulated	: the simulated values;
in	V	: the vector of the electric potential;
in	V_shift	: shift to the electric potential.

#### Returns

the value of the  $L^2$ -norm error.

Definition at line 75 of file numerics.c++.

## 6.3 utility Namespace Reference

Namespace for utilities and auxiliary functions.

## **Functions**

• std::string full\_path (const std::string &, const std::string &)

Auxiliary function to return the full path to a file.

void print\_block (const char \*, std::ostream &=std::cout)

Auxiliary function to print a string inside a block.

void print\_done (std::ostream &=std::cout)

Auxiliary function to print a "DONE!" string.

## 6.3.1 Detailed Description

Namespace for utilities and auxiliary functions.

## 6.3.2 Function Documentation

6.3.2.1 std::string full\_path ( const std::string & filename, const std::string & relative\_directory )

Auxiliary function to return the full path to a file.

### **Parameters**

in	filename	: the filename;
in	relative	: the directory for a relative path.
	directory	

### Returns

the variable *filename*, if it contains an absolute path; otherwise returns the concatenation of *relative\_directory* and *filename* (i.e. the relative path to *filename*).

Definition at line 3 of file typedefs.c++.

6.3.2.2 void print\_block ( const char \* string, std::ostream & os = std::cout )

Auxiliary function to print a string inside a block.

## **Parameters**

in	string	: the string to print;
out	os	: output stream.

Definition at line 8 of file typedefs.c++.

6.3.2.3 void print\_done ( std::ostream & os = std::cout )

Auxiliary function to print a "DONE!" string.

## **Parameters**

out	OS	: output stream.

Definition at line 31 of file typedefs.c++.

Namespace	Documer	ntation

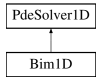
## **Class Documentation**

## 7.1 Bim1D Class Reference

Class derived from PdeSolver1D, providing a finite volume Box Integration Method (BIM) solver.

#include <solvers.h>

Inheritance diagram for Bim1D:



## **Public Member Functions**

• Bim1D ()=delete

Default constructor (deleted since it is required to specify the mesh).

• Bim1D (VectorXr &)

Constructor.

virtual ∼Bim1D ()=default

Destructor (defaulted).

- virtual void assembleAdvDiff (const VectorXr &, const VectorXr &, const VectorXr &, const VectorXr &, const VectorXr &) override

  Assemble the matrix for an advection-diffusion term.
- virtual void assembleStiff (const VectorXr &, const VectorXr &) override

Assemble the matrix for a diffusion term.

• virtual void assembleMass (const VectorXr &, const VectorXr &) override

Assemble the matrix for a reaction term.

## **Static Public Member Functions**

static VectorXr log\_mean (const VectorXr &, const VectorXr &)

Compute the element-wise logarithmic mean of two vectors.

• static std::pair< VectorXr,

VectorXr > bernoulli (const VectorXr &)

Compute the values of the Bernoulli function.

## **Additional Inherited Members**

## 7.1.1 Detailed Description

Class derived from PdeSolver1D, providing a finite volume Box Integration Method (BIM) solver.

Matrices are held in a sparse format.

Definition at line 111 of file solvers.h.

## 7.1.2 Constructor & Destructor Documentation

## 7.1.2.1 Bim1D ( VectorXr & mesh )

#### Constructor.

#### **Parameters**

in	mesh	: the mesh coordinates.

Definition at line 6 of file solvers.c++.

### 7.1.3 Member Function Documentation

7.1.3.1 VectorXr log\_mean ( const VectorXr & x1, const VectorXr & x2 ) [static]

Compute the element-wise logarithmic mean of two vectors.

$$M_{log}(x_1, x_2) = \frac{x_2 - x_1}{\log x_2 - \log x_1} = \frac{x_2 - x_1}{\log \left(\frac{x_2}{x_1}\right)}$$
.

## Parameters

in	x1	: the first vector;
in	x2	: the second vector.

#### Returns

the vector of the logarithmic means.

Definition at line 9 of file solvers.c++.

7.1.3.2 std::pair < VectorXr, VectorXr > bernoulli(const VectorXr & x) [static]

Compute the values of the Bernoulli function.

$$\mathfrak{B}(x) = \frac{x}{e^x - 1} \ .$$

## **Parameters**

in	X	: the vector of the values to compute the Bernoulli function at.

#### Returns

the pair 
$$(\mathfrak{B}(x),\mathfrak{B}(-x))$$
.

Definition at line 32 of file solvers.c++.

7.1.3.3 void assembleAdvDiff (const VectorXr & alpha, const VectorXr & gamma, const VectorXr & eta, const VectorXr & beta) [override], [virtual]

Assemble the matrix for an advection-diffusion term.

Build the Scharfetter-Gummel stabilized stiffness matrix for:  $-\nabla \cdot (\alpha \cdot \gamma(\eta \nabla u - \beta u)) = f$ .

#### **Parameters**

in	alpha	: $lpha$ , an element-wise constant function;
in	gamma	: $\gamma$ , an element-wise linear function;
in	eta	: $\eta$ , an element-wise linear function;
in	beta	: $eta$ , an element-wise constant function.

Implements PdeSolver1D.

Definition at line 79 of file solvers.c++.

7.1.3.4 void assembleStiff(const VectorXr & eps, const VectorXr & kappa) [override], [virtual]

Assemble the matrix for a diffusion term.

Build the standard finite element stiffness matrix for the diffusion problem:  $-\nabla \cdot (\varepsilon \cdot \kappa \nabla u) = f$ .

#### **Parameters**

in	eps	$: \mathcal{E},$ an element-wise constant function;
in	kappa	: $\kappa$ , an element-wise linear function.

Implements PdeSolver1D.

Definition at line 132 of file solvers.c++.

7.1.3.5 void assembleMass (const VectorXr & delta, const VectorXr & zeta) [override], [virtual]

Assemble the matrix for a reaction term.

Build the lumped finite element mass matrix for the reaction problem:  $\delta \cdot \zeta u = f$ .

### **Parameters**

in	delta	: $\delta$ , an element-wise constant function;
in	zeta	: $\zeta$ , an element-wise linear function.

Implements PdeSolver1D.

Definition at line 141 of file solvers.c++.

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

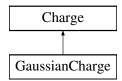
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/solvers.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/solvers.c++

## 7.2 Charge Class Reference

Abstract class providing methods to calculate total electric charge (the rhs in the Poisson equation).

#include <charge.h>

Inheritance diagram for Charge:



### **Public Member Functions**

• Charge ()=delete

Default constructor (deleted since it is required to specify a ParamList and a QuadratureRule).

Charge (const ParamList &, const QuadratureRule &)

Constructor.

virtual ∼Charge ()=default

Destructor (defaulted).

virtual VectorXr charge (const VectorXr &phi)=0

Compute the total charge.

• virtual VectorXr dcharge (const VectorXr &phi)=0

Compute the derivative of the total charge with respect to the electric potential.

## **Protected Attributes**

• const ParamList & params\_

Parameter list handler.

const QuadratureRule & rule

Quadrature rule handler.

## 7.2.1 Detailed Description

Abstract class providing methods to calculate total electric charge (the rhs in the Poisson equation).

Definition at line 25 of file charge.h.

### 7.2.2 Constructor & Destructor Documentation

## 7.2.2.1 Charge (const ParamList & params, const QuadratureRule & rule)

Constructor.

## **Parameters**

in	params	: the list of simulation parameters;
in	rule	: a quadrature rule.

Definition at line 5 of file charge.c++.

## 7.2.3 Member Function Documentation

**7.2.3.1 virtual VectorXr charge ( const VectorXr & phi )** [pure virtual]

Compute the total charge.

#### **Parameters**

in	phi	: the electric potential $oldsymbol{arphi}$ .

### Returns

the total charge q[C].

Implemented in GaussianCharge.

**7.2.3.2 virtual VectorXr dcharge ( const VectorXr & phi )** [pure virtual]

Compute the derivative of the total charge with respect to the electric potential.

#### **Parameters**

in	phi	: the electric potential $\phi$ .

#### Returns

the derivative:  $\frac{\mathrm{d}q}{\mathrm{d}\varphi}\left[C\cdot V^{-1}\right]$ .

Implemented in GaussianCharge.

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

- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/charge.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/charge.c++

### 7.3 CsvParser Class Reference

Class providing methods to read numeric content from a .csv file and to store it in Eigen matrices or vectors.

```
#include <csvParser.h>
```

### **Public Member Functions**

CsvParser ()=delete

Default constructor (deleted since it is required to specify at least a filename).

CsvParser (const std::string &, const bool &=true)

Constructor: load the input file and check its compatibility with the code.

virtual ∼CsvParser ()

Destructor: close the input file.

RowVectorXr importRow (const Index &)

Method to import a row from the input file.

MatrixXr importRows (const std::initializer\_list< Index > &)

Method to import multiple rows from the input file.

MatrixXr importFirstRows (const Index &)

Method to import the first nRows rows from the input file.

• VectorXr importCol (const Index &)

Method to import a column from the input file.

MatrixXr importCols (const std::initializer\_list< Index > &)

Method to import multiple columns from the input file.

MatrixXr importFirstCols (const Index &)

Method to import the first nCols columns from the input file.

• Real importCell (const Index &, const Index &)

Method to import a single cell from the input file.

· MatrixXr importAll ()

Method to import the whole input file.

### **Getter methods**

- const Index & nRows () const
- const Index & nCols () const

### **Private Member Functions**

• void reset ()

Reset all the flags for input\_ and go back to the beginning of file (possibly by ignoring headers).

## **Private Attributes**

bool hasHeaders

bool to determine if first row contains headers or not.

Index nRows

Number of rows in the input file.

Index nCols\_

Number of columns in the input file.

std::ifstream input\_

Input stream to input\_filename.

• std::string line\_

Auxiliary variable to store currently processed line.

· char separator\_

The separator character detected.

## 7.3.1 Detailed Description

Class providing methods to read **numeric** content from a .csv file and to store it in Eigen matrices or vectors.

Definition at line 29 of file csvParser.h.

## 7.3.2 Constructor & Destructor Documentation

7.3.2.1 CsvParser ( const std::string & input\_filename, const bool & hasHeaders = true )

Constructor: load the input file and check its compatibility with the code.

## Parameters

in	input_filename	: the name of the input file;
in	hasHeaders	: bool to specify if first row contains headers or not; if <b>true</b> , first row is always
		ignored.

Definition at line 10 of file csvParser.c++.

- 7.3.3 Member Function Documentation
- 7.3.3.1 RowVectorXr importRow ( const Index & index )

Method to import a row from the input file.

#### **Parameters**

in	index	: the row Index.
----	-------	------------------

### Returns

a row vector containing the content read.

Definition at line 65 of file csvParser.c++.

7.3.3.2 MatrixXr importRows ( const std::initializer\_list< Index > & indexes )

Method to import multiple rows from the input file.

## **Parameters**

in	indexes	: initializer list containing the row Indexes (e.g. something like {1, 3, 4}).
----	---------	--

#### Returns

a matrix containing the content read (row by row).

Definition at line 94 of file csvParser.c++.

7.3.3.3 MatrixXr importFirstRows ( const Index & nRows )

Method to import the first *nRows* rows from the input file.

#### **Parameters**

in	nRows	: the number of rows to import.
----	-------	---------------------------------

## Returns

a matrix containing the content read (row by row).

Definition at line 110 of file csvParser.c++.

7.3.3.4 VectorXr importCol ( const Index & index )

Method to import a column from the input file.

#### **Parameters**

in	index	: the column Index.
----	-------	---------------------

## Returns

a column vector containing the content read.

Definition at line 123 of file csvParser.c++.

7.3.3.5 MatrixXr importCols ( const std::initializer\_list< Index > & indexes )

Method to import multiple columns from the input file.

#### **Parameters**

in	indexes	: initializer list containing the column Indexes (e.g. something like {1, 3, 4}).

#### Returns

a matrix containing the content read (column by column).

Definition at line 148 of file csvParser.c++.

#### 7.3.3.6 MatrixXr importFirstCols ( const Index & nCols )

Method to import the first *nCols* columns from the input file.

#### **Parameters**

in	nCols	: the number of columns to import.
----	-------	------------------------------------

#### Returns

a matrix containing the content read (column by column).

Definition at line 164 of file csvParser.c++.

## 7.3.3.7 Real importCell ( const Index & rowIndex, const Index & colIndex )

Method to import a single cell from the input file.

#### **Parameters**

in	rowIndex	: the cell row Index.
in	collndex	: the cell column Index.

### Returns

a scalar containing the value read.

Definition at line 177 of file csvParser.c++.

## 7.3.3.8 MatrixXr importAll ( )

Method to import the whole input file.

#### Returns

a matrix containing the content read (cell by cell).

Definition at line 185 of file csvParser.c++.

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

- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/csvParser.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/csvParser.c++

## 7.4 DosModel Class Reference

Class providing methods to process a simulation to extract the Density of States starting from a parameter list.

#include <dosModel.h>

### **Public Member Functions**

· DosModel ()

Default constructor.

DosModel (const ParamList &)

Explicit conversion constructor.

virtual ~DosModel ()=default

Destructor (defaulted).

· const ParamList & params () const

Getter method.

void simulate (const GetPot &, const std::string &, const std::string &, const std::string &)
const

Perform the simulation.

 void post\_process (const GetPot &, const std::string &, std::ostream &, std::ostream &, const VectorXr &, const VectorXr &, const VectorXr &, const VectorXr &)

Perform post-processing.

• void gnuplot commands (const std::string &, std::ostream &) const

Defines commands to generate Gnuplot output files.

• void save\_plot (const std::string &, const std::string &, const std::string &, const std::string &) const Save the Gnuplot output files.

### **Private Attributes**

bool initialized

bool to determine if DosModel param\_ has been properly initialized.

· ParamList params\_

The parameter list.

## 7.4.1 Detailed Description

Class providing methods to process a simulation to extract the Density of States starting from a parameter list. Definition at line 35 of file dosModel.h.

#### 7.4.2 Constructor & Destructor Documentation

7.4.2.1 DosModel (const ParamList & params) [explicit]

Explicit conversion constructor.

## **Parameters**

in	params	: a parameter list.
----	--------	---------------------

Definition at line 10 of file dosModel.c++.

## 7.4.3 Member Function Documentation

7.4.3.1 void simulate ( const GetPot & config, const std::string & input\_experim, const std::string & output\_plot\_subdir, const std::string & output\_filename ) const

Perform the simulation.

#### **Parameters**

in	config	: the GetPot configuration object;
in	input_experim	: the file containing experimental data;
in	output_directory	: directory where to store output files;
in	output_plot	: sub-directory where to store Gnuplot files;
	subdir	
in	output_filename	: prefix for the output filename.

Definition at line 13 of file dosModel.c++.

7.4.3.2 void post\_process ( const GetPot & config, const std::string & input\_experim, std::ostream & output\_fitting, std::ostream & output\_CV, const VectorXr & x\_semic, const VectorXr & dens, const VectorXr & V\_simulated, const VectorXr & C\_simulated ) const

Perform post-processing.

### **Parameters**

in	config	: the GetPot configuration object;
in	input_experim	: the file containing experimental data;
out	output_fitting	: output file containing infos about fitting experimental data;
out	output_CV	: output file containing infos about capacitance-voltage data;
in	x_semic	: the mesh corresponding to the semiconductor domain;
in	dens	: charge density;
in	V_simulated	: simulated voltage values;
in	C_simulated	: simulated capacitance values.

Definition at line 176 of file dosModel.c++.

7.4.3.3 void gnuplot\_commands ( const std::string & output\_CV\_filename, std::ostream & os ) const

Defines commands to generate **Gnuplot** output files.

#### **Parameters**

in	output_CV filename	: output CV filename;
out	os	: output stream.

Definition at line 261 of file dosModel.c++.

7.4.3.4 void save\_plot ( const std::string & output\_directory, const std::string & output\_plot\_subdir, const std::string & output\_CV\_filename, const std::string & output\_filename ) const

Save the **Gnuplot** output files.

### **Parameters**

in	output_directory	: directory where to store output files;
in	output_plot	: sub-directory where to store Gnuplot files;
	subdir	
in	output_CV	: output CV filename;
	filename	

in output\_filename : prefix for the output filename.

Definition at line 290 of file dosModel.c++.

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

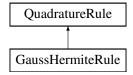
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/dosModel.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/dosModel.c++

## 7.5 GaussHermiteRule Class Reference

Class derived from QuadratureRule providing the Gauss-Hermite quadrature rule.

#include <quadratureRule.h>

Inheritance diagram for GaussHermiteRule:



#### **Public Member Functions**

• GaussHermiteRule ()=delete

Default constructor (deleted since it is required to specify the number of nodes).

GaussHermiteRule (const Index &)

Constructor.

virtual ∼GaussHermiteRule ()=default

Destructor (defaulted).

• virtual void apply () override

Apply the quadrature rule in order to compute the nodes and weights.

void apply (const GetPot &)

Apply the quadrature rule reading parameters from a configuration file.

void apply\_iterative\_algorithm (const Index &=1000, const Real &=1.0e-14)

Compute nodes and weights using an adapted version of the algorithm presented in: William H. Press, Saul A. Teukolsky, William T. Vetterling, and Brian P. Flannery. 2007. Numerical Recipes: The Art of Scientific Computing (3rd edition).

Cambridge University Press, New York, NY, USA.

• void apply\_using\_eigendecomposition ()

Compute nodes and weights using an eigendecomposition-based algorithm.

## **Additional Inherited Members**

## 7.5.1 Detailed Description

Class derived from QuadratureRule providing the Gauss-Hermite quadrature rule.

Compute nodes and weights for the nNodes -points approximation of

$$\int_{-\infty}^{+\infty} w(x) f(x) \, \mathrm{d}x$$

where  $w(x) = e^{-x^2}$ .

Definition at line 82 of file quadratureRule.h.

## 7.5.2 Constructor & Destructor Documentation

## 7.5.2.1 GaussHermiteRule (const Index & nNodes)

Constructor.

#### **Parameters**

in	nNodes	: the number of nodes to be used for the quadrature rule.
----	--------	---

Definition at line 14 of file quadratureRule.c++.

## 7.5.3 Member Function Documentation

## 7.5.3.1 void apply ( const GetPot & config )

Apply the quadrature rule reading parameters from a configuration file.

#### **Parameters**

in	config	: the GetPot configuration object.
----	--------	------------------------------------

Definition at line 22 of file quadratureRule.c++.

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

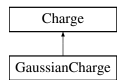
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/quadratureRule.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/quadratureRule.c++

## 7.6 GaussianCharge Class Reference

Class derived from Charge, under the hypothesis that Density of States is a combination of gaussians.

#include <charge.h>

Inheritance diagram for GaussianCharge:



#### **Public Member Functions**

• GaussianCharge ()=delete

Default constructor (deleted since it is required to specify a ParamList and a QuadratureRule).

• GaussianCharge (const ParamList &, const QuadratureRule &)

Constructor.

 $\bullet \ \ \text{virtual} \sim \\ \text{GaussianCharge ()=} \\ \text{default}$ 

Destructor (defaulted).

• virtual VectorXr charge (const VectorXr &) override

Compute the total charge.

• virtual VectorXr dcharge (const VectorXr &) override

Compute the derivative of the total charge with respect to the electric potential.

#### **Private Member Functions**

• Real n\_approx (const Real &, const Real &, const Real &) const

Compute electrons density (per unit volume).

Real dn approx (const Real &, const Real &, const Real &) const

Compute the approximate derivative of electrons density (per unit volume) with respect to the electric potential.

### **Additional Inherited Members**

### 7.6.1 Detailed Description

Class derived from Charge, under the hypothesis that Density of States is a combination of gaussians.

Provide methods to compute total electric charge and its derivative under the hypothesis that Density of States is a linear combination of multiple gaussians, whose parameters are read from a ParamList object.

Definition at line 70 of file charge.h.

## 7.6.2 Constructor & Destructor Documentation

## 7.6.2.1 GaussianCharge (const ParamList & params, const QuadratureRule & rule)

#### Constructor.

### **Parameters**

in	params	: the list of simulation parameters;
in	rule	: a quadrature rule.

Definition at line 8 of file charge.c++.

## 7.6.3 Member Function Documentation

7.6.3.1 VectorXr charge (const VectorXr & phi) [override], [virtual]

Compute the total charge.

### **Parameters**

in $phi$ : the electric potential $\varphi$ .	ı ın
---	------

## Returns

the total charge q[C].

Implements Charge.

Definition at line 37 of file charge.c++.

## 7.6.3.2 VectorXr dcharge (const VectorXr & phi) [override], [virtual]

Compute the derivative of the total charge with respect to the electric potential.

### **Parameters**

ı			
	in	phi	: the electric potential $\phi$ .

### Returns

the derivative:  $\frac{\mathrm{d}q}{\mathrm{d}\varphi}\left[C\cdot V^{-1}\right]$ .

Implements Charge.

Definition at line 60 of file charge.c++.

7.6.3.3 Real n\_approx ( const Real & phi, const Real & NO, const Real & sigma ) const [private]

Compute electrons density (per unit volume).

### **Parameters**

in	phi	: the electric potential $\phi$ ;
in	N0	: the gaussian mean $N_0$ ;
in	sigma	: the gaussian standard deviation $\sigma$ .

### Returns

the electrons density  $n(\varphi) \left[ m^{-3} \right]$ .

Definition at line 11 of file charge.c++.

7.6.3.4 Real dn\_approx ( const Real & phi, const Real & NO, const Real & sigma ) const [private]

Compute the approximate derivative of electrons density (per unit volume) with respect to the electric potential.

# **Parameters**

in	phi	: the electric potential $oldsymbol{arphi}$ ;
in	N0	: the gaussian mean $N_0$ ;
in	sigma	: the gaussian standard deviation $\sigma$ .

## Returns

the derivative:  $\frac{\mathrm{d}n}{\mathrm{d}\varphi}\left[m^{-3}\cdot V^{-1}\right]$ .

Definition at line 24 of file charge.c++.

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

- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/charge.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/charge.c++

# 7.7 NonLinearPoisson1D Class Reference

Provide a solver for a non-linear Poisson equation.

#include <solvers.h>

### **Public Member Functions**

• NonLinearPoisson1D ()=delete

Default constructor (deleted since it is required to specify the solver to be used).

NonLinearPoisson1D (const PdeSolver1D &, const Index &=100, const Real &=1.0e-6)

Constructor.

virtual ~NonLinearPoisson1D ()=default

Destructor (defaulted).

• void apply (const VectorXr &, const VectorXr &, Charge &)

Apply a Newton method to the equation and then discretize it using the solver specified.

### **Getter methods**

- const VectorXr & phi () const
- const VectorXr & norm () const
- · const Real & qTot () const
- · const Real & cTot () const

### **Private Member Functions**

SparseXr computeJac (const VectorXr &) const

Compute the Jacobi matrix.

### **Private Attributes**

const PdeSolver1D & solver\_

Solver handler.

Index maxIterationsNo

Maximum number of iterations.

· Real tolerance\_

Tolerance.

VectorXr phi\_

The electric potential.

VectorXr norm\_

Vector holding  $L^{\infty}$ -norm errors for each iteration.

Real qTot\_

Total charge.

• Real cTot\_

Total capacitance.

# 7.7.1 Detailed Description

Provide a solver for a non-linear Poisson equation.

A Newton method is applied in order to solve:

$$-\frac{\mathrm{d}}{\mathrm{d}z}\left(\varepsilon(z)\cdot\frac{\mathrm{d}\varphi}{\mathrm{d}z}(z)\right) = -q\cdot\frac{N_0}{\sqrt{\pi}}\int_{-\infty}^{+\infty}\exp\left(-\alpha^2\right)\left(1+\exp\left(\frac{\sqrt{2}\sigma\alpha-q\varphi(z)}{K_B\cdot T}\right)\right)^{-1}\mathrm{d}\alpha\;.$$

Definition at line 190 of file solvers.h.

- 7.7.2 Constructor & Destructor Documentation
- 7.7.2.1 NonLinearPoisson1D ( const PdeSolver1D & solver, const Index & maxlterationsNo = 100, const Real & tolerance = 1.0e-6)

Constructor.

### **Parameters**

in	solver	: the solver to be used;
in	maxIterationsNo	: maximum number of iterations desired;
in	tolerance	: tolerance desired.

Definition at line 162 of file solvers.c++.

# 7.7.3 Member Function Documentation

7.7.3.1 void apply ( const VectorXr & mesh, const VectorXr & init\_guess, Charge & charge\_fun )

Apply a Newton method to the equation and then discretize it using the solver specified.

### **Parameters**

in	mesh	: the mesh;
in	init_guess	: initial guess for the Newton algorithm;
in	charge_fun	: an object of class Charge specifying how to compute total electric charge.

Definition at line 165 of file solvers.c++.

# 7.7.3.2 SparseXr computeJac ( const VectorXr & x ) const [private]

Compute the Jacobi matrix.

### **Parameters**

in	X	: the vector where to start from.

### Returns

the Jacobi matrix in a sparse format.

Definition at line 273 of file solvers.c++.

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

- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/solvers.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/solvers.c++

# 7.8 ParamList Class Reference

Class providing methods to handle a list of parameters.

#include <paramList.h>

### **Public Member Functions**

• ParamList ()=default

Default constructor (defaulted).

ParamList (const RowVectorXr &)

Explicit conversion constructor.

• virtual  $\sim$ ParamList ()=default

Destructor (defaulted).

# **Getter methods**

```
• const Index & simulationNo () const
       • const Real & t_semic () const
       • const Real & t_ins () const
       • const Real & eps_semic () const
       • const Real & eps_ins () const
       • const Real & Wf () const
       · const Real & Ea () const

    const Real & N0 () const

       • const Real & sigma () const
       • const Real & N0_2 () const
       • const Real & sigma_2 () const
       • const Real & shift_2 () const
       • const Real & N0_3 () const
        • const Real & sigma_3 () const
       · const Real & shift_3 () const
       · const Real & N0_4 () const
       • const Real & sigma_4 () const
       • const Real & shift_4 () const
       • const Index & nNodes () const
       • const Index & nSteps () const

    const Real & V_min () const

       • const Real & V_max () const
Private Attributes

    Index simulationNo_

          Index of the simulation.
    · Real t_semic_
          Semiconductor layer thickness [m].
    · Real t_ins_
          Insulator layer thickness [m].

    Real eps_semic

          Semiconductor layer relative electrical permittivity [].
    · Real eps_ins_
          Insulator layer relative electrical permittivity [].

    Real Wf

          Work-function [V].

    Real Ea_

          Electron affinity [V].

    Real N0

           1st gaussian mean \lceil m^{-3} \rceil.
    · Real sigma_
           1st gaussian standard deviation (normalized by K_B \cdot T) [].
    • Real N0_2_
          2nd gaussian mean.
    · Real sigma_2_
          2nd gaussian standard deviation.

    Real shift 2

          2nd gaussian shift with respect to the 1st gaussian electric potential.

    Real N0 3

          3rd gaussian mean.

    Real sigma_3_

          3rd gaussian standard deviation.
```

3rd gaussian shift with respect to the 1st gaussian electric potential.

Real shift 3

Real N0\_4\_

4th gaussian mean.

· Real sigma\_4\_

4th gaussian standard deviation.

Real shift 4

4th gaussian shift with respect to the 1st gaussian electric potential.

Index nNodes\_

Number of nodes that form the mesh.

Index nSteps\_

Number of steps to simulate.

Real V min

Minimum voltage [V].

Real V\_max\_

Maximum voltage [V].

### **Friends**

- · class GaussianCharge
- · class DosModel

# 7.8.1 Detailed Description

Class providing methods to handle a list of parameters.

It can include up to 4 gaussians, later combined to compute total charge.

Definition at line 25 of file paramList.h.

# 7.8.2 Constructor & Destructor Documentation

# 7.8.2.1 ParamList (const RowVectorXr & list) [explicit]

Explicit conversion constructor.

### **Parameters**

in	list	: a row vector containing a parameters list (for example got by a CsvParser
		object). Parameters should be sorted in the same order as specified above.

Definition at line 5 of file paramList.c++.

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

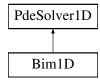
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/paramList.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/paramList.c++

# 7.9 PdeSolver1D Class Reference

Abstract class providing methods to assemble matrices to solve one-dimensional PDEs.

#include <solvers.h>

Inheritance diagram for PdeSolver1D:



### **Public Member Functions**

• PdeSolver1D ()=delete

Default constructor (deleted since it is required to specify the mesh).

PdeSolver1D (VectorXr &)

Constructor.

virtual ∼PdeSolver1D ()=default

Destructor (defaulted).

virtual void assembleAdvDiff (const VectorXr &alpha, const VectorXr &gamma, const VectorXr &eta, const VectorXr &beta)=0

Assemble the matrix for an advection-diffusion term.

virtual void assembleStiff (const VectorXr &eps, const VectorXr &kappa)=0

Assemble the matrix for a diffusion term.

• virtual void assembleMass (const VectorXr &delta, const VectorXr &zeta)=0

Assemble the matrix for a reaction term.

### **Getter methods**

- const SparseXr & AdvDiff () const
- const SparseXr & Stiff () const
- const SparseXr & Mass () const

# **Protected Attributes**

VectorXr mesh

The mesh.

Index nNodes

Number of nodes that form the mesh.

SparseXr AdvDiff\_

Matrix for an advection-diffusion term.

SparseXr Stiff

Stiffness matrix.

SparseXr Mass\_

Mass matrix.

# **Friends**

• class NonLinearPoisson1D

# 7.9.1 Detailed Description

Abstract class providing methods to assemble matrices to solve one-dimensional PDEs.

Matrices are held in a sparse format.

Definition at line 31 of file solvers.h.

### 7.9.2 Constructor & Destructor Documentation

# 7.9.2.1 PdeSolver1D ( VectorXr & mesh )

Constructor.

### **Parameters**

-			
	in	mesh	: the mesh.

Definition at line 3 of file solvers.c++.

### 7.9.3 Member Function Documentation

7.9.3.1 virtual void assembleAdvDiff ( const VectorXr & alpha, const VectorXr & gamma, const VectorXr & eta, const VectorXr & beta ) [pure virtual]

Assemble the matrix for an advection-diffusion term.

Build the matrix for the advection-diffusion problem:  $-\nabla \cdot (\alpha \cdot \gamma(\eta \nabla u - \beta u)) = f$ .

### **Parameters**

in	alpha	: $lpha$ , an element-wise constant function;
in	gamma	: $\gamma$ , an element-wise linear function;
in	eta	: $\eta$ , an element-wise linear function;
in	beta	: $eta$ , an element-wise constant function.

Implemented in Bim1D.

7.9.3.2 virtual void assembleStiff ( const VectorXr & eps, const VectorXr & kappa ) [pure virtual]

Assemble the matrix for a diffusion term.

Build the matrix for the diffusion problem:  $-\nabla \cdot (\varepsilon \cdot \kappa \nabla u) = f$ .

### **Parameters**

in	eps	$: \mathcal{E},$ an element-wise constant function;
in	kappa	: $\kappa$ , an element-wise linear function.

Implemented in Bim1D.

7.9.3.3 virtual void assembleMass (const VectorXr & delta, const VectorXr & zeta) [pure virtual]

Assemble the matrix for a reaction term.

Build the mass matrix for the reaction problem:  $\delta \cdot \zeta u = f$ .

### **Parameters**

in	delta	: $\delta$ , an element-wise constant function;
in	zeta	: $\zeta$ , an element-wise linear function.

Implemented in Bim1D.

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

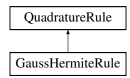
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/solvers.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/solvers.c++

#### 7.10 **QuadratureRule Class Reference**

Abstract class providing a quadrature rule.

#include <quadratureRule.h>

Inheritance diagram for QuadratureRule:



### **Public Member Functions**

• QuadratureRule ()=delete

Default constructor (deleted since it is required to specify the number of nodes).

• QuadratureRule (const Index &)

Constructor.

virtual ∼QuadratureRule ()=default

Destructor (defaulted).

virtual void apply ()=0

Apply the quadrature rule in order to compute the nodes and weights.

### **Getter methods**

- const Index & nNodes () const
- const VectorXr & nodes () const
- const VectorXr & weights () const

# **Protected Attributes**

Index nNodes

Number of nodes of the quadrature rule.

VectorXr nodes\_

Vector containing the computed nodes coordinates.

VectorXr weights\_

Vector containing the computed weights.

# **Friends**

• class GaussianCharge

#### 7.10.1 **Detailed Description**

Abstract class providing a quadrature rule.

Approximate the integral:

$$\int_{a}^{b} f(x) \, \mathrm{d}x$$

with the finite sum:

$$\int_{a}^{b} f(x) dx$$

$$\sum_{i=1}^{nNodes} w_{i} \cdot f(x_{i})$$

where  $\{x_i\}_{i=1}^{nNodes}$  and  $\{w_i\}_{i=1}^{nNodes}$  are called respectively nodes and weights.

Definition at line 29 of file quadratureRule.h.

# 7.10.2 Constructor & Destructor Documentation

# 7.10.2.1 QuadratureRule (const Index & nNodes)

Constructor.

### **Parameters**

in	nNodes	: the number of nodes to be used for the quadrature rule.

Definition at line 5 of file quadratureRule.c++.

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

- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/quadratureRule.h
- /home/Data/Dropbox/Progetto-PACS/C++/Source/src/quadratureRule.c++

# **Chapter 8**

# **File Documentation**

# 8.1 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/charge.h File Reference

Classes for computing total electric charge.

```
#include "paramList.h"
#include "quadratureRule.h"
#include "typedefs.h"
```

### **Classes**

· class Charge

Abstract class providing methods to calculate total electric charge (the rhs in the Poisson equation).

class GaussianCharge

Class derived from Charge, under the hypothesis that Density of States is a combination of gaussians.

# 8.1.1 Detailed Description

Classes for computing total electric charge.

**Author** 

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file charge.h.

# 8.2 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/csvParser.h File Reference

Tools to store content from a .csv file in matrices or vectors.

```
#include "typedefs.h"
#include <string>
#include <fstream>
#include <sstream>
#include <utility>
```

### Classes

class CsvParser

Class providing methods to read numeric content from a .csv file and to store it in Eigen matrices or vectors.

### 8.2.1 Detailed Description

Tools to store content from a .csv file in matrices or vectors.

**Author** 

```
Pasquale Claudio Africa pasquale.africa@gmail.com
```

Date

2014

Definition in file csvParser.h.

# 8.3 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/dosModel.h File Reference

Mathematical model for Density of States extraction.

```
#include "charge.h"
#include "csvParser.h"
#include "numerics.h"
#include "paramList.h"
#include "quadratureRule.h"
#include "solvers.h"
#include "typedefs.h"
#include "gnuplot-iostream.h"
#include <chrono>
#include <limits>
```

# Classes

· class DosModel

Class providing methods to process a simulation to extract the Density of States starting from a parameter list.

# 8.3.1 Detailed Description

Mathematical model for Density of States extraction.

**Author** 

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file dosModel.h.

# 8.4 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/numerics.h File Reference

### Generic numeric algorithms.

```
#include "typedefs.h"
#include <limits>
```

### **Namespaces**

numerics

Namespace for generic numeric algorithms.

### **Functions**

 $\bullet \ \ \text{template}{<} \text{typename ScalarType} >$ 

```
VectorX< ScalarType > sort (const VectorX< ScalarType > &vector)
```

Function to sort Eigen vectors.

template<typename ScalarType >

```
VectorXpair< ScalarType > sort_pair (const VectorX< ScalarType > &vector)
```

Function to sort Eigen vectors, keeping track of indexes.

Real trapz (const VectorXr &x, const VectorXr &y)

Function to compute approximate integral of y with spacing increment specified by x, using trapezoidal rule.

Real trapz (const VectorXr &y)

Compute the approximate integral of y with unit spacing, using trapezoidal rule.

VectorXr deriv (const VectorXr &, const VectorXr &)

Compute the numeric derivative:  $\frac{dy}{dx}$ .

• Real interp1 (const VectorXr &, const VectorXr &, const Real &)

Linear 1D interpolation. Interpolate y, defined at points x, at the point xNew.

VectorXr interp1 (const VectorXr &, const VectorXr &, const VectorXr &)

Linear 1D interpolation. Interpolate y, defined at points x, at the points xNew.

• Real error\_L2 (const VectorXr &, const VectorXr &, const VectorXr &, const Real &)

Compute the  $L^2$ -norm error between simulated and interpolated values, using trapz.

# 8.4.1 Detailed Description

Generic numeric algorithms.

Author

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file numerics.h.

# 8.5 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/paramList.h File Reference

List of simulation parameters.

```
#include "typedefs.h"
```

# Classes

class ParamList

Class providing methods to handle a list of parameters.

# 8.5.1 Detailed Description

List of simulation parameters.

**Author** 

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file paramList.h.

# 8.6 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/physicalConstants.h File Reference

```
Physical constants.
```

```
#include "typedefs.h"
```

# **Namespaces**

· constants

Numerical constants.

# **Variables**

```
• const Real Q = 1.60217653000000e-19 
 Electron charge [C].
```

• const Real Q2 = Q \* Q

Electron charge squared  $[C^2]$ .

• const Real K\_B = 1.38065050000000e-23

Boltzmann's constant  $[J \cdot K^{-1}]$ .

• const Real EPS0 = 8.854187817e-12

*Vacuum electrical permittivity*  $[C \cdot V^{-1} \cdot m^{-1}]$ .

• const Real T = 300

Reference temperature [K].

• const Real V\_TH = K\_B \* T / Q

Treshold voltage [V].

# 8.6.1 Detailed Description

Physical constants.

**Author** 

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file physicalConstants.h.

# 8.7 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/quadratureRule.h File Reference

```
Quadrature rules.
```

```
#include "typedefs.h"
```

### Classes

• class QuadratureRule

Abstract class providing a quadrature rule.

• class GaussHermiteRule

Class derived from QuadratureRule providing the Gauss-Hermite quadrature rule.

# 8.7.1 Detailed Description

Quadrature rules.

**Author** 

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file quadratureRule.h.

# 8.8 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/solvers.h File Reference

### Generic solvers for PDEs.

```
#include "charge.h"
#include "typedefs.h"
#include <utility>
#include <limits>
```

### Classes

class PdeSolver1D

Abstract class providing methods to assemble matrices to solve one-dimensional PDEs.

class Bim1D

Class derived from PdeSolver1D, providing a finite volume Box Integration Method (BIM) solver.

· class NonLinearPoisson1D

Provide a solver for a non-linear Poisson equation.

# 8.8.1 Detailed Description

Generic solvers for PDEs.

**Author** 

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file solvers.h.

# 8.9 /home/Data/Dropbox/Progetto-PACS/C++/Source/src/typedefs.h File Reference

# Typedefs and utility functions.

```
#include <Eigen/Dense>
#include <Eigen/Sparse>
#include "GetPot"
#include <iostream>
#include <fstream>
#include "physicalConstants.h"
```

# **Namespaces**

· constants

Numerical constants.

utility

Namespace for utilities and auxiliary functions.

# **Macros**

• #define Real double

Pre-processor macro for real numbers.

• #define Index ptrdiff\_t

Pre-processor macro for indexing variables.

# **Typedefs**

```
    typedef Matrix Real, Dynamic,
Dynamic > MatrixXr
```

Typedef for dense real-valued dynamic-sized matrices.

typedef Matrix< Real, Dynamic, 1 > VectorXr

Typedef for dense real-valued dynamic-sized column vectors.

typedef Matrix < Real, 1, Dynamic > RowVectorXr

Typedef for dense real-valued dynamic-sized row vectors.

typedef SparseMatrix< Real > SparseXr

Typedef for sparse real-valued dynamic-sized matrices.

template<typename ScalarType >
 using VectorX = Matrix< ScalarType, Dynamic, 1 >

Template alias for Eigen vectors.

template < typename T >
 using VectorXpair = VectorX < std::pair < T, Index > >

Template alias for an Eigen vector of pairs: (ScalarType, Index).

### **Functions**

• std::string full\_path (const std::string &, const std::string &)

Auxiliary function to return the full path to a file.

void print block (const char \*, std::ostream &=std::cout)

Auxiliary function to print a string inside a block.

void print\_done (std::ostream &=std::cout)

Auxiliary function to print a "DONE!" string.

### **Variables**

• const unsigned PARAMS\_NO = 22

Number of parameters required in input file.

• const Real PI = M\_PI

 $\pi$ 

const Real SQRT\_PI = std::sqrt(PI)

 $\sqrt{\pi}$ .

const Real PI\_M4 = 0.7511255444649425

 $\pi^{-\frac{1}{4}}$ 

• const Real SQRT\_2 = std::sqrt(2)

 $\sqrt{2}$ .

# 8.9.1 Detailed Description

Typedefs and utility functions.

**Author** 

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file typedefs.h.

# 8.9.2 Typedef Documentation

8.9.2.1 using VectorX = Matrix < ScalarType, Dynamic, 1>

Template alias for Eigen vectors.

**Template Parameters** 

```
ScalarType : the scalar type.
```

Definition at line 41 of file typedefs.h.

```
8.9.2.2 using VectorXpair = VectorX<std::pair<T, Index>>
```

Template alias for an Eigen vector of pairs: (ScalarType, Index).

**Template Parameters** 

```
ScalarType : the scalar type.
```

Definition at line 48 of file typedefs.h.

# 8.10 /home/Data/Dropbox/Progetto-PACS/C++/Source/test/simulate\_dos.c++ File Reference

A test file.

```
#include "src/dosModel.h"
```

# **Functions**

• int main (const int argc, const char \*const \*argv, const char \*const \*envp)

The main function.

# 8.10.1 Detailed Description

A test file.

Author

Pasquale Claudio Africa pasquale.africa@gmail.com

Date

2014

Definition in file simulate\_dos.c++.

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