libBorealis Documentation

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Chapter 1

Class Index

1.1 Class Hierarchy

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Chapter 2

Class Index

2.1 Class List

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

ырпана	rmonicOscillator	
	Harmonic Oscillator single particle wave function class. Uses the HarmonicOscillator BasisFunction subclasses auto-generated by - SymPy through the orbitalsGenerator tool	7
ASGD		
	Implementation for the Adaptive Stochastic Gradient Descent method (ASGD) Used to find optimal variational parameters using adaptive step lengths	10
AtomCor	re	
	Implementation of the Atom Core potentialZ/r	12
BasisFur	nctions	
	The Superclass shell for orbital basis functions	13
Blocking		14
Bosons		
	The Boson system class	16
Brute_Fo	orce	
	Implementation of the Brute Force QMC. Uses the Simle diffusion class. All methods are empty except for the energy necessities part which requires the gradients to be calculated (not using the Quantum Force)	18
Coulomb		
	Implementation of the Coulomb potential. 1/r_{ij}	20
Diffusion		
	Class containing rules for walker movement based on diffusion models. Serves as class member in the Sampling class. Brute force implies the Simple diffusion model, while Importance Sampling implies the Fokker Planck diffusion	21

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Distribution	
Class for calculating distribution functions such as the one-body der sity. Does not collect data itself, but works merely as a control orga for the QMC class, calling it's methods for storing position data .	
DMC	
Implementation of the Diffusion Monte-Carlo Method. Very little needs to be added when the QMC superclass holds all the generation functionality	
DMCparams	
Struct used to initialize DMC parameters	29
Error Estimator	20
Class handling error estimations of the QMC methods. The QMC class holds an object of this type, calling the update_data function i order to update the sampling pool. finalize() then either dumps th samples to file for later processing, or calulates an estimate	n e
ExpandedBasis	32
Fermions	
The Fermion system class	34
Fokker_Planck	
Anisotropic diffusion by the Fokker-Planck equation	37
GeneralParams	
Struct used to initialize general parameters	39
Harmonic osc	
Implementation of the Harmonic Oscillator potential. 0.5*w**2*r**	2 40
HartreeFock	41
hydrogenicOrbitals	
Hyrogen-like single particle wave function class. Uses the hydrogenic BasisFunction subclasses auto-generated by SymPy throug the orbitalsGenerator tool	
Importance	
Implementation of Importance sampled QMC. Using the Fokker-Planck diffusion class. Introduces the Quantum Force	44
Jastrow	
The class representing the Jastrow correlation functions Holds a data concerning the Jastrow function and it's influence on the QMO algorithm	
Minimizer	
Class for minimization methods used to obtain optimal variations	al 50
MinimizerParams	
Struct used to initialize Minimization parameters	52
No_Jastrow	_
Class loaded when no correlation factor is used	53
Class for suppressing standard output. Every node but the master has this. If-tests around cout is avoided	er 56
Orbitals Superclass for the single particle orbital classes. Handles everythin specific regarding choice of single particle basis	-

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OutputHa	andler	
	Class for handling output-methods. Designed to avoid rewriting code, as well as avoid if-tests if output is not desired	62
OutputPa	arams	
	Struct used to initialize output parameters	65
Pade_Ja	strow	
	The Pade Jastrow factor with a single variational parameter	65
ParParan	- · · · · · · · · · · · · · · · · · · ·	
	Struct used to initialize parallelization parameters	68
Potential	·	
QMC	Superclass for potentials. Potentials are stores in a vector in the system object	69
QIVIO	The QMC superclass. Holds implementations of general functions	
	for both VMC and DMC in order to avoid rewriting code and empha-	
	size the similarities	70
•		78
Sampling]	79
Simple		
	Simple Isotropic diffusion model	83
SimpleVa	ar	
	Calculates the simple variance of the sampled values	84
STDOUT	•	
	Class for handling standard output. Only the master node has this object	85
stdoutAS		
	Class for handling the output of ASGD. Ouputs values such as the variational gradients, step length, variational parameters, etc	86
stdoutDN	MC	
	Class for handling the output of DMC. Outputs values such as the trial energy, dmc energy, number of walkers, etc	87
System		
	The system class separating Fermions and Bosons. Designed to	
	generalize the solver in terms of particle species	88
SystemC	· · · ·	
-,	Struct used to initialize system objects	91
Variation	alParams	
variation	Struct used to initialize the varational parameters	92
VMC	Citati asca to initialize the varational parameters	52
VIVIC	Implementation of the Variational Manta Carlo Mathad Vary little	
	Implementation of the Variational Monte-Carlo Method. Very little needs to be added when the QMC superclass holds all the general	
	functionality	92
VMCpara		
	Struct used to initialize VMC parameters	94
Walker		
	Class representing a Random Walker. Holds position data, alive state, etc. Designed to lighten function arguments, and ease implementation of QMC methods involving multiple walkers. Alot of values are stored to avoid calculating the same value twice	95

6 Class Index

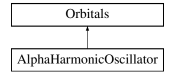
Chapter 3

Class Documentation

3.1 AlphaHarmonicOscillator Class Reference

Harmonic Oscillator single particle wave function class. Uses the HarmonicOscillator - BasisFunction subclasses auto-generated by SymPy through the orbitalsGenerator tool.

Inheritance diagram for AlphaHarmonicOscillator:



Public Member Functions

- AlphaHarmonicOscillator (GeneralParams &, VariationalParams &)
- void set_qnum_indie_terms (const Walker *walker, int i)

Protected Member Functions

- double get_variational_derivative (const Walker *walker, int n)
- void get_qnums ()
- double get coulomb element (const arma::uvec &qnum set)
- double **get_sp_energy** (int qnum) const
- double H (int n, double x) const

Method for calculating Hermite polynomials.

- double get_parameter (int n)
- void set_parameter (double parameter, int n)

Protected Attributes

· double w

The oscillator frequency.

• double * alpha

Pointer to the variational parameter alpha. Shared address with all the BasisFunction subclasses.

• double * k

Pointer to sqrt(alpha*w). Shared address with all the BasisFunction subclasses.

double * k2

Pointer to alpha*w. Shared address with all the BasisFunction subclasses.

- double * w over a
- double * exp_factor

Pointer to a factor precalculated by set_qnum_indie_terms(). Shared address with all the BasisFunction subclasses.

Friends

· class ExpandedBasis

3.1.1 Detailed Description

Harmonic Oscillator single particle wave function class. Uses the HarmonicOscillator - BasisFunction subclasses auto-generated by SymPy through the orbitalsGenerator tool.

3.1.2 Member Function Documentation

```
3.1.2.1 double AlphaHarmonicOscillator::get_coulomb_element( const arma::uvec & qnum_set ) [protected, virtual]
```

For Quantum Dots, closed form expressions for the matrix elements exist.

Reimplemented from Orbitals.

Returns

The variational parameter alpha.

Implements Orbitals.

```
3.1.2.3 void AlphaHarmonicOscillator::get_qnums() [protected]
```

Calculates the quantum numbers of the oscillator and stores them in the matrix qnums.

```
3.1.2.4 double AlphaHarmonicOscillator::get_variational_derivative ( const Walker * walker, int n ) [protected, virtual]
```

Overridden superclass method implementing closed form expressions using Hermite polynomials.

Reimplemented from Orbitals.

3.1.2.5 double AlphaHarmonicOscillator::H(int n, double x) const [protected]

Method for calculating Hermite polynomials.

Parameters

n	The degree of the Hermite polynomial.
X	The argument for evaluating the polynomial.

Sets a new value for the alpha and updates all the pointer values.

Implements Orbitals.

Calculates the exponential term shared by all oscillator function once pr. particle to save CPU-time.

See also

```
Orbitals::set_qnum_indie_terms()
```

Reimplemented from Orbitals.

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

- src/Orbitals/AlphaHarmonicOscillator/AlphaHarmonicOscillator.h
- src/Orbitals/AlphaHarmonicOscillator/AlphaHarmonicOscillator.cpp

3.2 ASGD Class Reference

Implementation for the Adaptive Stochastic Gradient Descent method (ASGD) Used to find optimal variational parameters using adaptive step lengths.

Inheritance diagram for ASGD:



Public Member Functions

- ASGD (VMC *, MinimizerParams &, const ParParams &)
- · void minimize ()

Method for executing the minimization main solver.

Protected Member Functions

void get_total_grad ()

Method for calculating the total gradient.

• void update parameters ()

Calculates the step and updates parameters.

void output_cycle ()

Standard output of the progress.

• void thermalize_walkers ()

Thermalizes a set of walkers before the main loop.

• double f (double x)

Function for calculating the adaptive step.

• void get_variational_gradients (Walker *walker, double e_local)

Method for updating the vectors needed to calculate the total variational derivative.

Protected Attributes

• int n_c

The correlation length between storing two walkers after thermalization.

• int n_c_SGD

The number of samples used to estimate expectation values.

• int SGDsamples

The number of ASGD cycles.

· int n walkers

The number of walkers.

· int thermalization

The number of thermalization cycles used on walkers.

• int sample

The current ASGD cycle.

double t_prev

The previous t.

double t

The current t.

double step

The current calculates step.

• double max step

The maximum threshold on a step.

• double E

The energy summation variable used to calculate the mean.

• double a

ASGD step parameter.

• double A

ASGD step parameter.

• double f_min

ASGD step parameter.

• double f_max

ASGD step parameter.

double w

ASGD step parameter.

Walker ** walkers

The walkers used to sample expectation values.

- Walker ** trial_walkers
- arma::rowvec parameter
- arma::rowvec gradient

Sumamtion vector for the trial wave function's variational derivatives.

• arma::rowvec gradient_local

Sumamtion vector for the trial wave function's variational derivatives times the energy.

· arma::rowvec gradient_old

The previous total gradient.

• arma::rowvec gradient_tot

The current total gradient.

Friends

· class stdoutASGD

3.2.1 Detailed Description

Implementation for the Adaptive Stochastic Gradient Descent method (ASGD) Used to find optimal variational parameters using adaptive step lengths.

3.2.2 Member Function Documentation

```
3.2.2.1 void ASGD::get_total_grad() [protected]
```

Method for calculating the total gradient.

Updates the error estimator with statistics.

Method for updating the vectors needed to calculate the total variational derivative.

Calculates the single particle variational derivatives V and accumulates V and V*e_local.

Parameters

```
e_local The local energy of the current walker at the current time step.
```

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

- src/Minimizer/ASGD/ASGD.h
- src/Minimizer/ASGD/ASGD.cpp

3.3 AtomCore Class Reference

Implementation of the Atom Core potential. -Z/r.

Inheritance diagram for AtomCore:



Public Member Functions

- AtomCore (GeneralParams &)
- double get_pot_E (const Walker *walker) const
 Method for calculating a walker's potential energy.

Protected Attributes

• int Z

The core charge.

3.3.1 Detailed Description

Implementation of the Atom Core potential. -Z/r.

3.3.2 Member Function Documentation

3.3.2.1 double AtomCore::get_pot_E(const Walker * walker) const [virtual]

Method for calculating a walker's potential energy.

Method overridden by subclasses.

Parameters

walker The walker for which the potential energy should be calculated.

Returns

The potential energy.

Implements Potential.

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

- src/Potential/AtomCore/AtomCore.h
- src/Potential/AtomCore/AtomCore.cpp

3.4 BasisFunctions Class Reference

The Superclass shell for orbital basis functions.

Public Member Functions

virtual double eval (const Walker *walker, int i)=0
 The method representing the orbitals functional expression.

3.4.1 Detailed Description

The Superclass shell for orbital basis functions.

Each single particle orbital has it's own implementation as a subclass of this class. A set of orbitals can then be loaded into the Orbitals basis_function vectors. An orbital-Generator script is supplied to autogenerate CPP files using this class.

See also

```
Orbitals::basis_functions
Orbitals::dell_basis_functions
Orbitals::lapl_basis_functions
```

3.4.2 Member Function Documentation

```
3.4.2.1 virtual double BasisFunctions::eval ( const Walker * walker, int i ) [pure virtual]
```

The method representing the orbitals functional expression.

Parameters

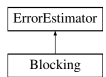
walker	The Walker whose position the orbital is to be evaluted at.
i	The particle to be evaluated (Single particle function).

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

- · src/BasisFunctions/BasisFunctions.h
- src/BasisFunctions/BasisFunctions.cpp

3.5 Blocking Class Reference

Inheritance diagram for Blocking:



Public Member Functions

- **Blocking** (int n_c, ParParams &pp, std::string filename="blocking_out", std::string path="./", int n_b=100, int maxb=10000, int minb=10, bool rerun=false)
- **Blocking** (int n_c, std::string filename="blocking_out", std::string path="./", int n-b=100, int maxb=10000, int minb=10)
- double estimate_error ()

Estimates the error based on the subclass implementation.

- void get_initial_error ()
- void get_unique_blocks (arma::Row< int > &block_sizes, int &n)

Calculates the block sizes.

Protected Member Functions

void block_data (int block_size, double &var, double &mean)
 Calculates the variance and mean of the dataset with the specified block size.

Protected Attributes

- arma::rowvec local block
- · int min block size

The minimum amount of samples in one block.

int max_block_size

The maximum amount of samples in one block.

int n_block_samples

The total amount of different block sizes.

3.5.1 Member Function Documentation

3.5.1.1 void Blocking::block_data (int block_size, double & var, double & mean) [protected]

Calculates the variance and mean of the dataset with the specified block size.

Parameters

block_size	The number of samples in each block.
var	Reference to the variance of the block's means.
mean	Reference to the mean of the block's means. Needed to combine the
	variances from different processes.

3.5.1.2 void Blocking::get_initial_error()

Calculates the variance as in SimpleVar

3.5.1.3 void Blocking::get_unique_blocks (arma::Row < int > & block_sizes, int & n)

Calculates the block sizes.

Due to integer division, alot of sizes becomes equal. Only unique block sizes are returned.

Parameters

block_sizes	Vector containing the block sizes
n	The number of unquue block sizes

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

- · src/ErrorEstimator/Blocking/Blocking.h
- · src/ErrorEstimator/Blocking/Blocking.cpp

3.6 Bosons Class Reference

The Boson system class.

Inheritance diagram for Bosons:



Public Member Functions

- Bosons (GeneralParams &, Orbitals *)
- void get_spatial_grad (Walker *walker, int particle) const

Method for calculating the changed part of the spatial gradient.

void get_spatial_grad_full (Walker *walker) const

Method for calculating the full spatial gradient.

double get_spatial_ratio (const Walker *walker_post, const Walker *walker_pre, int particle)

Method for calculating the spatial wave function ratios between two subsequent time steps.

• double get_spatial_lapl_sum (const Walker *walker) const

Method for calculating the sum of all Laplacians for a given walker.

• bool allow_transition ()

Infinite potential to simulate bosonic behaviour.

- void copy_walker (const Walker *parent, Walker *child) const
- void update_walker (Walker *walker_pre, const Walker *walker_post, int particle)
 const
- void reset_walker (const Walker *walker_pre, Walker *walker_post, int particle) const
- double get_spatial_wf (const Walker *walker)
- void initialize (Walker *walker)
- void calc_for_newpos (const Walker *walker_old, Walker *walker_new, int i)

Protected Attributes

• int a

The hard core radius for the infinite potential.

bool overlap

True if the relative distance is less than the hard core radius.

3.6.1 Detailed Description

The Boson system class.

3.6.2 Member Function Documentation

```
3.6.2.1 void Bosons::calc_for_newpos ( const Walker * walker_old, Walker * walker_new, int i) [inline, virtual]
```

Does nothing.

Implements System.

```
3.6.2.2 void Bosons::copy_walker ( const Walker * parent, Walker * child ) const [inline, virtual]
```

Does nothing.

Implements System.

```
3.6.2.3 void Bosons::get_spatial_grad ( Walker * walker, int particle ) const [virtual]
```

Method for calculating the changed part of the spatial gradient.

Depending on which particle we moved, one of the spatial wave function parts (it is split) will be unchanged.

Implements System.

```
3.6.2.4 double Bosons::get_spatial_wf(const Walker * walker) [virtual]
```

The single particle states of each particle multiplied. Assumes the trial wave function initializes every particle in the same single particle state.

Implements System.

```
3.6.2.5 void Bosons::initialize ( Walker * walker ) [inline, virtual]
```

Does nothing.

Implements System.

3.6.2.6 void Bosons::reset_walker (const Walker * walker_pre, Walker * walker_post, int particle) const [inline, virtual]

Does nothing.

Implements System.

3.6.2.7 void Bosons::update_walker (Walker * walker_pre, const Walker * walker_post, int particle) const [inline, virtual]

Does nothing.

Implements System.

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

- src/System/Bosons/Bosons.h
- src/System/Bosons/Bosons.cpp

3.7 Brute Force Class Reference

Implementation of the Brute Force QMC. Uses the Simle diffusion class. All methods are empty except for the energy necessities part which requires the gradients to be calculated (not using the Quantum Force).

Inheritance diagram for Brute_Force:



Public Member Functions

- Brute Force (GeneralParams &)
- void update_walker (Walker *walker_pre, const Walker *walker_post, int particle)
 const
- void get_necessities (Walker *walker)

Method for calculating the sampling specific necessary values.

void update_necessities (const Walker *walker_pre, Walker *walker_post, int particle) const

Method for updating the sampling specific necessary values.

· void calculate energy necessities (Walker *walker) const

Method for calculating the sampling specific necessary values in order to compute the local energy.

- void copy_walker (const Walker *parent, Walker *child) const
 Method for copying the sampling specific parts of a walker.
- void reset_walker (const Walker *walker_pre, Walker *walker_post, int particle)
 const

3.7.1 Detailed Description

Implementation of the Brute Force QMC. Uses the Simle diffusion class. All methods are empty except for the energy necessities part which requires the gradients to be calculated (not using the Quantum Force).

3.7.2 Member Function Documentation

```
3.7.2.1 void Brute_Force::copy_walker ( const Walker * parent, Walker * child ) const [inline, virtual]
```

Method for copying the sampling specific parts of a walker.

See also

```
QMC::copy_walker()
```

Implements Sampling.

Method for calculating the sampling specific necessary values.

Called after a trial position is set.

Implements Sampling.

```
3.7.2.3 void Brute_Force::reset_walker ( const Walker * walker_pre, Walker * walker_post, int particle ) const [inline, virtual]
```

See also

```
QMC::reset_walker()
```

Implements Sampling.

```
3.7.2.4 void Brute_Force::update_walker( Walker * walker_pre, const Walker * walker_post, int particle) const [inline, virtual]
```

See also

QMC::update_walker()

Implements Sampling.

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

- src/Sampling/Brute_Force/Brute_Force.h
- src/Sampling/Brute_Force/Brute_Force.cpp

3.8 Coulomb Class Reference

Implementation of the Coulomb potential. 1/r_{ij}.

Inheritance diagram for Coulomb:



Public Member Functions

- Coulomb (GeneralParams &)
- double get_pot_E (const Walker *walker) const
 Method for calculating a walker's potential energy.

3.8.1 Detailed Description

Implementation of the Coulomb potential. 1/r_{ij}.

3.8.2 Member Function Documentation

3.8.2.1 double Coulomb::get_pot_E (const Walker * walker) const [virtual]

Method for calculating a walker's potential energy.

Method overridden by subclasses.

Parameters

walker The walker for which the potential energy should be calculated.

Returns

The potential energy.

Implements Potential.

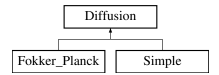
The documentation for this class was generated from the following files:

- src/Potential/Coulomb/Coulomb.h
- src/Potential/Coulomb/Coulomb.cpp

3.9 Diffusion Class Reference

Class containing rules for walker movement based on diffusion models. Serves as class member in the Sampling class. Brute force implies the Simple diffusion model, while - Importance Sampling implies the Fokker Planck diffusion.

Inheritance diagram for Diffusion:



Public Member Functions

- **Diffusion** (int n_p, int dim, double timestep, seed_type random_seed, double D)
- virtual double get_new_pos (const Walker *walker, int i, int j)

Virtual function returning the new position.

 virtual double get_g_ratio (const Walker *walker_post, const Walker *walker_pre) const =0

Calculates the Diffusion Green's function ratio needed by metropolis.

• double call_RNG ()

Calls a uniform random number generator.

- void set_qmc_ptr (QMC *qmc)
- void set_dt (double dt)

Function for altering the time step.

- double get_dt () const
- double $\mathbf{get_std}$ () const

Protected Attributes

- int n_p
- · int dim

• QMC * qmc

The qmc main solver object. Not needed?

· double timestep

The discrete time step.

double D

The diffusion constant.

· long random seed

The random seed. Needs to be stored for some RNGs to work.

double std

The standard deviation from QMC stored for efficiency. sqrt(2D*timestep).

3.9.1 Detailed Description

Class containing rules for walker movement based on diffusion models. Serves as class member in the Sampling class. Brute force implies the Simple diffusion model, while - Importance Sampling implies the Fokker Planck diffusion.

See also

Brute_Force, Importance.

3.9.2 Member Function Documentation

```
3.9.2.1 double Diffusion::call_RNG()
```

Calls a uniform random number generator.

Returns a random uniform number on [0,1).

```
3.9.2.2 virtual double Diffusion::get_g_ratio ( const Walker * walker_post, const Walker * walker_pre ) const [pure virtual]
```

Calculates the Diffusion Green's function ratio needed by metropolis.

Parameters

walker_post	Walker at current time step.
walker_pre	Walker at previous time step.

Returns

The Diffusion Green's function ratio.

Implemented in Simple, and Fokker_Planck.

Virtual function returning the new position.

Returns the simple diffusion step if not overridden.

Parameters

i	Particle number.
j	dimension (x,y,z).

Returns

The new position (relative to the old).

Reimplemented in Fokker_Planck, and Simple.

```
3.9.2.4 void Diffusion::set_dt ( double dt ) [inline]
```

Function for altering the time step.

Takes care of consequences. Time step should only be altered using this function.

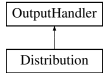
The documentation for this class was generated from the following files:

- src/Diffusion/Diffusion.h
- src/Diffusion/Diffusion.cpp

3.10 Distribution Class Reference

Class for calculating distribution functions such as the one-body density. Does not collect data itself, but works merely as a control organ for the QMC class, calling it's methods for storing position data.

Inheritance diagram for Distribution:



Public Member Functions

Distribution (ParParams &, std::string path, std::string name)
 Constructor.

- void dump ()
- · void finalize ()

Method for calculating the distribution.

• void rerun (int n_p, int N, double bin_edge=0)

Method for re-calculating the distribution given a stores set of position data.

3.10.1 Detailed Description

Class for calculating distribution functions such as the one-body density. Does not collect data itself, but works merely as a control organ for the QMC class, calling it's methods for storing position data.

3.10.2 Constructor & Destructor Documentation

3.10.2.1 Distribution::Distribution (ParParams & pp, std::string path, std::string name)

Constructor.

Parameters

path	The path where output is stored (or read).
name	Name of the file.

3.10.3 Member Function Documentation

```
3.10.3.1 void Distribution::dump() [virtual]
```

Signals QMC solver to store position data.

Implements OutputHandler.

```
3.10.3.2 void Distribution::finalize() [virtual]
```

Method for calculating the distribution.

Overrides the superclass implementation.

Reimplemented from OutputHandler.

```
3.10.3.3 void Distribution::rerun ( int n_p, int N, double bin_edge = 0 )
```

Method for re-calculating the distribution given a stores set of position data.

Scatters the data across nodes.

Parameters

n_p	Number of particles in the set.
N	Number of mesh points used in the histogram.
bin_edge	The Cartesian position of the end points of the histogram.

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

- src/OutputHandler/Distribution/Distribution.h
- src/OutputHandler/Distribution/Distribution.cpp

3.11 DMC Class Reference

Implementation of the Diffusion Monte-Carlo Method. Very little needs to be added when the QMC superclass holds all the general functionality.

Inheritance diagram for DMC:



Public Member Functions

 DMC (GeneralParams &, DMCparams &, SystemObjects &, ParParams &, VMC *vmc, bool dist_out)

Constructor.

void run_method ()

Method used for executing the solver.

• void output ()

Method for standard output.

Static Public Attributes

• static const int K = 50

Factor of empty space for walkers over initial walkers.

- static const int check_thresh = 25
- static const int sendcount_thresh = 20

Protected Member Functions

• void set trial positions ()

Method for setting the trial position of all DMC walkers.

• void iterate walker (int k)

Method for iterating a walker one time step.

void Evolve_walker (int k, double GB)

Method for the birth/death process of a walker.

void bury_the_dead ()

Method for cleaning up the dead walkers and compact the list.

void update_energies ()

Method for updating the DMC energy and calculating the new trial energy.

- bool move authorized (double A)
- void reset_parameters ()
- void node_comm ()
- void save_distribution ()

Method for storing positional data for the Distribtuon.

void switch souls (int root, int root id, int dest, int dest id)

Method for sending a walker between two nodes.

• void normalize_population ()

Method for evening out the number of walkers on each node.

• void free_walkers ()

Method which deletes all walkers.

Protected Attributes

bool thermalized

Flag used to indicate whether to start sampling or not.

• int n w last

The amount of walkers at the time the walker loop was initiated.

• int n_w_tot

The total number of walkers across all nodes.

• arma::uvec n_w_list

List of the number of walkers of each node.

• bool force_comm

Flag set true if population should be renormalized.

- int deaths
- · int block size
- int samples
- double dmc E

The DMC energy.

double dmc_E_unscaled

The accumulative DMC energy: The sum of all previous trial energies.

• double E T

The trial energy at the current cycle.

• double E

The accumulative energy for each cycle.

Friends

· class stdoutDMC

3.11.1 Detailed Description

Implementation of the Diffusion Monte-Carlo Method. Very little needs to be added when the QMC superclass holds all the general functionality.

3.11.2 Member Function Documentation

```
3.11.2.1 void DMC::Evolve_walker(int k, double GB) [protected]
```

Method for the birth/death process of a walker.

Parameters

GB	The branching Green's Function.
k	The index of the walker.

3.11.2.2 void DMC::iterate_walker(int k) [protected]

Method for iterating a walker one time step.

Parameters

thermalized	Flag to indicate whether to start sampling or not.
k	The index of the walker.

In case of DMC, we must let the system have the possibility to override the metropolis test (fixed node approximation in case of a Fermion system)

Implements QMC.

```
3.11.2.4 void DMC::node_comm() [protected, virtual]
```

For each process: -Calculates the total number of walkers. -Sums up the energies sampled. -Sums up the total number of samples made.

Implements QMC.

```
3.11.2.5 void DMC::save_distribution() [protected, virtual]
```

Method for storing positional data for the Distribtuon.

Stores the position data from all currently alive $\ensuremath{\mathsf{DMC}}$ walkers every dist_tresh cycle.

Implements QMC.

```
3.11.2.6 void DMC::set_trial_positions() [protected, virtual]
```

Method for setting the trial position of all DMC walkers.

In case VMC is not run prior to DMC, trial positions must be set.

Implements QMC.

```
3.11.2.7 void DMC::switch_souls ( int root, int root_id, int dest, int dest_id ) [protected]
```

Method for sending a walker between two nodes.

Parameters

root	The node from which the walker is sent.
root_id	The index of the walker being sent from root.
dest	The node which receives the walker.
dest_id	The index where the walker is to be received.

See also

```
Walker::send_soul(), Walker::recv_soul()
```

3.11.3 Member Data Documentation

```
3.11.3.1 const int DMC::K = 50 [static]
```

Factor of empty space for walkers over initial walkers.

See also

QMC::QMC()

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

- src/QMC/DMC/DMC.h
- src/QMC/DMC/DMC.cpp

3.12 DMCparams Struct Reference

Struct used to initialize DMC parameters.

Public Attributes

• int n_c

The number of cycles.

int therm

Thermalization cycles.

• int n b

Number of block samples pr. walker pr. cycle.

• int n w

Number of walkers.

• double dt

Time step.

3.12.1 Detailed Description

Struct used to initialize DMC parameters.

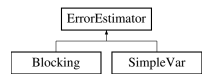
The documentation for this struct was generated from the following file:

• src/QMCheaders.h

3.13 ErrorEstimator Class Reference

Class handling error estimations of the QMC methods. The QMC class holds an object of this type, calling the update_data function in order to update the sampling pool. finalize() then either dumps the samples to file for later processing, or calulates an estimate.

Inheritance diagram for ErrorEstimator:



Public Member Functions

• ErrorEstimator (int n_c, std::string filename, std::string path, bool parallel, int node, int n_nodes, bool rerun=false)

Constructor.

• double combine_variance (double var, double mean=0, int n=0)

Calculates the combined variance of n_nodes variances.

- void finalize ()
- void node comm gather data ()
- void node_comm_scatter_data ()
- void init_file ()

Opens a file with filename at path supplied in constructor.

• virtual double estimate_error ()=0

Estimates the error based on the subclass implementation.

• virtual void update_data (double val)

Adds values to the data vector.

Static Public Member Functions

• static double combine_mean (double mean, int n, int n_tot=0)

Calculates the combined mean of n_nodes means.

Public Attributes

· bool data_to_file

If true, the data vector are stored to file.

· bool output_to_file

If init_file() method is called, this flag is true.

Protected Attributes

• int n_c

Size of the data vector.

• int i

Count variable for the data vector.

- bool parallel
- bool is_master
- int node
- int n nodes
- bool rerun

If false, data is assumed to already exist.

- std::string filename
- · std::string path

- std::ofstream file
- · arma::rowvec data

The vector containing the samples used in error calculation.

3.13.1 Detailed Description

Class handling error estimations of the QMC methods. The QMC class holds an object of this type, calling the update_data function in order to update the sampling pool. finalize() then either dumps the samples to file for later processing, or calulates an estimate.

3.13.2 Constructor & Destructor Documentation

3.13.2.1 ErrorEstimator::ErrorEstimator (int n_c , std::string filename, std::string path, bool parallel, int node, int n_n nodes, bool rerun = false)

Constructor.

Parameters

n_c	The expected number of samples to be stored.
filename	The name of the file. Only necessary if init_file() is called.
path	The path where the data and/or the file is stored (or read).

3.13.3 Member Function Documentation

3.13.3.1 double ErrorEstimator::combine_mean (double mean, int n, int n_tot = 0) [static]

Calculates the combined mean of n_nodes means.

Only useful for parallel calls.

Parameters

mean	The local mean on an individual node
n	The number of samples used to calculate the local mean.
n_tot	The total number of samples on all nodes. Calculated if not supplied.

3.13.3.2 double ErrorEstimator::combine_variance (double var, double mean = 0, int n = 0)

Calculates the combined variance of n_nodes variances.

Only useful for parallel calls.

Parameters

var	The local variance on an individual node
mean	The local mean on an individual node
n_tot	The total number of samples used on all nodes.

3.13.3.3 void ErrorEstimator::finalize ()

if [output_to_file]: Closes opened files if [data_to_file]: Stores accumulated data. if data vector was used, it's memory is freed.

```
3.13.3.4 void ErrorEstimator::init_file()
```

Opens a file with filename at path supplied in constructor.

Subclass implementations can call this function. Superclass does not.

```
3.13.3.5 void ErrorEstimator::node_comm_gather_data()
```

Gathers the data vectors from all processes into a single one on the master node.

```
3.13.3.6 void ErrorEstimator::node_comm_scatter_data( )
```

Exact reverse of node_comm_gather_data()

3.13.3.7 void ErrorEstimator::update_data(double *val***)** [virtual]

Adds values to the data vector.

Can be overridden if storage is not wanted.

Parameters

val A local sample of the quantity of which the error is calculated	
---	--

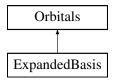
Reimplemented in SimpleVar.

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

- src/ErrorEstimator/ErrorEstimator.h
- src/ErrorEstimator/ErrorEstimator.cpp

3.14 ExpandedBasis Class Reference

Inheritance diagram for ExpandedBasis:



Public Member Functions

- ExpandedBasis (GeneralParams &gp, Orbitals *basis, int m, std::string coeff-Path)
- double phi (const Walker *walker, int particle, int q_num)

Calculates the single particle wave function for a given walker's particle.

double del_phi (const Walker *walker, int particle, int q_num, int d)

Calculates the single particle wave function derivative for a given walker's particle and dimension.

double lapl_phi (const Walker *walker, int particle, int q_num)

Calculates the single particle wave function for a given walker's particle.

• void set_qnum_indie_terms (const Walker *walker, int i)

Calculates single particle wave function terms which are independent of the quantum numbers.

Protected Attributes

- int basis_size
- · arma::mat coeffs
- Orbitals * basis

3.14.1 Member Function Documentation

3.14.1.1 double ExpandedBasis::del_phi (const Walker * walker, int particle, int q_num, int d) [virtual]

Calculates the single particle wave function derivative for a given walker's particle and dimension.

Parameters

Γ	q_num	The quantum number index.
ľ	d	The dimension for which the derivative should be calculated (x,y,z).

Reimplemented from Orbitals.

3.14.1.2 double ExpandedBasis::lapl_phi (const Walker * walker, int particle, int q_num) [virtual]

Calculates the single particle wave function for a given walker's particle.

Parameters

```
q_num The quantum number index.
```

Reimplemented from Orbitals.

3.14.1.3 double ExpandedBasis::phi (const Walker * walker, int particle, int q_num) [virtual]

Calculates the single particle wave function for a given walker's particle.

Parameters

```
q_num The quantum number index.
```

Reimplemented from Orbitals.

3.14.1.4 void ExpandedBasis::set_qnum_indie_terms (const Walker * walker, int i) [inline, virtual]

Calculates single particle wave function terms which are independent of the quantum numbers.

If a term in the single particle functions are independent of the quantum number, this function can be overridden to calculate them beforehand (for each particle), and rather extract the value instead of recalculating.

Parameters

```
i Particle number.
```

Reimplemented from Orbitals.

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

- $\bullet \ src/Orbitals/ExpandedBasis/ExpandedBasis.h$
- src/Orbitals/ExpandedBasis/ExpandedBasis.cpp

3.15 Fermions Class Reference

The Fermion system class.

Inheritance diagram for Fermions:



Public Member Functions

- Fermions (GeneralParams &, Orbitals *)
- void get_spatial_grad (Walker *walker, int particle) const

Method for calculating the changed part of the spatial gradient.

void get_spatial_grad_full (Walker *walker) const

Method for calculating the full spatial gradient.

double get_spatial_ratio (const Walker *walker_post, const Walker *walker_pre, int particle)

Method for calculating the spatial wave function ratios between two subsequent time steps.

double get spatial lapl sum (const Walker *walker) const

Method for calculating the sum of all Laplacians for a given walker.

• bool allow transition ()

Fixed node approximation.

- void copy_walker (const Walker *parent, Walker *child) const
- void update_walker (Walker *walker_pre, const Walker *walker_post, int particle)
- void reset_walker (const Walker *walker_pre, Walker *walker_post, int particle) const
- double get_spatial_wf (const Walker *walker)
- void initialize (Walker *walker)
- void calc_for_newpos (const Walker *walker_old, Walker *walker_new, int i)

Protected Member Functions

• void make_merged_inv (Walker *walker)

Method for calculating the Slater matrix inverse.

• void update_inverse (const Walker *walker_old, Walker *walker_new, int particle)

Protected Attributes

• arma::rowvec |

The diagonal of the new slater matrix times the old slater inverse.

Method for updating the inverse given that we moved one particle.

· bool node_crossed

True if the spatial ratio is negative.

3.15.1 Detailed Description

The Fermion system class.

3.15.2 Member Function Documentation

```
3.15.2.1 void Fermions::calc_for_newpos ( const Walker * walker_old, Walker * walker_new, int i) [inline, virtual]
```

When a particle is moved, the inverse is updated.

Implements System.

```
3.15.2.2 void Fermions::copy_walker ( const Walker * parent, Walker * child ) const [inline, virtual]
```

Copies the inverse.

Implements System.

```
3.15.2.3 void Fermions::get_spatial_grad ( Walker * walker, int particle ) const [virtual]
```

Method for calculating the changed part of the spatial gradient.

Depending on which particle we moved, one of the spatial wave function parts (it is split) will be unchanged.

Implements System.

The determinant of each spin value multiplied.

Implements System.

```
3.15.2.5 void Fermions::initialize ( Walker * walker ) [inline, virtual]
```

Calculates the inverse.

Implements System.

```
3.15.2.6 void Fermions::make_merged_inv( Walker * walker) [protected]
```

Method for calculating the Slater matrix inverse.

The merged inverse is made by concatenating the two slater matrix inverses. This way we can sum freely over particles without having to if-test on the spin.

```
3.15.2.7 void Fermions::reset_walker ( const Walker * walker_pre, Walker * walker_post, int particle ) const [inline, virtual]
```

Resets the inverse.

Implements System.

```
3.15.2.8 void Fermions::update_walker ( Walker * walker_pre, const Walker * walker_post, int particle ) const [inline, virtual]
```

Updates the inverse.

Implements System.

3.15.3 Member Data Documentation

```
3.15.3.1 arma::rowvec Fermions::I [protected]
```

The diagonal of the new slater matrix times the old slater inverse.

Needed for updating the inverse. Stored because only half of the vector is changed when moving one particle.

See also

```
System::set_spin_state()
```

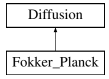
The documentation for this class was generated from the following files:

- src/System/Fermions/Fermions.h
- src/System/Fermions/Fermions.cpp

3.16 Fokker Planck Class Reference

Anisotropic diffusion by the Fokker-Planck equation.

Inheritance diagram for Fokker_Planck:



Public Member Functions

- Fokker_Planck (int n_p, int dim, double timestep, seed_type random_seed, double D=0.5)
- double get_g_ratio (const Walker *walker_post, const Walker *walker_pre)
 const

Calculates the Diffusion Green's function ratio needed by metropolis.

• double get_new_pos (const Walker *walker, int i, int j)

Virtual function returning the new position.

3.16.1 Detailed Description

Anisotropic diffusion by the Fokker-Planck equation.

3.16.2 Member Function Documentation

```
3.16.2.1 double Fokker_Planck::get_g_ratio ( const Walker * walker_post, const Walker * walker_pre ) const [virtual]
```

Calculates the Diffusion Green's function ratio needed by metropolis.

Parameters

walker_post	Walker at current time step.
walker_pre	Walker at previous time step.

Returns

The Diffusion Green's function ratio.

Implements Diffusion.

```
3.16.2.2 double Fokker_Planck::get_new_pos ( const Walker * walker, int i, int j ) [inline, virtual]
```

Virtual function returning the new position.

Returns the simple diffusion step if not overridden.

Parameters

i	Particle number.
j	dimension (x,y,z).

Returns

The new position (relative to the old).

Reimplemented from Diffusion.

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

- src/Diffusion/Fokker_Planck/Fokker_Planck.h
- src/Diffusion/Fokker_Planck/Fokker_Planck.cpp

3.17 GeneralParams Struct Reference

Struct used to initialize general parameters.

Public Attributes

• int n_p

The number of particles.

• int dim

The dimension.

• seed_type random_seed

The random number generator's seed.

• double systemConstant

The constant used in systems.

- bool doMIN
- bool doVMC
- bool doDMC
- bool do_blocking
- bool use_jastrow
- bool use_coulomb
- std::string system

String specifing the system type, e.g. "Atoms".

• std::string sampling

String specifing the sampling type, e.g. "IS".

• std::string runpath

The directory which the simulation is set to run.

3.17.1 Detailed Description

Struct used to initialize general parameters.

3.17.2 Member Data Documentation

3.17.2.1 double GeneralParams::systemConstant

The constant used in systems.

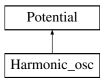
e.g. charge for atoms and oscillator frequency for quantum dots.

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

· src/QMCheaders.h

3.18 Harmonic_osc Class Reference

Implementation of the Harmonic Oscillator potential. 0.5*w**2*r**2. Inheritance diagram for Harmonic_osc:



Public Member Functions

- Harmonic_osc (GeneralParams &)
- double get_pot_E (const Walker *walker) const
 Method for calculating a walker's potential energy.

Protected Attributes

• double w

The oscillator frequency.

3.18.1 Detailed Description

Implementation of the Harmonic Oscillator potential. 0.5*w**2*r**2.

3.18.2 Member Function Documentation

Method for calculating a walker's potential energy.

Method overridden by subclasses.

Parameters

walker	The walker for which the potential energy should be calculated.
--------	---

Returns

The potential energy.

Implements Potential.

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

- src/Potential/Harmonic_osc/Harmonic_osc.h
- src/Potential/Harmonic_osc/Harmonic_osc.cpp

3.19 HartreeFock Class Reference

Public Member Functions

- HartreeFock (int m, Orbitals *sp_basis, double thresh=1e-5)
- void run_method ()

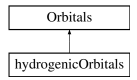
The documentation for this class was generated from the following files:

- src/HartreeFock/HartreeFock.h
- src/HartreeFock/HartreeFock.cpp

3.20 hydrogenicOrbitals Class Reference

Hyrogen-like single particle wave function class. Uses the hydrogenic BasisFunction subclasses auto-generated by SymPy through the orbitalsGenerator tool.

Inheritance diagram for hydrogenicOrbitals:



Public Member Functions

- hydrogenicOrbitals (GeneralParams &, VariationalParams &)
- void set_qnum_indie_terms (const Walker *walker, int i)

Protected Member Functions

- double get variational derivative (const Walker *walker, int n)
 - ! Overridden superclass method implementing closed form expressions generated by SymPy.
- double get_dell_alpha_phi (const Walker *walker, int qnum, int i)

Method for calculating a single particle wave functions variational derivative.

- · double get_sp_energy (int qnum) const
- double get_coulomb_element (const arma::uvec &qnum_set)

Method for calculating the anti-symmetrized Coulumb matrix elements.

- · void get_qnums ()
- double get_parameter (int n)
- void set parameter (double parameter, int n)

Protected Attributes

• int Z

The charge of the core.

double * alpha

Pointer to the variational parameter alpha. Shared address with all the BasisFunction subclasses.

double * k

Pointer to alpha*Z. Shared address with all the BasisFunction subclasses.

double * k2

Pointer (alpha*Z) $^{\wedge}$ 2. Shared address with all the BasisFunction subclasses.

double * exp factor n1

Pointer to a factor precalculated by set_qnum_indie_terms(). Shared address with all the BasisFunction subclasses.

double * exp_factor_n2

Pointer to a factor precalculated by set_qnum_indie_terms(). Shared address with all the BasisFunction subclasses.

double * exp_factor_n3

Pointer to a factor precalculated by set_qnum_indie_terms(). Shared address with all the BasisFunction subclasses.

double * exp_factor_n4

Pointer to a factor precalculated by set_qnum_indie_terms(). Shared address with all the BasisFunction subclasses.

Friends

· class ExpandedBasis

3.20.1 Detailed Description

Hyrogen-like single particle wave function class. Uses the hydrogenic BasisFunction subclasses auto-generated by SymPy through the orbitalsGenerator tool.

3.20.2 Member Function Documentation

Method for calculating the anti-symmetrized Coulumb matrix elements.

Used by Hartree Fock

Reimplemented from Orbitals.

```
3.20.2.2 double hydrogenicOrbitals::get_dell_alpha_phi(const Walker * walker, int qnum, int i) [protected]
```

Method for calculating a single particle wave functions variational derivative.

Parameters

```
i The particle number.
```

Returns

The variational parameter alpha.

Implements Orbitals.

```
3.20.2.4 void hydrogenicOrbitals::set_parameter ( double parameter, int n ) [inline, protected, virtual]
```

Sets a new value for the alpha and updates all the pointer values.

Implements Orbitals.

Calculates the exponential terms exp(-r/n) for all needed n once pr. particle to save CPU-time.

See also

Orbitals::set_qnum_indie_terms()

Reimplemented from Orbitals.

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

- src/Orbitals/hydrogenicOrbitals/hydrogenicOrbitals.h
- src/Orbitals/hydrogenicOrbitals/hydrogenicOrbitals.cpp

3.21 Importance Class Reference

Implementation of Importance sampled QMC. Using the Fokker-Planck diffusion class. Introduces the Quantum Force.

Inheritance diagram for Importance:



Public Member Functions

- Importance (GeneralParams &)
- void update_walker (Walker *walker_pre, const Walker *walker_post, int particle)
- void reset_walker (const Walker *walker_pre, Walker *walker_post, int particle)
- void get_necessities (Walker *walker)
- void update_necessities (const Walker *walker_pre, Walker *walker_post, int particle) const
- void calculate_energy_necessities (Walker *walker) const
- void copy walker (const Walker *parent, Walker *child) const

3.21.1 Detailed Description

Implementation of Importance sampled QMC. Using the Fokker-Planck diffusion class. Introduces the Quantum Force.

3.21.2 Member Function Documentation

```
3.21.2.1 void Importance::calculate_energy_necessities ( Walker * walker ) const [inline, virtual]
```

No energy necessities (they are already calculated).

Implements Sampling.

```
3.21.2.2 void Importance::copy_walker ( const Walker * parent, Walker * child ) const [inline, virtual]
```

The gradients and the Quantum force is copied.

Implements Sampling.

The gradients and the Quantum force are calculated.

Implements Sampling.

```
3.21.2.4 void Importance::reset_walker ( const Walker * walker_pre, Walker * walker_post, int particle ) const [virtual]
```

The parts of the gradients with the same spin as the moved particle are reset.

Implements Sampling.

```
3.21.2.5 void Importance::update_necessities ( const Walker * walker_pre, Walker * walker_post, int particle ) const [inline, virtual]
```

The gradients are updated and the Quantum force is re-calculated.

Implements Sampling.

```
3.21.2.6 void Importance::update_walker ( Walker * walker_pre, const Walker * walker_post, int particle ) const [virtual]
```

The parts of the gradients with the same spin as the moved particle are updated.

Implements Sampling.

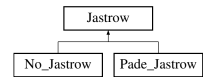
The documentation for this class was generated from the following files:

- src/Sampling/Importance/Importance.h
- src/Sampling/Importance/Importance.cpp

3.22 Jastrow Class Reference

The class representing the Jastrow correlation functions Holds all data concerning the Jastrow function and it's influence on the QMC algorithm.

Inheritance diagram for Jastrow:



Public Member Functions

- Jastrow (int n_p, int dim)
- virtual void initialize ()=0
- virtual double get_val (const Walker *walker) const =0
- virtual double get_j_ratio (const Walker *walker_new, const Walker *walker_old, int i) const =0

Calculates the ratio of the Jastrow factor needed by metropolis.

- virtual void get_grad (Walker *walker) const =0
- virtual void get_grad (const Walker *walker_pre, Walker *walker_post, int i) const
 =0

Updates the gradient for a new particle move.

virtual void get_dJ_matrix (Walker *walker, int i) const =0

Updates the summation factors of Jastrow factors closed form expressions.

void get_dJ_matrix (Walker *walker) const

Calculates the summation factors of Jastrow factors closed form expressions.

• virtual double get_lapl_sum (Walker *walker) const =0

Method for calculating the Laplacian.

Protected Member Functions

virtual double get parameter (int n)=0

Returns variational parameters.

• virtual void set_parameter (double param, int n)=0

Sets variational parameters.

virtual double get variational derivative (const Walker *walker, int n)

Calculates the derivative of the Jastrow factor with respect to a variational parameter.

• double get_derivative_num (Walker *walker, int i, int d) const

Numerical Cartesian derivative.

double get_laplaciansum_num (Walker *walker) const

Numerical Cartesian Laplacian.

Protected Attributes

- int n p
- int **n2**
- int dim
- · bool active

Parameter false if No_Jastrow is loaded.

Friends

- · class Minimizer
- · class ASGD
- · class stdoutASGD

3.22.1 Detailed Description

The class representing the Jastrow correlation functions Holds all data concerning the Jastrow function and it's influence on the QMC algorithm.

3.22.2 Member Function Documentation

3.22.2.1 double Jastrow::get_derivative_num (Walker * walker, int i, int d) const [protected]

Numerical Cartesian derivative.

For use in get_grad() when no closed form expression is implemented.

Parameters

i	Particle number.
d	Dimension (x,y,z).

3.22.2.2 virtual void Jastrow::get_dJ_matrix (Walker * *walker*, int *i*) const [pure virtual]

Updates the summation factors of Jastrow factors closed form expressions.

Used to optimize the calculations as few of these terms change as we move a particle.

Parameters

i	Particle number.

Implemented in Pade_Jastrow, and No_Jastrow.

3.22.2.3 void Jastrow::get_dJ_matrix (Walker * walker) const

Calculates the summation factors of Jastrow factors closed form expressions.

Used to optimize the calculations as few of these terms change as we move a particle.

3.22.2.4 virtual void Jastrow::get_grad (Walker * *walker* **) const** [pure virtual]

Calculates the entire Cartesian gradient.

Implemented in Pade Jastrow, and No Jastrow.

Updates the gradient for a new particle move.

Parameters

walker_post	Walker at current time step
walker_pre	Walker at previous time step
i	Particle number.

Implemented in Pade_Jastrow, and No_Jastrow.

3.22.2.6 virtual double Jastrow::get_j_ratio (const Walker * walker_new, const Walker * walker_old, int i) const [pure virtual]

Calculates the ratio of the Jastrow factor needed by metropolis.

Parameters

walker_new	Walker at current time step
walker_old	Walker at previous time step
i	The particle number.

Implemented in Pade_Jastrow, and No_Jastrow.

3.22.2.7 virtual double Jastrow::get_lapl_sum (Walker * walker) const [pure virtual]

Method for calculating the Laplacian.

Calculates the sum of all particles Laplacians.

Implemented in No_Jastrow, and Pade_Jastrow.

3.22.2.8 double Jastrow::get_laplaciansum_num (Walker * walker) const [protected]

Numerical Cartesian Laplacian.

For use in get_lapl_sum() when no closed form expression is implemented.

Returns variational parameters.

Parameters

n The index of the sought variational parameter

Returns

Variational parameter with index [n]

Implemented in Pade_Jastrow, and No_Jastrow.

3.22.2.10 virtual double Jastrow::get_val (const Walker * walker) const [pure virtual]

Calculates the value of the Jastrow Factor at the walker's position.

Implemented in Pade_Jastrow, and No_Jastrow.

Calculates the derivative of the Jastrow factor with respect to a variational parameter.

Parameters

n	The index of the variational parameter for which the derivative is to be taken
walker	The walker holds the positions etc. needed to evaluate the derivative

Reimplemented in No_Jastrow, and Pade_Jastrow.

3.22.2.12 virtual void Jastrow::initialize() [pure virtual]

Initializes the non-variational parameters needed by the Jastrow Factor.

Implemented in Pade_Jastrow, and No_Jastrow.

Sets variational parameters.

Parameters

n	The index of the sought variational parameter
param	The new value of parameter [n]

Implemented in No_Jastrow, and Pade_Jastrow.

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

- · src/Jastrow/Jastrow.h
- src/Jastrow/Jastrow.cpp

3.23 Minimizer Class Reference

Class for minimization methods used to obtain optimal variational parameters.

Inheritance diagram for Minimizer:



Public Member Functions

 Minimizer (VMC *vmc, const ParParams &, const arma::rowvec &alpha, const arma::rowvec &beta)

Constructor.

void add_output (OutputHandler *output_handler)

Method used for loading the stdoutASGD object.

- Orbitals * get_orbitals ()
- Jastrow * get_jastrow ()
- virtual void minimize ()=0

Method for executing the minimization main solver.

• void output (std::string message, double number=-1)

Method for dumping variational parameter values to screen.

• void add_error_estimator (ErrorEstimator *error_estimator)

Method used to add error estimators.

Protected Member Functions

- void dump output ()
- void finalize_output ()
- void error_output ()
- virtual void update_parameters ()=0

Method for updating the variational parameters based on the previous step.

Protected Attributes

- int n nodes
- bool is_master
- VMC * vmc

Uses VMC methods to calculate stochastic variational gradients.

• STDOUT * std out

Output object. Wraps and replaces std::cout.

- std::stringstream s
- int Nspatial_params

The number of variational parameters in the spatial trial wave function.

• int Njastrow_params

The number of variational parameters in the Jastrow factor.

• int Nparams

The total number of variational parameters.

std::vector< OutputHandler * > output_handler

Either contains a stdoutASGD object or not.

• std::vector< ErrorEstimator * > error_estimators

One ErrorEstimator object pr. variational parameter.

3.23.1 Detailed Description

Class for minimization methods used to obtain optimal variational parameters.

3.23.2 Constructor & Destructor Documentation

3.23.2.1 Minimizer::Minimizer (VMC * vmc, const ParParams & pp, const arma::rowvec & alpha, const arma::rowvec & beta)

Constructor.

Parameters

	The VMC object used for storing variational parameters and calculating stochastic gradients.
alpha	Vector of initial conditions of spatial variational parameters
beta	Vector of initial conditions of Jastrow variational parameters

3.23.3 Member Function Documentation

```
3.23.3.1 void Minimizer::dump_output() [protected]
```

Iterates over the output objects in the output_handler vector. No if-tests.

```
3.23.3.2 void Minimizer::error_output() [protected]
```

Estimates and finalizes the ErrorEstimator objects initialized in the error_estimators vector.

```
3.23.3.3 void Minimizer::finalize_output( ) [protected]
```

Calls the finalize function for the object in the output_handler vector.

```
3.23.3.4 virtual void Minimizer::update_parameters ( ) [protected, pure virtual]
```

Method for updating the variational parameters based on the previous step.

Needs to be implemented by a subclass.

Implemented in ASGD.

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

- · src/Minimizer/Minimizer.h
- src/Minimizer/Minimizer.cpp

3.24 MinimizerParams Struct Reference

Struct used to initialize Minimization parameters.

Public Attributes

- double max_step
- double f_max
- · double f min
- · double omega
- · double A
- double a
- · int SGDsamples
- int **n_w**
- · int therm
- int **n** c

- int n c SGD
- arma::rowvec alpha

Initial condition for the spatial variational parameter(s).

arma::rowvec beta

Initial condition for the Jastrow variational parameter(s).

3.24.1 Detailed Description

Struct used to initialize Minimization parameters.

See also

ASGD

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

· src/QMCheaders.h

3.25 No Jastrow Class Reference

Class loaded when no correlation factor is used.

Inheritance diagram for No_Jastrow:



Public Member Functions

- void get grad (Walker *walker) const
- void get_grad (const Walker *walker_pre, Walker *walker_post, int i) const Updates the gradient for a new particle move.
- void initialize ()
- void get_dJ_matrix (Walker *walker, int i) const

Updates the summation factors of Jastrow factors closed form expressions.

 double get_j_ratio (const Walker *walker_post, const Walker *walker_pre, int i) const

Calculates the ratio of the Jastrow factor needed by metropolis.

- double get_val (const Walker *walker) const
- double get_lapl_sum (Walker *walker) const

Method for calculating the Laplacian.

Protected Member Functions

• double get_parameter (int n)

Returns variational parameters.

• void set_parameter (double param, int n)

Sets variational parameters.

• double get_variational_derivative (const Walker *walker, int n)

Calculates the derivative of the Jastrow factor with respect to a variational parameter.

3.25.1 Detailed Description

Class loaded when no correlation factor is used.

3.25.2 Member Function Documentation

Updates the summation factors of Jastrow factors closed form expressions.

Used to optimize the calculations as few of these terms change as we move a particle.

Parameters

i	Particle number.

Implements Jastrow.

Calculates the entire Cartesian gradient.

Implements Jastrow.

```
3.25.2.3 void No_Jastrow::get_grad ( const Walker * walker_pre, Walker * walker_post, int i) const [inline, virtual]
```

Updates the gradient for a new particle move.

Parameters

walker_post	Walker at current time step
walker_pre	Walker at previous time step
i	Particle number.

Implements Jastrow.

```
3.25.2.4 double No_Jastrow::get_j_ratio ( const Walker * walker_new, const Walker * walker_old, int i ) const [inline, virtual]
```

Calculates the ratio of the Jastrow factor needed by metropolis.

Parameters

wal	lker_new	Walker at current time step
Wá	alker_old	Walker at previous time step
	i	The particle number.

Implements Jastrow.

Method for calculating the Laplacian.

Calculates the sum of all particles Laplacians.

Implements Jastrow.

Returns variational parameters.

Parameters

n The index of the sought variational parameter

Returns

Variational parameter with index [n]

Implements Jastrow.

Calculates the value of the Jastrow Factor at the walker's position.

Implements Jastrow.

```
3.25.2.8 double No_Jastrow::get_variational_derivative ( const Walker * walker, int n
) [inline, protected, virtual]
```

Calculates the derivative of the Jastrow factor with respect to a variational parameter.

Parameters

n	The index of the variational parameter for which the derivative is to be
	taken
walker	The walker holds the positions etc. needed to evaluate the derivative

Reimplemented from Jastrow.

```
3.25.2.9 void No_Jastrow::initialize( ) [inline, virtual]
```

Initializes the non-variational parameters needed by the Jastrow Factor.

Implements Jastrow.

Sets variational parameters.

Parameters

n	The index of the sought variational parameter
param	The new value of parameter [n]

Implements Jastrow.

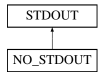
The documentation for this class was generated from the following files:

- src/Jastrow/No_Jastrow/No_Jastrow.h
- src/Jastrow/No Jastrow/No Jastrow.cpp

3.26 NO_STDOUT Class Reference

Class for suppressing standard output. Every node but the master has this. If-tests around cout is avoided.

Inheritance diagram for NO_STDOUT:



Public Member Functions

• virtual void cout (std::stringstream &a)

3.26.1 Detailed Description

Class for suppressing standard output. Every node but the master has this. If-tests around cout is avoided.

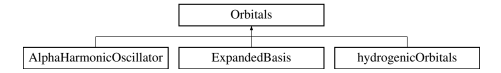
The documentation for this class was generated from the following file:

· src/QMCheaders.h

3.27 Orbitals Class Reference

Superclass for the single particle orbital classes. Handles everything specific regarding choice of single particle basis.

Inheritance diagram for Orbitals:



Public Member Functions

- Orbitals (int n_p, int dim)
- virtual void set_qnum_indie_terms (const Walker *walker, int i)

Calculates single particle wave function terms which are independent of the quantum numbers.

- virtual double phi (const Walker *walker, int particle, int q_num)
 - Calculates the single particle wave function for a given walker's particle.
- virtual double del_phi (const Walker *walker, int particle, int q_num, int d)
 Calculates the single particle wave function derivative for a given walker's particle and
- virtual double lapl_phi (const Walker *walker, int particle, int q_num)
 - Calculates the single particle wave function for a given walker's particle.
- void set_qmc_ptr (QMC *qmc)

Protected Member Functions

dimension.

- virtual double get_parameter (int n)=0
 - A method for retrieving variational parameters.
- virtual void set_parameter (double parameter, int n)=0
 - A method for setting variational parameters.
- virtual double get variational derivative (const Walker *walker, int n)

A method for calculating the variational derivative.

double num_diff (const Walker *walker, int particle, int q_num, int d)

Method for calculating the single particle derivative using a finite difference scheme.

• double num_ddiff (const Walker *walker, int particle, int q_num)

Method for calculating the single particle Laplacian using a finite difference scheme.

void testLaplace (const Walker *walker, int particle, int q_num)

Method for validating closed form expressions for laplacians by comparing them to numerical calculations.

void testDell (const Walker *walker, int particle, int q_num, int d)

Method for validating closed form expressions for derivatives by comparing them to numerical calculations.

• virtual double get_coulomb_element (const arma::uvec &qnum_set)

Method for calculating the anti-symmetrized Coulumb matrix elements.

virtual double get_sp_energy (int qnum) const

Protected Attributes

- int **n_p**
- int **n2**
- int dim
- · int max implemented

The maximum number basis size supported for any system ##RYDD OPP.

• QMC * qmc

A pointer to the QMC solver object. Needed for numerical variational derivatives.

double h

The step length for finite difference schemes.

- double h2
- double two h
- arma::imat qnums

Quantum number matrix needed by Hartree-Fock and the variational derivatives.

BasisFunctions ** basis_functions

A vector maping a quantum number index to a single particle wave function.

• BasisFunctions *** dell basis functions

A maxtrix maping a quantum number- and dimension index to a single particle wave function derivative.

• BasisFunctions ** lapl basis functions

A vector maping a quantum number index to a single particle wave function Laplacian.

Friends

- · class HartreeFock
- · class Minimizer
- · class ASGD
- · class stdoutASGD

3.27.1 Detailed Description

Superclass for the single particle orbital classes. Handles everything specific regarding choice of single particle basis.

3.27.2 Member Function Documentation

```
3.27.2.1 double Orbitals::del_phi ( const Walker * walker, int particle, int q_num, int d ) [virtual]
```

Calculates the single particle wave function derivative for a given walker's particle and dimension.

Parameters

q_num	The quantum number index.
d	The dimension for which the derivative should be calculated (x,y,z).

Reimplemented in ExpandedBasis.

Method for calculating the anti-symmetrized Coulumb matrix elements.

Used by Hartree Fock

Reimplemented in AlphaHarmonicOscillator, and hydrogenicOrbitals.

A method for retrieving variational parameters.

Parameters

n	Index of the sought variational parameter.

Implemented in AlphaHarmonicOscillator, and hydrogenicOrbitals.

3.27.2.4 double Orbitals::get_variational_derivative (const Walker
$$*$$
 walker, int n) [protected, virtual]

A method for calculating the variational derivative.

By default uses a finite difference scheme. Can be overridden to evaluate a closed form expression.

Parameters

n Index of the sought variational parameter.

Reimplemented in AlphaHarmonicOscillator, and hydrogenicOrbitals.

```
3.27.2.5 double Orbitals::lapl_phi( const Walker * walker, int particle, int q_num )
[virtual]
```

Calculates the single particle wave function for a given walker's particle.

Parameters

```
q_num The quantum number index.
```

Reimplemented in ExpandedBasis.

```
3.27.2.6 double Orbitals::num_ddiff ( const Walker * walker, int particle, int q_num ) [protected]
```

Method for calculating the single particle Laplacian using a finite difference scheme.

Method lapl_phi() can be overridden to use this method in case no closed form expressions are implemented.

Parameters

```
q_num The quantum number index.
```

3.27.2.7 double Orbitals::num_diff (const Walker * walker, int particle, int q_num, int d) [protected]

Method for calculating the single particle derivative using a finite difference scheme.

Method del_phi() can be overridden to use this method in case no closed form expressions are implemented.

Parameters

d The dimension for which the derivative should be calculated (x,y,z).

```
3.27.2.8 double Orbitals::phi ( const Walker * walker, int particle, int q_num ) [virtual]
```

Calculates the single particle wave function for a given walker's particle.

Parameters

q_num	The quantum number index.

Reimplemented in ExpandedBasis.

A method for setting variational parameters.

Parameters

n	Index of the sought variational parameter.
parameter	The new value of the variational parameter.

Implemented in AlphaHarmonicOscillator, and hydrogenicOrbitals.

Calculates single particle wave function terms which are independent of the quantum numbers.

If a term in the single particle functions are independent of the quantum number, this function can be overridden to calculate them beforehand (for each particle), and rather extract the value instead of recalculating.

Parameters

i	Particle number.

Reimplemented in AlphaHarmonicOscillator, hydrogenicOrbitals, and ExpandedBasis.

Method for validating closed form expressions for derivatives by comparing them to numerical calculations.

Parameters

q_num	The quantum number index.
d	The dimension for which the derivative should be calculated (x,y,z) .

3.27.2.12 void Orbitals::testLaplace (const Walker * walker, int particle, int q_n um) [protected]

Method for validating closed form expressions for laplacians by comparing them to numerical calculations.

Parameters

```
q_num The quantum number index.
```

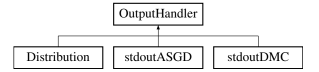
The documentation for this class was generated from the following files:

- · src/Orbitals/Orbitals.h
- src/Orbitals/Orbitals.cpp

3.28 OutputHandler Class Reference

Class for handling output-methods. Designed to avoid rewriting code, as well as avoid if-tests if output is not desired.

Inheritance diagram for OutputHandler:



Public Member Functions

 OutputHandler (std::string filename, std::string path, bool parallel, int node, int n_nodes)

Constructor.

• virtual void dump ()=0

Methods for updating the output.

• virtual void finalize ()

Finalizes the output.

- void set_qmc_ptr (QMC *qmc)
- void set_min_ptr (Minimizer *min)

Protected Member Functions

- void init file ()
- virtual void post_pointer_init ()

Method for initialization requires once the correct QMC/Min pointer type is set.

Protected Attributes

· bool is vmc

Switch used to typecast the QMC object to a VMC object.

· bool is dmc

Switch used to typecast the QMC object to a DMC object.

bool is ASGD

Switch used to typecast the Min object to an ASGD object.

- bool parallel
- int node
- int n nodes
- · bool use_file

If init_file() is called, this flag is set true. Assures correct finalization.

- std::stringstream s
- · std::string filename
- · std::string path
- std::ofstream file
- QMC * qmc
- DMC * dmc
- VMC * vmc
- Minimizer * min
- ASGD * asgd

3.28.1 Detailed Description

Class for handling output-methods. Designed to avoid rewriting code, as well as avoid if-tests if output is not desired.

See also

QMC::output_handler, Minimizer::output_handler

3.28.2 Constructor & Destructor Documentation

3.28.2.1 OutputHandler::OutputHandler (std::string *filename*, std::string *path*, bool *parallel*, int *node*, int *n_nodes*)

Constructor.

Parameters

ĺ	filename	The name of the output file.
	path	The path of the output.

3.28.3 Member Function Documentation

```
3.28.3.1 virtual void OutputHandler::dump() [pure virtual]
```

Methods for updating the output.

Typically retrieves information through the solver pointers (given correct accessibility levels/friend)

Implemented in Distribution, stdoutDMC, and stdoutASGD.

```
3.28.3.2 void OutputHandler::finalize() [virtual]
```

Finalizes the output.

Closes file if use_file flag is true. Can be overridden if more complex tasks needs to be done, such as calculating histograms etc.

See also

```
Distribution::finalize()
```

Reimplemented in Distribution.

```
3.28.3.3 void OutputHandler::init_file() [protected]
```

Opens a file with filename at path supplied in constructor. Subclass implementations can call this function. Superclass does not.

Method for initialization requires once the correct QMC/Min pointer type is set.

Defaults to nothing.

Reimplemented in stdoutASGD.

```
3.28.3.5 void OutputHandler::set_min_ptr ( Minimizer * min )
```

Sets the Min pointer and typecasts it according to the minimizer flags.

```
3.28.3.6 void OutputHandler::set_qmc_ptr ( QMC * qmc )
```

Sets the QMC pointer and typecasts it according to the solver flags.

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

- src/OutputHandler/OutputHandler.h
- src/OutputHandler/OutputHandler.cpp

3.29 OutputParams Struct Reference

Struct used to initialize output parameters.

Public Attributes

· bool dist out

If true, distributions are calculated for VMC/DMC.

• bool dmc_out

If true, DMC outputs data to file each cycle.

bool ASGD_out

If true, ASGD outputs data to file each cycle.

3.29.1 Detailed Description

Struct used to initialize output parameters.

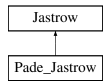
The documentation for this struct was generated from the following file:

· src/QMCheaders.h

3.30 Pade_Jastrow Class Reference

The Pade Jastrow factor with a single variational parameter.

Inheritance diagram for Pade_Jastrow:



Public Member Functions

- Pade_Jastrow (GeneralParams &, VariationalParams &)
- void initialize ()
- void get_grad (Walker *walker) const
- void get_grad (const Walker *walker_pre, Walker *walker_post, int i) const

Updates the gradient for a new particle move.

• void get_dJ_matrix (Walker *walker, int i) const

Updates the summation factors of Jastrow factors closed form expressions.

 double get_j_ratio (const Walker *walker_new, const Walker *walker_old, int i) const Calculates the ratio of the Jastrow factor needed by metropolis.

- double get_val (const Walker *walker) const
- double get_lapl_sum (Walker *walker) const

Method for calculating the Laplacian.

Protected Member Functions

- double get_variational_derivative (const Walker *walker, int n)
 Calculates the derivative of the Jastrow factor with respect to a variational parameter.
- void set parameter (double param, int n)

Sets variational parameters.

• double get_parameter (int n)

Returns variational parameters.

Protected Attributes

· double beta

The variational parameter.

• arma::mat a

The spin-dependent variables taking care of the cusp condition.

3.30.1 Detailed Description

The Pade Jastrow factor with a single variational parameter.

3.30.2 Member Function Documentation

```
3.30.2.1 void Pade_Jastrow::get_dJ_matrix ( Walker * walker, int i ) const [virtual]
```

Updates the summation factors of Jastrow factors closed form expressions.

Used to optimize the calculations as few of these terms change as we move a particle.

Parameters

```
i Particle number.
```

Implements Jastrow.

```
3.30.2.2 void Pade_Jastrow::get_grad ( Walker * walker ) const [virtual]
```

Calculates the entire Cartesian gradient.

Implements Jastrow.

3.30.2.3 void Pade_Jastrow::get_grad (const Walker * walker_pre, Walker * walker_post, int i) const [virtual]

Updates the gradient for a new particle move.

Parameters

ĺ	walker_post	Walker at current time step
	walker_pre	Walker at previous time step
ĺ	i	Particle number.

Implements Jastrow.

3.30.2.4 double Pade_Jastrow::get_j_ratio (const Walker * walker_new, const Walker * walker_old, int i) const [virtual]

Calculates the ratio of the Jastrow factor needed by metropolis.

Parameters

walker_new	Walker at current time step
walker_old	Walker at previous time step
i	The particle number.

Implements Jastrow.

3.30.2.5 double Pade_Jastrow::get_lapl_sum (Walker * walker) const [virtual]

Method for calculating the Laplacian.

Calculates the sum of all particles Laplacians.

Implements Jastrow.

Returns variational parameters.

Parameters

n	The index of the sought variational parameter

Returns

Variational parameter with index [n]

Implements Jastrow.

3.30.2.7 double Pade_Jastrow::get_val(const Walker * walker) const [virtual]

Calculates the value of the Jastrow Factor at the walker's position.

Implements Jastrow.

Calculates the derivative of the Jastrow factor with respect to a variational parameter.

Parameters

n	The index of the variational parameter for which the derivative is to be
	taken
walker	The walker holds the positions etc. needed to evaluate the derivative

Reimplemented from Jastrow.

```
3.30.2.9 void Pade_Jastrow::initialize() [virtual]
```

In case of Pade Jastrow, initializing means seting up the a matrix.

Implements Jastrow.

Sets variational parameters.

Parameters

n	The index of the sought variational parameter
param	The new value of parameter [n]

Implements Jastrow.

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

- · src/Jastrow/Pade Jastrow/Pade Jastrow.h
- src/Jastrow/Pade_Jastrow/Pade_Jastrow.cpp

3.31 ParParams Struct Reference

Struct used to initialize parallelization parameters.

Public Attributes

bool is_master

True for the master node.

bool parallel

True if $n_nodes > 1$.

• int node

The process' rank.

• int n_nodes

The total number of processes.

3.31.1 Detailed Description

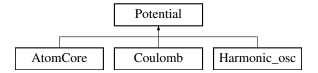
Struct used to initialize parallelization parameters.

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

· src/QMCheaders.h

3.32 Potential Class Reference

Superclass for potentials. Potentials are stores in a vector in the system object. Inheritance diagram for Potential:



Public Member Functions

- Potential (int n_p, int dim)
- virtual double get_pot_E (const Walker *walker) const =0

Method for calculating a walker's potential energy.

• std::string get_name ()

Public Attributes

Sampler pot_sampler

Protected Attributes

- int **n p**
- int dim
- · std::string name

3.32.1 Detailed Description

Superclass for potentials. Potentials are stores in a vector in the system object.

See also

System::potentials, System::get_potential_energy()

3.32.2 Member Function Documentation

3.32.2.1 virtual double Potential::get_pot_E (const Walker * walker) const [pure virtual]

Method for calculating a walker's potential energy.

Method overridden by subclasses.

Parameters

walker The walker for which the potential energy should be calculated.

Returns

The potential energy.

Implemented in Harmonic_osc, AtomCore, and Coulomb.

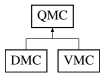
The documentation for this class was generated from the following files:

- src/Potential/Potential.h
- src/Potential/Potential.cpp

3.33 QMC Class Reference

The QMC superclass. Holds implementations of general functions for both VMC and DMC in order to avoid rewriting code and emphasize the similarities.

Inheritance diagram for QMC:



Public Member Functions

QMC (GeneralParams &, int n_c, SystemObjects &, ParParams &, int n_w, int K=1)

Constructor.

virtual void run_method ()=0

Method used for executing the solver.

• void get_QF (Walker *walker) const

Method for calculating the Quantum Force.

void get_gradients (const Walker *walker_pre, Walker *walker_post, int particle)
 const

Method for calculating the gradients after moving a particle.

• void get_gradients (Walker *walker) const

Method for calculating the full gradients.

• void get_laplsum (Walker *walker) const

Method for calculating the Laplacian of all walkers.

• double get_wf_value (const Walker *walker) const

Method for calculating the wave functions value at a given walker's position.

• double calculate_local_energy (const Walker *walker)

Method for calculating the local energy.

void get_accepted_ratio ()

Method for calculating the acceptance ratio.

void add_output (OutputHandler *output_handler)

Method used for loading the output_handler with objects.

void set_error_estimator (ErrorEstimator *error_estimator)

Method for setting the error estimator.

• virtual void output ()=0

Method for standard output.

- System * get_system_ptr () const
- Sampling * get_sampling_ptr () const
- Jastrow * get_jastrow_ptr () const
- Orbitals * get_orbitals_ptr () const

Protected Member Functions

virtual void set trial positions ()=0

Method for setting the trial position of the QMC method's walkers.

void diffuse_walker (Walker *original, Walker *trial)

Method for diffusing a walker one time step.

double get_acceptance_ratio (const Walker *walker_pre, const Walker *walker_post, int particle) const

Method for calculating the acceptance ratio used in the Metropolis test.

• virtual bool move_autherized (double A)=0

Method for deciding whether or not to accept a move.

bool metropolis_test (double A)

Method for performing the metropolis test after when diffusing a walker.

 void update_walker (Walker *walker_pre, const Walker *walker_post, int particle) const

Method for updating the walker after an accepted step.

void reset_walker (const Walker *walker_pre, Walker *walker_post, int particle)
 const

Method for reseting the walker after a rejected step.

• void copy_walker (const Walker *parent, Walker *child) const

Method for (hard) copying a walker object.

• void calculate_energy_necessities (Walker *walker) const

Method for calculating the necessary quantities needed in order to calculate the local energy.

double get KE (const Walker *walker)

Method for calculating the kinetic energy of a walker.

- void **update_subsamples** (double weight=1.0)
- void push_subsamples ()
- void dump_subsamples (bool mean_of_means=false)
- virtual void save_distribution ()=0

Method for storing positional data.

• virtual void node_comm ()=0

Method for performing node communication.

- void dump output ()
- void finalize_output ()
- void estimate_error () const
- void set_spin_state (int particle) const
- void test_ratios (const Walker *walker_pre, const Walker *walker_post, int particle, double R_qmc) const

Method used for testing the optimized ratio calculation.

void test_gradients (Walker *walker)

Method for testing the optimized gradients calculation.

Protected Attributes

• STDOUT * std out

Output object. Wraps and replaces std::cout.

- std::stringstream s
- int output_tresh
- int n_w_size

The total number of allocated walkers.

• std::string runpath

The directory which the simulation is set to run.

std::string dist_path

The path where the distribution are saved.

· arma::mat dist

Matrix holding positional data for the distribution.

· int last inserted

Index of last inserted positional data.

· int dist tresh

Amount of cycles to skip in between storing position data.

- bool is_master
- bool parallel
- int node
- int n_nodes
- int n_c

The number of Monte-Carlo cycles.

· int thermalization

The number of thermalization steps.

- int cycle
- int n_w

The number of walkers.

- int **n_p**
- int **n2**
- int dim
- · unsigned long int accepted

Number of accepted moves.

• unsigned long int total_samples

Total number of moves.

• double local_E

The last calculated local energy.

Walker * trial_walker

The trial walker used to test a move.

Walker ** original_walkers

A list of n_w walkers used in DMC.

Jastrow * jastrow

The Jastrow object.

Sampling * sampling

The Sampling object.

System * system

The system object.

• ErrorEstimator * error estimator

The error estimator.

- Sampler kinetic_sampler
- std::vector< OutputHandler * > output_handler

Can hold stdoutDMC (in case of DMC), Distribution, both or none.

Friends

· class Distribution

3.33.1 Detailed Description

The QMC superclass. Holds implementations of general functions for both VMC and DMC in order to avoid rewriting code and emphasize the similarities.

3.33.2 Constructor & Destructor Documentation

```
3.33.2.1 QMC::QMC ( GeneralParams & gP, int n_{-}c, SystemObjects & sO, ParParams & pp, int n_{-}w, int K = 1 )
```

Constructor.

K K times n_w walkers are initialized. K != 0 only sensible in DMC.

3.33.3 Member Function Documentation

```
3.33.3.1 void QMC::calculate_energy_necessities ( Walker * walker ) const [protected]
```

Method for calculating the necessary quantities needed in order to calculate the local energy.

See also

Sampling::calculate_energy_necessities()

3.33.3.2 double QMC::calculate_local_energy (const Walker * walker) [inline]

Method for calculating the local energy.

See also

```
get_KE(), System::get_potential_energy()
```

```
3.33.3.3 void QMC::copy_walker ( const Walker * parent, Walker * child ) const [protected]
```

Method for (hard) copying a walker object.

Parameters

```
parent,child The parent is copied to the child.
```

```
3.33.3.4 void QMC::diffuse_walker ( Walker * original, Walker * trial ) [protected]
```

Method for diffusing a walker one time step.

The trial walker must equal the original walker in input. The original walker is updated on output.

```
3.33.3.5 void QMC::dump_output( ) [protected]
```

Iterates over the output objects in the output_handler vector. No if-tests.

```
3.33.3.6 void QMC::estimate_error() const [protected]
```

 ${\bf Estimates\ and\ finalizes\ the\ Error Estimator\ object\ initialized\ in\ the\ error_estimator\ vector.}$

```
3.33.3.7 void QMC::finalize_output() [protected]
```

Calls the finalize function for the object in the output_handler vector.

```
3.33.3.8 void QMC::get_gradients ( const Walker * walker_pre, Walker * walker_post, int particle ) const
```

Method for calculating the gradients after moving a particle.

See also

```
Jastrow::get_grad(), System::get_spatial_grad()
```

```
3.33.3.9 void QMC::get_gradients ( Walker * walker ) const
```

Method for calculating the full gradients.

See also

```
Jastrow::get_grad(), System::get_spatial_grad()
```

```
3.33.3.10 void QMC::get_laplsum ( Walker * walker ) const [inline]
```

Method for calculating the Laplacian of all walkers.

See also

```
System::get_spatial_lapl_sum(), Jastrow::get_lapl_sum()
```

```
3.33.3.11 double QMC::get_wf_value ( const Walker * walker ) const [inline]
```

Method for calculating the wave functions value at a given walker's position.

See also

```
System::get_spatial_wf(), Jastrow::get_val()
```

```
3.33.3.12 bool QMC::metropolis_test(double A) [protected]
```

Method for performing the metropolis test after when diffusing a walker.

Parameters

```
A The acceptance ratio calulated by get_acceptance_ratio().
```

```
3.33.3.13 virtual bool QMC::move_autherized ( double A ) [protected, pure virtual]
```

Method for deciding whether or not to accept a move.

Wraps the metropolis sampling with possibilities of overriding.

See also

```
System::allow_transition()
```

Implemented in DMC, and VMC.

```
3.33.3.14 void QMC::reset_walker( const Walker * walker_pre, Walker * walker_post, int particle) const [protected]
```

Method for reseting the walker after a rejected step.

Given a particle number, the method only resets the changed values.

Parameters

walker_post	Walker at current time step
walker_pre	Walker at previous time step

```
3.33.3.15 virtual void QMC::save_distribution() [protected, pure virtual]
```

Method for storing positional data.

Stored in the dist matrix. Used by OutputHandler::Distribution.

See also

```
Distribution::dump(), VMC::save_distribution(), DMC::save_distribution()
```

Implemented in DMC, and VMC.

```
3.33.3.16 void QMC::set_spin_state(int particle) const [protected]
```

Since the spatial wave function is split, certain values are unchanged if the moved particle has opposite spin. Assuming a two-level system, the first half of the particles are assumed to have one spin value, and the second half the other.

This method sets the start and end position of the block that needs to be changed.

See also

```
System::start, System::end
```

```
3.33.3.17 virtual void QMC::set_trial_positions() [protected, pure virtual]
```

Method for setting the trial position of the QMC method's walkers.

See also

```
Sampling::set_trial_pos()
```

Implemented in DMC, and VMC.

```
3.33.3.18 void QMC::test_gradients ( Walker * walker ) [protected]
```

Method for testing the optimized gradients calculation.

Compares with finite difference calculation.

```
3.33.3.19 void QMC::test_ratios ( const Walker * walker_pre, const Walker * walker_post, int particle, double R_qmc ) const [protected]
```

Method used for testing the optimized ratio calculation.

Compares to brute force computation of the wave function values. R_qmc The optimized trial wave function ratio (spatial and Jastrow).

```
3.33.3.20 void QMC::update_walker ( Walker * walker_pre, const Walker * walker_post, int particle ) const [protected]
```

Method for updating the walker after an accepted step.

Given a particle number, the method only updates the changed values.

Parameters

walker_post	Walker at current time step
walker_pre	Walker at previous time step

3.33.4 Member Data Documentation

```
3.33.4.1 int QMC::cycle [protected]
```

The current Monte-Carlo cycle.

```
3.33.4.2 int QMC::n_w [protected]
```

The number of walkers.

VMC stores this many cycles in case of DMC

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

- src/QMC/QMC.h
- src/QMC/QMC.cpp

3.34 Sampler Class Reference

Public Member Functions

· void queue_value (const double &value)

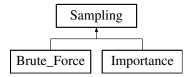
- · void update mean (const double &weight)
- · void push_mean ()
- const double extract_mean ()
- const double extract mean of means ()

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

· src/Sampler/Sampler.h

3.35 Sampling Class Reference

Inheritance diagram for Sampling:



Public Member Functions

- Sampling (int n_p, int dim)
- void update_pos (const Walker *walker_pre, Walker *walker_post, int particle)
 const

Method for updating the position of a walker's particle.

virtual void update_necessities (const Walker *walker_pre, Walker *walker_post, int particle) const =0

Method for updating the sampling specific necessary values.

- virtual void update_walker (Walker *walker_pre, const Walker *walker_post, int particle) const =0
- virtual void reset_walker (const Walker *walker_pre, Walker *walker_post, int particle) const =0
- void set_trial_pos (Walker *walker)

Method for setting the trial position for a given walker.

void set_trial_states (Walker *walker)

Method for setting up the single particle orbitals and it's derivatives for a given walker.

virtual void get_necessities (Walker *walker)=0

Method for calculating the sampling specific necessary values.

virtual void calculate_energy_necessities (Walker *walker) const =0

Method for calculating the sampling specific necessary values in order to compute the local energy.

virtual void copy_walker (const Walker *parent, Walker *child) const =0

Method for copying the sampling specific parts of a walker.

virtual double get_g_ratio (const Walker *walker_post, const Walker *walker_pre) const

Method for calculating the diffusion Green's function ratios.

- double get_branching_Gfunc (double E_x, double E_y, double E_T) const Calculates the Branching Green's function ratio needed by DMC.
- double get_spatialjast_ratio (const Walker *walker_post, const Walker *walker_pre, int particle) const
- void set_qmc_ptr (QMC *qmc)
- · void set_dt (double dt)
- double get_dt () const
- double get_std () const
- double call_RNG ()

Calls a uniform random number generator.

• void set_spin_state (int start, int end)

Protected Attributes

- int n p
- int **n2**
- int dim
- int start
- · int end
- Diffusion * diffusion

The Diffusion object.

• QMC * qmc

The QMC main solver object. Needed to access e.g. the system object.

3.35.1 Member Function Documentation

```
3.35.1.1 double Sampling::call_RNG() [inline]
```

Calls a uniform random number generator.

Returns a random uniform number on [0,1).

```
3.35.1.2 virtual void Sampling::copy_walker ( const Walker * parent, Walker * child ) const [pure virtual]
```

Method for copying the sampling specific parts of a walker.

See also

```
QMC::copy_walker()
```

Implemented in Importance, and Brute Force.

```
3.35.1.3 double Sampling::get_branching_Gfunc ( double E_-x, double E_-y, double E_-T ) const [inline]
```

Calculates the Branching Green's function ratio needed by DMC.

Parameters

E	_ <i>X</i>	Energy at current time step
E	<u>_</u> y	Energy at previous time step

Returns

The Branching Green's function ratio

```
3.35.1.4 virtual double Sampling::get_g_ratio ( const Walker * walker_post, const Walker * walker_pre ) const [inline, virtual]
```

Method for calculating the diffusion Green's function ratios.

See the Diffusion class for documentation.

```
3.35.1.5 virtual void Sampling::get_necessities ( Walker * walker ) [pure
     virtual]
```

Method for calculating the sampling specific necessary values.

Called after a trial position is set.

Implemented in Importance, and Brute_Force.

```
3.35.1.6 virtual void Sampling::reset_walker( const Walker * walker_pre, Walker * walker_post, int particle) const [pure virtual]
```

See also

```
QMC::reset_walker()
```

Implemented in Brute Force, and Importance.

```
3.35.1.7 void Sampling::set_spin_state (int start, int end) [inline]
```

See also

```
QMC::set_spin_state()
```

3.35.1.8 void Sampling::update_pos (const Walker * walker_pre, Walker * walker_post, int particle) const

Method for updating the position of a walker's particle.

Sets a new position according to the diffusion rules, and calls all the functions necessary to get all the values updates, e.g. System::calc_for_new_pos()

3.35.1.9 virtual void Sampling::update_walker(Walker * walker_pre, const Walker * walker_post, int particle) const [pure virtual]

See also

```
QMC::update_walker()
```

Implemented in Importance, and Brute_Force.

```
3.35.2 Member Data Documentation
```

```
3.35.2.1 Diffusion* Sampling::diffusion [protected]
```

The Diffusion object.

See also

Diffusion

```
3.35.2.2 int Sampling::end [protected]
```

See also

System::end

```
3.35.2.3 int Sampling::start [protected]
```

See also

System::start

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

- src/Sampling/Sampling.h
- src/Sampling/Sampling.cpp

3.36 Simple Class Reference

Simple Isotropic diffusion model.

Inheritance diagram for Simple:



Public Member Functions

- **Simple** (int n_p, int dim, double timestep, seed_type random_seed, double D=0.-5)
- double get_new_pos (const Walker *walker, int i, int j)

Virtual function returning the new position.

double get_g_ratio (const Walker *walker_post, const Walker *walker_pre) const

Calculates the Diffusion Green's function ratio needed by metropolis.

3.36.1 Detailed Description

Simple Isotropic diffusion model.

3.36.2 Member Function Documentation

```
3.36.2.1 double Simple::get_g_ratio ( const Walker * walker_post, const Walker * walker_pre ) const [inline, virtual]
```

Calculates the Diffusion Green's function ratio needed by metropolis.

Parameters

[walker post	Walker at current time step.
İ	walker_pre	Walker at previous time step.

Returns

The Diffusion Green's function ratio.

Implements Diffusion.

Virtual function returning the new position.

Returns the simple diffusion step if not overridden.

Parameters

i	Particle number.
j	dimension (x,y,z).

Returns

The new position (relative to the old).

Reimplemented from Diffusion.

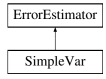
The documentation for this class was generated from the following files:

- src/Diffusion/Simple/Simple.h
- src/Diffusion/Simple/Simple.cpp

3.37 SimpleVar Class Reference

Calculates the simple variance of the sampled values.

Inheritance diagram for SimpleVar:



Public Member Functions

- SimpleVar (ParParams &)
- double estimate_error ()

Estimates the error based on the subclass implementation.

- void update_data (double val)
- void normalize ()

Protected Attributes

· double f

sum variable used to calculate the mean

• double f2

sum variable used to calulate the mean of squares.

3.37.1 Detailed Description

Calculates the simple variance of the sampled values.

3.37.2 Member Function Documentation

3.37.2.1 void SimpleVar::update_data (double *val* **)** [virtual]

Overrides the default described in the superclass. Does not store values in memory, but rather use sum variables.

Reimplemented from ErrorEstimator.

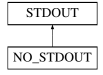
The documentation for this class was generated from the following files:

- src/ErrorEstimator/SimpleVar/SimpleVar.h
- src/ErrorEstimator/SimpleVar/SimpleVar.cpp

3.38 STDOUT Class Reference

Class for handling standard output. Only the master node has this object.

Inheritance diagram for STDOUT:



Public Member Functions

virtual void cout (std::stringstream &a)

3.38.1 Detailed Description

Class for handling standard output. Only the master node has this object.

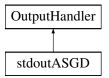
The documentation for this class was generated from the following file:

· src/QMCheaders.h

3.39 stdoutASGD Class Reference

Class for handling the output of ASGD. Ouputs values such as the variational gradients, step length, variational parameters, etc.

Inheritance diagram for stdoutASGD:



Public Member Functions

- stdoutASGD (std::string path, std::string filename="ASGD_out")
- void dump ()

Methods for updating the output.

void post_pointer_init ()

3.39.1 Detailed Description

Class for handling the output of ASGD. Ouputs values such as the variational gradients, step length, variational parameters, etc.

3.39.2 Member Function Documentation

```
3.39.2.1 void stdoutASGD::dump() [virtual]
```

Methods for updating the output.

Typically retrieves information through the solver pointers (given correct accessibility levels/friend)

Implements OutputHandler.

```
3.39.2.2 void stdoutASGD::post_pointer_init( ) [inline, virtual]
```

Initializes the correct size of the variational gradient once the min pointer has been cast to ASGD.

Reimplemented from OutputHandler.

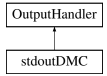
The documentation for this class was generated from the following files:

- src/OutputHandler/stdoutASGD/stdoutASGD.h
- src/OutputHandler/stdoutASGD/stdoutASGD.cpp

3.40 stdoutDMC Class Reference

Class for handling the output of DMC. Outputs values such as the trial energy, dmc energy, number of walkers, etc.

Inheritance diagram for stdoutDMC:



Public Member Functions

- stdoutDMC (std::string path, std::string filename="DMC_out")
- void dump ()

Methods for updating the output.

Protected Attributes

• int n

Number of times the dump() method has been called.

double sumE

Sum of the DMC energy used to calculate the trailing average.

double sumN

Sum of the number of walkers used to calculate the trailing average.

3.40.1 Detailed Description

Class for handling the output of DMC. Outputs values such as the trial energy, dmc energy, number of walkers, etc.

3.40.2 Member Function Documentation

```
3.40.2.1 void stdoutDMC::dump() [virtual]
```

Methods for updating the output.

Typically retrieves information through the solver pointers (given correct accessibility levels/friend)

Implements OutputHandler.

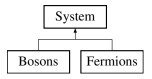
The documentation for this class was generated from the following files:

- src/OutputHandler/stdoutDMC/stdoutDMC.h
- src/OutputHandler/stdoutDMC/stdoutDMC.cpp

3.41 System Class Reference

The system class separating Fermions and Bosons. Designed to generalize the solver in terms of particle species.

Inheritance diagram for System:



Public Member Functions

- System (int n p, int dim, Orbitals *orbital)
- virtual void initialize (Walker *walker)=0

Initializes the system before the main solver loop starts.

void add potential (Potential *pot)

Method for adding a potential to the system.

• double get_potential_energy (const Walker *walker)

Method for calculating the total potential energy.

- virtual void update_walker (Walker *walker_pre, const Walker *walker_post, int particle) const =0
- virtual void reset_walker (const Walker *walker_pre, Walker *walker_post, int particle) const =0
- virtual void calc_for_newpos (const Walker *walker_old, Walker *walker_new, int particle)=0

Method for calculating the necessary values needed by the walker after a new step is made.

virtual double get_spatial_ratio (const Walker *walker_pre, const Walker *walker_post, int particle)=0

Method for calculating the spatial wave function ratios between two subsequent time steps.

• virtual double get_spatial_wf (const Walker *walker)=0

Method for calculating the spatial wave function's value at a given walkers position.

virtual void get_spatial_grad (Walker *walker, int particle) const =0

Method for calculating the changed part of the spatial gradient.

virtual void get_spatial_grad_full (Walker *walker) const =0

Method for calculating the full spatial gradient.

virtual double get spatial lapl sum (const Walker *walker) const =0

Method for calculating the sum of all Laplacians for a given walker.

• virtual void copy_walker (const Walker *parent, Walker *child) const =0

Method for copying the system specific parts of a walker.

• virtual bool allow_transition ()=0

Method allowing the system to override the Metropolis test.

- void update_potential_samples (double weight=1.0)
- void push_potential_samples ()
- std::string dump_samples (bool mean_of_means=false)
- Orbitals * get_orbital_ptr ()
- void set_spin_state (int start, int end)

Protected Attributes

- int **n_p**
- int **n2**
- int dim
- · int start

The start point of separable calculations.

• int end

The end point of separable calculations.

std::vector< Potential * > potentials

A vector of potentials.

Orbitals * orbital

The single particle wave function object.

3.41.1 Detailed Description

The system class separating Fermions and Bosons. Designed to generalize the solver in terms of particle species.

3.41.2 Member Function Documentation

3.41.2.1 virtual void System::calc_for_newpos (const Walker * walker_old, Walker * walker_new, int particle) [pure virtual]

Method for calculating the necessary values needed by the walker after a new step is made.

Given a particle number, the method does not recompute unchanged values.

Parameters

walker_old	Walker at current time step.	
walker_new	Walker at previous time step.	

Implemented in Fermions, and Bosons.

```
3.41.2.2 virtual void System::copy_walker ( const Walker * parent, Walker * child ) const [pure virtual]
```

Method for copying the system specific parts of a walker.

See also

```
QMC::copy_walker()
```

Implemented in Fermions, and Bosons.

```
3.41.2.3 double System::get_potential_energy ( const Walker * walker )
```

Method for calculating the total potential energy.

Iterates over all objects in the potentials vector and accumulates their potential energies for the given walker.

```
3.41.2.4 virtual void System::get_spatial_grad ( Walker * walker, int particle ) const [pure virtual]
```

Method for calculating the changed part of the spatial gradient.

Depending on which particle we moved, one of the spatial wave function parts (it is split) will be unchanged.

Implemented in Fermions, and Bosons.

```
3.41.2.5 virtual void System::initialize ( Walker * walker ) [pure virtual]
```

Initializes the system before the main solver loop starts.

Called by the Sampling class when trial positions are set.

Implemented in Fermions, and Bosons.

```
3.41.2.6 virtual void System::reset_walker ( const Walker * walker_pre, Walker * walker_post, int particle ) const [pure virtual]
```

See also

```
QMC::reset_walker()
```

Implemented in Fermions, and Bosons.

```
3.41.2.7 void System::set_spin_state(int start, int end) [inline]
```

See also

```
QMC::set_spin_state()
```

3.41.2.8 virtual void System::update_walker (Walker * walker_pre, const Walker * walker_post, int particle) const [pure virtual]

See also

```
QMC::update_walker()
```

Implemented in Fermions, and Bosons.

3.41.3 Member Data Documentation

```
3.41.3.1 int System::end [protected]
```

The end point of separable calculations.

Either N/2 or N. Since the spatial wave function is split, particles with spin not equal that of the moved particle is unchanged and does not need to be recalculated.

```
3.41.3.2 int System::start [protected]
```

The start point of separable calculations.

Either 0 or N/2. Since the spatial wave function is split, particles with spin not equal that of the moved particle is unchanged and does not need to be recalculated.

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

- · src/System/System.h
- src/System/System.cpp

3.42 SystemObjects Struct Reference

Struct used to initialize system objects.

Public Attributes

- Orbitals * SP_basis
- Potential * onebody_pot
- System * SYSTEM
- Sampling * sample_method
- Jastrow * jastrow

3.42.1 Detailed Description

Struct used to initialize system objects.

The memory addresses allocated here will not change throughout the run.

See also

Orbitals, Potential, System, Sampling, Jastrow

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

· src/QMCheaders.h

3.43 VariationalParams Struct Reference

Struct used to initialize the varational parameters.

Public Attributes

· double alpha

The spatial variational parameter.

• double beta

The Jastrow variational parameter.

3.43.1 Detailed Description

Struct used to initialize the varational parameters.

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

• src/QMCheaders.h

3.44 VMC Class Reference

Implementation of the Variational Monte-Carlo Method. Very little needs to be added when the QMC superclass holds all the general functionality.

Inheritance diagram for VMC:



Public Member Functions

 VMC (GeneralParams &, VMCparams &, SystemObjects &, ParParams &, int n-_w, bool dist_out)

Constructor.

- void set_e (double E)
- double get_energy () const
- void run_method ()

Method used for executing the solver.

void output ()

Method for standard output.

Protected Member Functions

- void set_trial_positions ()
- void store_walkers ()

Method for storing walkers for DMC.

• void save_distribution ()

Method for storing positional data for the Distribtuon.

- bool move_autherized (double A)
- void scale_values ()
- void node_comm ()

Method for performing node communication.

Protected Attributes

· int pop_tresh

The amount of cycles between storing walkers for DMC.

· int offset

The amount of cycles before starting to store walkers for DMC.

· int last walker

Count variable for the last walker stores for DMC.

• double vmc_E

The VMC energy.

• Walker * original_walker

The VMC walker.

Friends

- class DMC
- · class Minimizer
- · class ASGD
- class BlockingData

3.44.1 Detailed Description

Implementation of the Variational Monte-Carlo Method. Very little needs to be added when the QMC superclass holds all the general functionality.

3.44.2 Member Function Documentation

In VMC, only the metropolis test is performed.

Implements QMC.

```
3.44.2.2 void VMC::save_distribution() [protected, virtual]
```

Method for storing positional data for the Distribtuon.

Stores the position data of the single VMC walker every dist_tresh cycle after thermalization.

Implements QMC.

```
3.44.2.3 void VMC::set_trial_positions() [protected, virtual]
```

Sets the trial position for the single walker.

Implements QMC.

```
3.44.2.4 void VMC::store_walkers() [protected]
```

Method for storing walkers for DMC.

Stores the single VMC walker every pop_thresh cycle after offset cycles.

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

- src/QMC/VMC/VMC.h
- src/QMC/VMC/VMC.cpp

3.45 VMCparams Struct Reference

Struct used to initialize VMC parameters.

Public Attributes

• int n c

The number of cycles.

double dt

The time step.

3.45.1 Detailed Description

Struct used to initialize VMC parameters.

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

· src/QMCheaders.h

3.46 Walker Class Reference

Class representing a Random Walker. Holds position data, alive state, etc. Designed to lighten function arguments, and ease implementation of QMC methods involving multiple walkers. Alot of values are stored to avoid calculating the same value twice.

Public Member Functions

```
• Walker (int n p, int dim, bool alive=true)
```

Constructor.

void calc_r_i2 (int i)

Method for calculating the radius squared for one particle.

• void calc_r_i2 ()

Method for calculating the radius squared for all particles.

• double calc_r_rel (int i, int j) const

Method for calculating the relative distance between two particles.

- void make_rel_matrix ()
- void send_soul (int dest)
- void recv_soul (int root)
- double get_r_i2 (int i) const

Method for fetching the squared radius of a particle.

• double get_r_i (int i) const

Method for calculating the radius of a particle.

- void kill ()
- bool is_dead ()
- bool is_alive ()
- void ressurect ()
- void set E (double E)
- double get_E () const
- void print (std::string header="----") const

Prints out all the walkers information.

Public Attributes

double spatial_ratio

The ratio of the spatial wave function (stored in the newest walker).

• double lapl_sum

The sum of the Laplacians of all particles.

double E

The energy of the given configuration (stored to speed up DMC).

· arma::mat r

The positions of all particles.

arma::mat r_rel

The relative positions of all particles.

arma::mat qforce

The Quantum Force for all particles.

· arma::mat spatial_grad

The gradient of the Spatial Wave function for all particles.

· arma::mat jast_grad

The gradient of the Jastrow Factor for all particles.

· arma::mat inv

The inverse of the Slater matrix (given fermion system)

· arma::mat phi

The single particle wave functions for all particles and quantum numbers.

arma::field< arma::mat > dell phi

The derivatives of the single particle wave functions for all particles and quantum numbers

· arma::cube dJ

Cube used for storing sum terms for the Jastrow Factor's closed form expressions.

arma::rowvec r2

The radius squared for all particles.

Protected Attributes

- int **n p**
- int **n2**
- int dim
- · bool is_murdered

If true, the walker will be deleted and removed (DMC only).

3.46.1 Detailed Description

Class representing a Random Walker. Holds position data, alive state, etc. Designed to lighten function arguments, and ease implementation of QMC methods involving multiple walkers. Alot of values are stored to avoid calculating the same value twice.

3.46.2 Constructor & Destructor Documentation

3.46.2.1 Walker::Walker (int n_p, int dim, bool alive = true)

Constructor.

Parameters

alive If false, the walker is initialized dead.

3.46.3 Member Function Documentation

3.46.3.1 void Walker::calc_r_i2 (int *i*)

Method for calculating the radius squared for one particle.

Parameters

i The particle number.

3.46.3.2 double Walker::calc_r_rel (int i, int j) const

Method for calculating the relative distance between two particles.

Parameters

i,j The particle numbers.

3.46.3.3 double Walker::get_r_i (int i) const [inline]

Method for calculating the radius of a particle.

Parameters

i Particle number.

3.46.3.4 double Walker::get_r_i2 (int *i* **) const** [inline]

Method for fetching the squared radius of a particle.

Used in order to avoid calculating the same radius twice.

Parameters

i Particle number.

```
3.46.3.5 void Walker::kill() [inline]
```

Flags the walker for destruction.

See also

```
DMC::bury_the_dead()
```

```
3.46.3.6 void Walker::make_rel_matrix()
```

Creates the relative position matrix.

```
3.46.3.7 void Walker::print ( std::string header = "----" ) const
```

Prints out all the walkers information.

Extremely handy for debugging.

Parameters

header A header for the printout in order to distinguish several printouts easily.

```
3.46.3.8 void Walker::recv_soul ( int root )
```

Receives a walker from a different node.

Parameters

root The rank of the node from which the walker was sent.

```
3.46.3.9 void Walker::ressurect() [inline]
```

Sets the destruction flag to false.

```
3.46.3.10 void Walker::send_soul (int dest)
```

Send a walker to a different node.

Parameters

```
dest The receiving node's rank.
```

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

- src/Walker/Walker.h
- src/Walker/Walker.cpp