oepdev 0.0.0

Generated by Doxygen 1.8.6

Tue Jan 23 2018 18:56:13

# **Contents**

1	Mair	n Page																				1
2	Lice	nse																				3
3	OEF	Desigr	1.																			5
	3.1	OEP C	classes .																 		 	5
		3.1.1	Structure	of po	ssible	OEF	²-base	ed ex	pres	sior	ns a	nd 1	thei	r un	ifica	atio	n .		 		 	5
4	List	of One-	Electron	Poten	tals																	7
	4.1	Electro	static Ene	ergy O	EP's														 		 	7
	4.2	Pauli F	Repulsion (	OEP's															 		 	7
		4.2.1	First-ord	er con	tributi	on in	overl	lap m	atrix	exp	ans	ion							 		 	7
		4.2.2	Second-	order (	contril	bution	ı in o	verla	p ma	trix	exp	ans	ion.						 		 	8
	4.3	Excito	nic Energy	Trans	fer Ol	EP's													 		 	8
		4.3.1	ET contr	ibution	18														 		 	8
		4.3.2	HT contr	ibutior	ns														 		 	8
		4.3.3	CT contr	ibutior	18														 		 	8
	4.4	Full H	- Interaction	on OEI	P's .														 		 	8
5	Nam	nespace	Index																			9
	5.1	Names	space List																 		 	9
6	Hier	archica	l Index																			11
	6.1	Class	Hierarchy																 		 	11
7	Clas	s Index																				13
	7.1	Class	List																 		 	13
8	Nam	nespace	Docume	ntatio	n																	15
	8.1	oepde	v Namesp	ace Re	eferen	ice .													 		 	15
		8.1.1	Detailed	Descr	ription	١													 		 	16
		8.1.2	Function	Docu	menta	ation													 		 	17
			8.1.2.1	crea	ıte_su	perfu	nction	nal .											 		 	17
			8.1.2.2	extra	act_m	onom	ner .											 	 		 	17

iv CONTENTS

			8.1.2.3 solve_scf	17
	8.2	psi Nar	nespace Reference	17
		8.2.1	Detailed Description	18
•	Clas	a Daaw		19
9				
	9.1	•		19
		9.1.1	·	19
		9.1.2		20
			<u> </u>	20
			Ç	20
		9.1.3	Member Function Documentation	20
			9.1.3.1 index	20
	9.2	oepdev	::AllAOShellCombinationsIterator Class Reference	20
		9.2.1	Detailed Description	21
		9.2.2	Constructor & Destructor Documentation	21
			9.2.2.1 AllAOShellCombinationsIterator	21
			9.2.2.2 AllAOShellCombinationsIterator	21
		9.2.3	Member Function Documentation	22
			9.2.3.1 compute_shell	22
	9.3	oepdev	::CPHF Class Reference	22
		9.3.1	Detailed Description	23
		9.3.2	Constructor & Destructor Documentation	23
			9.3.2.1 CPHF	23
	9.4	oepdev	::CubePoints3DIterator Class Reference	24
		9.4.1	Detailed Description	24
	9.5	oepdev	::CubePointsCollection3D Class Reference	25
		9.5.1		25
	9.6	oepdev	::DIISManager Class Reference	25
		9.6.1	Detailed Description	26
		9.6.2	Constructor & Destructor Documentation	26
			9.6.2.1 DIISManager	26
		9.6.3		26
				26
				26
			•	26
	9.7	oendev		- 26
	···	9.7.1		-0 27
		9.7.2		-, 27
		J		- ' 27
	9.8	nende		-, 27
	0.0	Johna	in Liout de la	-1

CONTENTS

	9.8.1	Detailed I	Description	. 28
	9.8.2	Member I	Function Documentation	. 28
		9.8.2.1	compute_3D	. 28
9.9	oepdev	::Electrost	taticEnergySolver Class Reference	. 28
	9.9.1	Member I	Function Documentation	. 29
		9.9.1.1	compute_benchmark	. 29
		9.9.1.2	compute_oep_based	. 29
9.10	oepdev	::Electrost	taticPotential3D Class Reference	. 29
	9.10.1	Detailed I	Description	. 30
9.11	oepdev	:::ESPSolv	ver Class Reference	. 30
	9.11.1	Detailed I	Description	. 31
	9.11.2	Construc	stor & Destructor Documentation	. 31
		9.11.2.1	ESPSolver	. 31
		9.11.2.2	ESPSolver	. 31
9.12	oepdev	::OEPDev	/Solver Class Reference	. 32
	9.12.1	Detailed I	Description	. 33
	9.12.2	Construc	stor & Destructor Documentation	. 33
		9.12.2.1	OEPDevSolver	. 33
	9.12.3	Member I	Function Documentation	. 33
		9.12.3.1	build	. 33
		9.12.3.2	compute_benchmark	. 33
		9.12.3.3	compute_oep_based	. 33
9.13	oepdev	::OEPoten	ntial Class Reference	. 33
	9.13.1	Detailed I	Description	. 35
	9.13.2	Construc	stor & Destructor Documentation	. 35
		9.13.2.1	OEPotential	. 35
		9.13.2.2	OEPotential	. 35
	9.13.3	Member I	Function Documentation	. 36
		9.13.3.1	build	. 36
		9.13.3.2	build	. 36
		9.13.3.3	compute_3D	. 36
		9.13.3.4	write_cube	. 36
9.14	oepdev	::OEPoten	ntial3D< T > Class Template Reference	. 36
	9.14.1	Detailed I	Description	. 37
	9.14.2	Construc	stor & Destructor Documentation	. 38
		9.14.2.1	OEPotential3D	. 38
		9.14.2.2	OEPotential3D	. 38
9.15	oepdev	::Points3D	Olterator::Point Struct Reference	. 38
9.16	oepdev	::Points3D	Olterator Class Reference	. 39
	9.16.1	Detailed I	Description	. 40

vi CONTENTS

	9.16.2	Constructor & Destructor Documentation	40
		9.16.2.1 Points3DIterator	40
	9.16.3	Member Function Documentation	40
		9.16.3.1 build	40
		9.16.3.2 build	41
		9.16.3.3 build	42
9.17	oepdev	r::PointsCollection3D Class Reference	42
	9.17.1	Detailed Description	43
	9.17.2	Constructor & Destructor Documentation	43
		9.17.2.1 PointsCollection3D	43
	9.17.3	Member Function Documentation	44
		9.17.3.1 build	44
		9.17.3.2 build	44
		9.17.3.3 build	44
9.18	oepdev	::PotentialInt Class Reference	44
	9.18.1	Detailed Description	45
	9.18.2	Constructor & Destructor Documentation	45
		9.18.2.1 PotentialInt	45
		9.18.2.2 PotentialInt	45
		9.18.2.3 PotentialInt	46
	9.18.3	Member Function Documentation	46
		9.18.3.1 set_charge_field	46
9.19	oepdev	::RandomPoints3DIterator Class Reference	46
	9.19.1	Detailed Description	47
9.20	oepdev	::RandomPointsCollection3D Class Reference	48
	9.20.1	Detailed Description	48
9.21	oepdev	::RepulsionEnergyOEPotential Class Reference	48
	9.21.1	Detailed Description	49
	9.21.2	Member Function Documentation	49
		9.21.2.1 compute_3D	49
9.22	oepdev	:::ScalarField3D Class Reference	49
	9.22.1	Detailed Description	51
9.23	oepdev	:::WavefunctionUnion Class Reference	51
		Detailed Description	53
		Constructor & Destructor Documentation	54
		9.23.2.1 WavefunctionUnion	54

Index

55

## **Main Page**

## oep-dev

Generalized One-Electron Potentials: Development Platform.

Contact: Bartosz Błasiak (blasiak.bartosz@gmail.com)

#### Overview

Test various models of the intermolecular interaction that is based on the application of the **One-Electron Potentials** (**OEP's**) technique.

Currently, the interaction between two molecules described by the Hartree-Fock-Roothaan-Hall theory or the configuration interaction with singles theory is considered. In particular, the plugin tests the models of:

- 1. the Pauli exchange-repulsion interaction energy (Project II)
- 2. the Induction interaction energy (Project III)
- 3. the excitation energy transfer couplings (Project I )

against reference solutions (exact or other approximations).

Places to go:

- https://github.com/globulion/oepdev/blob/master/doc/git/doc\_oep\_design.md "OEP Design"
- https://github.com/globulion/oepdev/blob/master/doc/git/doc\_implemented\_models.md "Implemented Models"
- https://github.com/globulion/oepdev/blob/master/doc/git/doc\_programming\_etiquette.md "Programming Etiquette"
- Current Issues

#### References

2 Main Page

## License

Copyright (c) 2018, Bartosz Błasiak

All rights reserved.

Usage, copy or redistribution is allowed only after obtaining written consent of the Repository Administrator.

License

## **OEP Design.**

OEP (One-Electron Potential) is associated with certain quantum one-electron operator  $\hat{v}^A$  that defines the ability of molecule A to interact in a particular way with other molecules.

Technically, OEP can be understood as a **container object** (associated with the molecule in question) that stores the information about the above mentioned quantum operator. Here, it is assumed that similar OEP object is also defined for all other molecules in a molecular aggregate.

In case of interaction between molecules A and B, OEP object of molecule A interacts directly with wavefunction object of the molecule B. Defining a Solver class that handles such interaction Wavefunction class and OEP class the universal design of OEP-based approaches can be established and developed.

**Important:** OEP and Wavefunction classes should not be restricted to Hartree-Fock; in generall any correlated wavefunction and derived OEP's should be allowed to work with each other.

## 3.1 OEP Classes

There are many types of OEP's, but the underlying principle is the same and independent of the type of intermolecular interaction. Therefore, the OEP's should be implemented by using a multi-level class design. In turn, this design depends on the way OEP's enter the mathematical expressions, i.e., on the types of matrix elements of the one-electron effective operator  $\hat{v}^A$ .

#### 3.1.1 Structure of possible OEP-based expressions and their unification

Structure of OEP-based mathematical expressions is listed below:

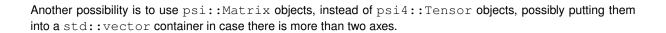
In the above table, I, J and K indices correspond to basis functions or molecular orbitals. Basis functions can be primary or auxiliary OEP-specialized density-fitting. Depending on the type of function and matrix element, there are many subtypes of resulting matrix elements that differ in their dimensionality. Examples are given below:

```
| Matrix Element | DF-based form | ESP-based form | |---|--|
```

In the formulae above, the OEP-part (stored by OEP instances) is shown in blue whereas the Solver-part (to be computed by the Solver) is shown in brown. It is apparent that all OEP-parts have the form of 2nd- or 3rd-rank tensors with different class of axes (molecular orbitals, primary/auxiliary basis, atomic space). Therefore, they can be uniquely defined by a unified *tensor object* (storing double precision numbers) and unified *dimension object* storing the information of the axes classes.

In Psi4, a perfect candidate for the above is psi4::Tensor class declared in psi4/libthce/thce.h. Except from the numeric content its instances also store the information of the dimensions in a form of a vector of psi4::Dimension instances.

6 OEP Design.



## **List of One-Electron Potentals**

Here I provide the list of OEP's that have been already derived within the scope of the OEPDev project.

## 4.1 Electrostatic Energy OEP's

For electrostatic energy calculations, OEP is simply the electrostatic potential due to nuclei and electrons. 3D form:

$$v(\mathbf{r}) = \sum_{x} \frac{Z_{x}}{|\mathbf{r} - \mathbf{r}_{x}|} + \sum_{\mu\nu \in A} P_{\nu\mu} \int d\mathbf{r}' \frac{\phi_{\mu}^{*}(\mathbf{r}')\phi_{\nu}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}$$

Matrix form:

$$v_{ik} = \sum_{x \in A} Z_x V_{ik}^{(x)} + \sum_{\mu \nu \in A} P_{\nu \mu} \left( \mu \nu | ik \right)$$

## 4.2 Pauli Repulsion OEP's

The following potentials are derived for the evaluation of the Pauli repulsion energy based on Murrel's expressions.

### 4.2.1 First-order contribution in overlap matrix expansion.

This contribution is simply the electrostatic potential coming from all nuclei and electron density except\* from electron density from molecular orbital i that interacts with the generalized overlap density between i of molecule A and j of molecule B.

3D forms:

$$v(\mathbf{r})_{S^{-1}}^{A[i]} = -\sum_{x \in A} \frac{Z_x}{|\mathbf{r} - \mathbf{r}_x|} + \sum_{\mu\nu \in A} \left\{ D_{\nu\mu} - C_{\mu i}^* C_{\nu i} \right\} \int d\mathbf{r}' \frac{\phi_\mu^*(\mathbf{r}')\phi_\nu(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}$$

Matrix forms:

$$v_{\xi_i}(S^{-1}) = \sum_{\kappa \in A} C_{i\kappa} \left\{ -\sum_{x \in A} V_{\kappa\xi}^{(x)} + \sum_{\mu\nu \in A} \left\{ D_{\nu\mu} - C_{\mu i}^* C_{\nu i} \right\} (\mu\nu | \xi \kappa) \right\}$$

4.2.2 Second-order contribution in overlap matrix expansion.

To be added here!

## 4.3 Excitonic Energy Transfer OEP's

The following potentials are derived for the evaluation of the short-range EET couplings based on Fujimoto's TDFI-TI method.

#### 4.3.1 ET contributions.

3D forms:

$$\begin{split} &v(\mathbf{r})_{1}^{A[\mu]} = -C_{\mu L}^{*} \sum_{x \in A} \frac{Z_{x}}{|\mathbf{r} - \mathbf{r}_{x}|} + \sum_{v_{K} \in A} \left\{ C_{\mu L}^{*} D_{v_{K}} - \frac{1}{2} C_{vL}^{*} D_{\mu_{K}} \right\} \int d\mathbf{r}' \frac{\phi_{v}^{*}(\mathbf{r}') \phi_{K}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} \\ &v(\mathbf{r})_{2}^{A[\mu]} = C_{\kappa H} \sum_{v_{K} \in A} \left\{ 2C_{vL}^{*} C_{\mu H}^{*} - C_{vH}^{*} C_{\mu L}^{*} \right\} \int d\mathbf{r}' \frac{\phi_{v}^{*}(\mathbf{r}') \phi_{K}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} \\ &v(\mathbf{r})_{3}^{A[\mu]} = v(\mathbf{r})_{1}^{A[\mu]} + v(\mathbf{r})_{1}^{A[\mu]} \end{split}$$

Matrix forms:

$$\begin{split} v_{\mu\xi}(1) &= -C_{\mu L}^* \sum_{\mathbf{x} \in A} V_{\mu\xi}^{\mathbf{x}} + \sum_{\mathbf{v} \kappa \in A} \left\{ C_{\mu L}^* D_{\mathbf{v} \kappa} - \frac{1}{2} C_{\mathbf{v} L}^* D_{\mu \kappa} \right\} (\mathbf{v} \kappa | \mu \xi) \\ v_{\mu\xi}(2) &= C_{\kappa H} \sum_{\mathbf{v} \kappa \in A} \left\{ 2 C_{\mathbf{v} L}^* C_{\mu H}^* - C_{\mathbf{v} H}^* C_{\mu L}^* \right\} (\mathbf{v} \kappa | \mu \xi) \\ v_{\mu\xi}(3) &= v_{\mu\xi}(1) + v_{\mu\xi}(2) \end{split}$$

#### 4.3.2 HT contributions.

Do be derived.

### 4.3.3 CT contributions.

To be derived.

#### 4.4 Full HF Interaction OEP's

The following potentials are derived for the evaluation of the full Hartree-Fock interaction energy based on the OEPDev equations.

# Namespace Index

## 5.1 Namespace List

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

oepdev																			
psi	Oepdev module namespace																	7	??
ры	Psi4 package namespace .																	7	??

10 Namespace Index

## **Hierarchical Index**

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

## 6.1 Class Hierarchy

Wavefunction

oepdev::AllAOIntegralsIterator	??
oepdev::AllAOShellCombinationsIterator	??
oepdev::CPHF	??
CubicScalarGrid	
oepdev::CubePointsCollection3D	??
oepdev::DIISManager	??
enable_shared_from_this	
oepdev::OEPDevSolver	??
oepdev::ElectrostaticEnergySolver	??
oepdev::OEPotential	??
oepdev::EETCouplingOEPotential	??
oepdev::ElectrostaticEnergyOEPotential	??
oepdev::RepulsionEnergyOEPotential	
oepdev::ESPSolver	??
oepdev::Points3DIterator::Point	
oepdev::Points3DIterator	
oepdev::CubePoints3DIterator	
oepdev::RandomPoints3DIterator	
oepdev::PointsCollection3D	
·	
oepdev::CubePointsCollection3D	
oepdev::RandomPointsCollection3D	??
PotentialInt	

12 **Hierarchical Index** 

# **Class Index**

## 7.1 Class List

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

oepdev::AllAOIntegralsIterator	
Loop over all possible ERI within a particular shell	??
oepdev::AllAOShellCombinationsIterator	
Loop over all possible ERI shells	??
oepdev::CPHF	
CPHF solver class	??
oepdev::CubePoints3DIterator	
Iterator over a collection of points in 3D space. g09 Cube-like order	??
oepdev::CubePointsCollection3D	
G09 cube-like ordered collection of points in 3D space	??
oepdev::DIISManager	
DIIS manager	??
oepdev::EETCouplingOEPotential	
Generalized One-Electron Potential for EET coupling calculations	??
oepdev::ElectrostaticEnergyOEPotential	
Generalized One-Electron Potential for Electrostatic Energy calculations	??
oepdev::ElectrostaticEnergySolver	??
oepdev::ElectrostaticPotential3D	
Electrostatic potential of a molecule	??
oepdev::ESPSolver	
Charges from Electrostatic Potential (ESP). A solver-type class	??
oepdev::OEPDevSolver	
Solver of properties of molecular aggregates. Abstract base	??
oepdev::OEPotential	
Generalized One-Electron Potential: Abstract base	??
oepdev::OEPotential3D< T >	
Class template for OEP scalar fields	??
oepdev::Points3Dlterator::Point	??
oepdev::Points3DIterator	
Iterator over a collection of points in 3D space. Abstract base	??
oepdev::PointsCollection3D	
Collection of points in 3D space. Abstract base	??
oepdev::PotentialInt	
Computes potential integrals	??
oepdev::RandomPoints3DIterator	
Iterator over a collection of points in 3D space. Random collection	??
oepdev::RandomPointsCollection3D	_
Collection of random points in 3D space	??

14 Class Index

oepdev::RepulsionEnergyOEPotential	
Generalized One-Electron Potential for Pauli repulsion energy calculations	??
oepdev::ScalarField3D	
Scalar field in 3D space. Abstract base	??
oepdev::WavefunctionUnion	
Union of two Wavefunction objects	??

## **Namespace Documentation**

## 8.1 oepdev Namespace Reference

oepdev module namespace.

#### Classes

· class OEPotential

Generalized One-Electron Potential: Abstract base.

• class ElectrostaticEnergyOEPotential

Generalized One-Electron Potential for Electrostatic Energy calculations.

class RepulsionEnergyOEPotential

Generalized One-Electron Potential for Pauli repulsion energy calculations.

class EETCouplingOEPotential

Generalized One-Electron Potential for EET coupling calculations.

· class PotentialInt

Computes potential integrals.

· class CPHF

CPHF solver class.

· class DIISManager

DIIS manager.

class ESPSolver

Charges from Electrostatic Potential (ESP). A solver-type class.

• class AllAOShellCombinationsIterator

Loop over all possible ERI shells.

class AllAOIntegralsIterator

Loop over all possible ERI within a particular shell.

class OEPDevSolver

Solver of properties of molecular aggregates. Abstract base.

- class ElectrostaticEnergySolver
- · class Points3DIterator

Iterator over a collection of points in 3D space. Abstract base.

· class CubePoints3DIterator

Iterator over a collection of points in 3D space. g09 Cube-like order.

· class RandomPoints3DIterator

Iterator over a collection of points in 3D space. Random collection.

class PointsCollection3D

Collection of points in 3D space. Abstract base.

class RandomPointsCollection3D

Collection of random points in 3D space.

class CubePointsCollection3D

G09 cube-like ordered collection of points in 3D space.

class ScalarField3D

Scalar field in 3D space. Abstract base.

· class ElectrostaticPotential3D

Electrostatic potential of a molecule.

class OEPotential3D

Class template for OEP scalar fields.

· class WavefunctionUnion

Union of two Wavefunction objects.

### **Typedefs**

- using **SharedWavefunction** = std::shared ptr< Wavefunction >
- using SharedBasisSet = std::shared\_ptr< BasisSet >
- using **SharedTensor** = std::shared ptr< Tensor >
- using SharedMatrix = std::shared\_ptr< Matrix >
- using SharedVector = std::shared\_ptr< Vector >
- using SharedScalarField3D = std::shared ptr< ScalarField3D >
- using SharedIntegralFactory = std::shared\_ptr< IntegralFactory >
- using SharedTwoBodyAOInt = std::shared ptr< TwoBodyAOInt >
- using SharedWavefunctionUnion = std::shared\_ptr< WavefunctionUnion >
- using SharedOEPotential = std::shared\_ptr< OEPotential >
- using SharedMolecule = std::shared\_ptr< Molecule >
- using SharedSuperFunctional = std::shared\_ptr< SuperFunctional >
- using SharedMOSpace = std::shared ptr< MOSpace >
- using **SharedMOSpaceVector** = std::vector< std::shared ptr< MOSpace >>
- using SharedIntegralTransform = std::shared\_ptr< IntegralTransform >
- using SharedLocalizer = std::shared\_ptr< Localizer >

#### **Functions**

void preambule (void)

Print preambule for module OEPDEV.

- std::shared\_ptr< SuperFunctional > create\_superfunctional (std::string name, Options & options)
  - Set up DFT functional.
- std::shared\_ptr< Molecule > extract\_monomer (std::shared\_ptr< const Molecule > molecule\_dimer, int id)
   Extract molecule from dimer.
- std::shared\_ptr< Wavefunction > solve\_scf (std::shared\_ptr< Molecule > molecule, std::shared\_ptr< Basis-Set > primary, std::shared\_ptr< SuperFunctional > functional, Options &options, std::shared\_ptr< PSIO > psio)

Solve RHF-SCF equations for a given molecule in a given basis set.

### 8.1.1 Detailed Description

oepdev module namespace. Contains:

#### 8.1.2 Function Documentation

8.1.2.1 std::shared\_ptr< SuperFunctional > oepdev::create\_superfunctional ( std::string name, Options & options )

Set up DFT functional.

Now it accepts only pure HF functional.

#### **Parameters**

name	name of the functional ("HF" is now only available)
options	psi::Options object

#### Returns

psi::SharedSuperFunctional object with functional.

8.1.2.2 std::shared\_ptr< Molecule > oepdev::extract\_monomer ( std::shared\_ptr< const Molecule > molecule\_dimer, int id )

Extract molecule from dimer.

#### **Parameters**

molecule_dimer	psi::SharedMolecule object with dimer
id	index of a molecule (starts from 1)

#### Returns

psi::SharedMolecule object with indicated monomer

8.1.2.3 std::shared\_ptr< Wavefunction > oepdev::solve\_scf ( std::shared\_ptr< Molecule > molecule, std::shared\_ptr< BasisSet > primary, std::shared\_ptr< SuperFunctional > functional, Options & options, std::shared\_ptr< PSIO > psio )

Solve RHF-SCF equations for a given molecule in a given basis set.

#### **Parameters**

molecule	psi::SharedMolecule object with molecule
primary	shared primary basis set
functional	DFT functional
options	psi::Options object
psio	psi::PSIO object

### Returns

psi::SharedWavefunction SCF wavefunction of the molecule

## 8.2 psi Namespace Reference

Psi4 package namespace.

### **Typedefs**

- using SharedVetor = std::shared\_ptr< Vector >
- using SharedBasisSet = std::shared\_ptr< BasisSet >

- using **SharedMolecule** = std::shared\_ptr< Molecule >
- using  $SharedMatrix = std::shared\_ptr < Matrix >$
- using SharedWavefunction = std::shared\_ptr< Wavefunction >

## 8.2.1 Detailed Description

Psi4 package namespace. Contains all Psi4 functionalities.

## **Class Documentation**

## 9.1 oepdev::AllAOIntegralsIterator Class Reference

Loop over all possible ERI within a particular shell.

```
#include <integrals_iter.h>
```

#### **Public Member Functions**

• AllAOIntegralsIterator (const AllAOShellCombinationsIterator &shellIter)

Construct by shell iterator (const object)

AllAOIntegralsIterator (std::shared\_ptr< AllAOShellCombinationsIterator > shellIter)

Construct by shell iterator (pointed by shared pointer)

• void first ()

First iteration.

• void next ()

Next iteration.

• bool is\_done ()

Check status of iterations.

· int i () const

Grab the current integral i index.

• int j () const

Grab the current integral j index.

• int k () const

Grab the current integral k index.

• int I () const

Grab the current integral I index.

• int index () const

## 9.1.1 Detailed Description

Loop over all possible ERI within a particular shell.

Constructed by providing a const reference or shared pointer to an AllAOShellCombinationsIterator object.

Suggested usage:

#### See Also

AllAOShellCombinationsIterator

20 Class Documentation

#### 9.1.2 Constructor & Destructor Documentation

9.1.2.1 oepdev::AllAOIntegralsIterator::AllAOIntegralsIterator ( const AllAOShellCombinationsIterator & shellIter )

Construct by shell iterator (const object)

**Parameters** 

```
shellIter - shell iterator object
```

9.1.2.2 oepdev::AllAOIntegralsIterator::AllAOIntegralsIterator ( std::shared\_ptr< AllAOShellCombinationsIterator > shellIter )

Construct by shell iterator (pointed by shared pointer)

**Parameters** 

```
shellIter - shell iterator object
```

#### 9.1.3 Member Function Documentation

9.1.3.1 int oepdev::AllAOIntegralsIterator::index ( ) const [inline]

Grab the current index of integral value stored in the buffer

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

- · oepdev/libutil/integrals\_iter.h
- · oepdev/libutil/integrals\_iter.cc

## 9.2 oepdev::AllAOShellCombinationsIterator Class Reference

Loop over all possible ERI shells.

```
#include <integrals_iter.h>
```

#### **Public Member Functions**

AllAOShellCombinationsIterator (SharedBasisSet bs\_1, SharedBasisSet bs\_2, SharedBasisSet bs\_3, SharedBasisSet bs\_4)

Construct by providing basis sets for each axis. The basis sets must be defined for the same molecule.

• AllAOShellCombinationsIterator (SharedIntegralFactory integrals)

Construct by providing integral factory.

• void first ()

First iteration.

· void next ()

Next iteration.

• bool is done ()

Check status of iterations.

• int P () const

Grab the current shell P index.

• int Q () const

Grab the current shell Q index.

· int R () const

Grab the current shell R index.

• int S () const

Grab the current shell S index.

• SharedBasisSet bs\_1 () const

Grab the basis set of axis 1.

• SharedBasisSet bs 2 () const

Grab the basis set of axis 2.

SharedBasisSet bs\_3 () const

Grab the basis set of axis 3.

SharedBasisSet bs\_4 () const

Grab the basis set of axis 4.

void compute\_shell (SharedTwoBodyAOInt tei) const

Compute ERI's for the current shell. The eris are stored in the buffer of the argument object.

#### 9.2.1 Detailed Description

Loop over all possible ERI shells.

Constructed by providing shared pointer to IntegralFactory object or shared pointers to four basis set spaces.

Suggested usage:

```
SharedIntegralFactory ints = std::make_shared<IntegralFactory>(bs1, bs2, bs3, bs4);
SharedTwoBodyAoInt tei(ints->eri());
AllAoShellCombinationsIterator shellIter(ints);
const double * buffer = tei->buffer();
for (shellIter.first(); shellIter.is_done()==false; shellIter.next())
{
    shellIter.compute_shell(tei);
    AllAoIntegralsIterator intsIter(shellIter);
    for (intsIter.first(); intsIter.is_done()==false; intsIter.next())
    {
        // Grab (ij|kl) integrals and indices here
        int i = intsIter.i();
        int j = intsIter.k();
        int k = intsIter.k();
        int l = intsIter.l();
        double integral = buffer[intsIter.index()];
    }
}
```

#### 9.2.2 Constructor & Destructor Documentation

9.2.2.1 oepdev::AllAOShellCombinationsIterator::AllAOShellCombinationsIterator ( SharedBasisSet bs\_1, SharedBasisSet bs\_2, SharedBasisSet bs\_3, SharedBasisSet bs\_4)

Construct by providing basis sets for each axis. The basis sets must be defined for the same molecule.

#### **Parameters**

bs_1	- basis set of axis 1
bs_2	- basis set of axis 2
bs_3	- basis set of axis 3
bs_4	- basis set of axis 4

9.2.2.2 oepdev::AllAOShellCombinationsIterator::AllAOShellCombinationsIterator ( SharedIntegralFactory integrals )

Construct by providing integral factory.

22 Class Documentation

#### **Parameters**

integrals - integral factory object

#### 9.2.3 Member Function Documentation

9.2.3.1 void oepdev::AllAOShellCombinationsIterator::compute\_shell ( SharedTwoBodyAOInt tei ) const

Compute ERI's for the current shell. The eris are stored in the buffer of the argument object.

**Parameters** 

tei - two electron AO integral

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

- · oepdev/libutil/integrals iter.h
- · oepdev/libutil/integrals\_iter.cc

## 9.3 oepdev::CPHF Class Reference

**CPHF** solver class.

#include <cphf.h>

#### **Public Member Functions**

• CPHF (SharedWavefunction ref\_wfn, Options &options)

orbital-associated polarizabilities tensors

•  $\sim$ CPHF ()

Desctructor.

void compute (void)

run the calculations

· void print (void) const

print to output file

• std::shared\_ptr< Matrix > get\_molecular\_polarizability (void) const

retrieve the molecular (total) polarizability

#### **Protected Attributes**

const int \_no

Number of occupied orbitals.

· const int \_nv

Number of virtual orbitals.

const int \_nn

Number of basis functions.

· long int \_memory

Memory.

int \_maxiter

Maximum number of iterations.

double <u>conv</u>

CPHF convergence threshold.

· bool \_with\_diis

whether use DIIS or not

• const int \_diis\_dim

Size of subspace.

std::shared\_ptr< BasisSet > \_primary

Primary Basis Set.

std::shared\_ptr< Matrix > \_cocc

Occupied orbitals.

std::shared ptr< Matrix > cvir

Virtual orbitals.

std::shared\_ptr< Vector > \_eps\_occ

Occupied orbital energies.

std::shared ptr< Vector > eps vir

Virtual orbital energies.

• std::vector< std::shared ptr

< oepdev::DIISManager > > \_diis

the DIIS managers for each perturbation operator x, y and z

• Options options

Options.

std::shared\_ptr< Matrix > \_molecular\_polarizability

Total (molecular) polarizability tensor.

#### 9.3.1 Detailed Description

**CPHF** solver class.

Solves CPHF equations (now only for RHF wavefunction). Computes molecular and orbital-associated polarizabilities

Suggested usage:

std::shared\_ptr<CPHF> cphf(new CPHF(ref\_wfn, options)); cphf->compute(); std::shared\_ptr<Matrix> polarizability = cphf->get\_molecular\_polarizability(); std::shared\_ptr<Tensor> orbital\_polars = cphf->get\_orbital\_polarizabilities();

#### 9.3.2 Constructor & Destructor Documentation

9.3.2.1 oepdev::CPHF::CPHF ( SharedWavefunction ref\_wfn, Options & options )

orbital-associated polarizabilities tensors

Constructor

Parameters

ref_wfn	reference HF wavefunction
options	set of Psi4 options

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

- · oepdev/libutil/cphf.h
- · oepdev/libutil/cphf.cc

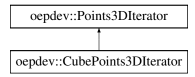
24 Class Documentation

## 9.4 oepdev::CubePoints3DIterator Class Reference

Iterator over a collection of points in 3D space. g09 Cube-like order.

```
#include <space3d.h>
```

Inheritance diagram for oepdev::CubePoints3DIterator:



#### **Public Member Functions**

- CubePoints3Dlterator (const int &nx, const int &ny, const int &nz, const double &dx, const double &dy, const double &dx, const double &dx, const double &ox)
- virtual void first ()

Initialize first iteration.

virtual void next ()

Step to next iteration.

#### **Protected Attributes**

- const int nx
- · const int ny\_
- · const int nz\_
- · const double dx\_
- · const double dy\_
- · const double dz\_
- const double ox\_
- const double oy\_
- const double oz\_int ii\_
- int **jj**\_
- int kk

#### **Additional Inherited Members**

#### 9.4.1 Detailed Description

Iterator over a collection of points in 3D space. g09 Cube-like order.

**Note:** Always create instances by using static factory method from Points3Dlterator. Do not use constructor of this class.

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

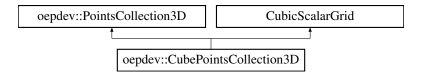
- · oepdev/libutil/space3d.h
- · oepdev/libutil/space3d.cc

## 9.5 oepdev::CubePointsCollection3D Class Reference

G09 cube-like ordered collection of points in 3D space.

#include <space3d.h>

Inheritance diagram for oepdev::CubePointsCollection3D:



#### **Public Member Functions**

- CubePointsCollection3D (Collection collectionType, const int &nx, const int &ny, const int &nz, const double &px, const double &px, const double &px, psi::SharedBasisSet bs, psi::Options &options)
- · virtual void print () const

Print the information to Psi4 output file.

virtual void write\_cube\_file (psi::SharedMatrix v, const std::string &name)

#### **Additional Inherited Members**

### 9.5.1 Detailed Description

G09 cube-like ordered collection of points in 3D space.

**Note:** Do not use constructors of this class explicitly. Instead, use static factory methods of the superclass to create instances.

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

- · oepdev/libutil/space3d.h
- · oepdev/libutil/space3d.cc

## 9.6 oepdev::DIISManager Class Reference

#### DIIS manager.

#include <diis.h>

#### **Public Member Functions**

- DIISManager (int dim, int na, int nb)
- ∼DIISManager ()

Destructor.

- void put (const std::shared\_ptr< const Matrix > &error, const std::shared\_ptr< const Matrix > &vector)
- void compute (void)
- void update (std::shared\_ptr< Matrix > &other)

26 Class Documentation

### 9.6.1 Detailed Description

#### DIIS manager.

Instance can interact directly with the process of solving vector quantities in iterative manner. One needs to pass the dimensions of solution vector as well as the DIIS subspace size. The iterative procedure requires providing the current vector and also an estimate of the error vector. The updated DIIS vector can be copied to an old vector through the Instance.

#### 9.6.2 Constructor & Destructor Documentation

9.6.2.1 oepdev::DIISManager::DIISManager ( int dim, int na, int nb )

#### Constructor.

#### **Parameters**

dim	Size of DIIS subspace
na	Number of solution rows
nb	Number of solution columns

#### 9.6.3 Member Function Documentation

9.6.3.1 void oepdev::DIISManager::compute (void)

Perform DIIS interpolation.

9.6.3.2 void oepdev::DIISManager::put ( const std::shared\_ptr< const Matrix > & error, const std::shared\_ptr< const Matrix > & vector )

Put the current solution to the DIIS manager.

#### **Parameters**

error	Shared matrix with current solution error
vector	Shared matrix with current solution vector

#### 9.6.3.3 void oepdev::DIISManager::update ( std::shared\_ptr< Matrix > & other )

Update solution vector. Pass the Shared pointer to current solution. Then it will be overriden by the updated DIIS solution.

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

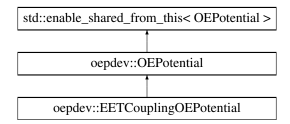
- · oepdev/libutil/diis.h
- · oepdev/libutil/diis.cc

## 9.7 oepdev::EETCouplingOEPotential Class Reference

Generalized One-Electron Potential for EET coupling calculations.

#include <oep.h>

Inheritance diagram for oepdev::EETCouplingOEPotential:



#### **Public Member Functions**

- EETCouplingOEPotential (SharedWavefunction wfn, SharedBasisSet auxiliary, Options & options)
- **EETCouplingOEPotential** (SharedWavefunction wfn, Options &options)
- virtual void compute (const std::string &oepType) override
- virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, double &v) override
- virtual void print\_header () const override

#### **Additional Inherited Members**

### 9.7.1 Detailed Description

Generalized One-Electron Potential for EET coupling calculations.

Contains the following OEP types: "ET1" "ET2" "HT1" "HT1" "HT2" "CT1" "CT2"

### 9.7.2 Member Function Documentation

9.7.2.1 void EETCouplingOEPotential::compute\_3D ( const std::string & oepType, const double & x, const double & y, const double & y, const double & z, double & v ) [override], [virtual]

Compute value of potential in point x, y, z and save at v

Implements oepdev::OEPotential.

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

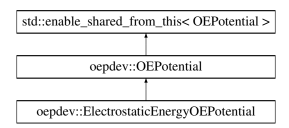
- · oepdev/liboep/oep.h
- · oepdev/liboep/oep.cc

## 9.8 oepdev::ElectrostaticEnergyOEPotential Class Reference

Generalized One-Electron Potential for Electrostatic Energy calculations.

#include <oep.h>

Inheritance diagram for oepdev::ElectrostaticEnergyOEPotential:



28 Class Documentation

#### **Public Member Functions**

- ElectrostaticEnergyOEPotential (SharedWavefunction wfn, Options & options)
  - Only ESP-based potential is worth implementing.
- virtual void compute (const std::string &oepType) override
- virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, double &v) override
- virtual void print\_header () const override

#### **Additional Inherited Members**

#### 9.8.1 Detailed Description

Generalized One-Electron Potential for Electrostatic Energy calculations.

Contains the following OEP types: "V"

#### 9.8.2 Member Function Documentation

9.8.2.1 void ElectrostaticEnergyOEPotential::compute\_3D ( const std::string & oepType, const double & x, const double & y, const double & v ) [override], [virtual]

Compute value of potential in point x, y, z and save at v

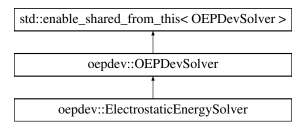
Implements oepdev::OEPotential.

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

- · oepdev/liboep/oep.h
- oepdev/liboep/oep.cc

## 9.9 oepdev::ElectrostaticEnergySolver Class Reference

 $Inheritance\ diagram\ for\ oepdev:: Electrostatic Energy Solver:$ 



#### **Public Member Functions**

- ElectrostaticEnergySolver (SharedWavefunctionUnion wfn\_union)
- virtual double compute\_oep\_based (const std::string &method="DEFAULT")
   Compute property by using OEP's.
- virtual double compute\_benchmark (const std::string &method="DEFAULT")
   Compute property by using benchmark method.

#### **Additional Inherited Members**

#### 9.9.1 Member Function Documentation

9.9.1.1 double ElectrostaticEnergySolver::compute\_benchmark ( const std::string & method = "DEFAULT" )

[virtual]

Compute property by using benchmark method.

#### **Parameters**

```
method - benchmark method
```

Implements oepdev::OEPDevSolver.

```
9.9.1.2 double ElectrostaticEnergySolver::compute_oep_based ( const std::string & method = "DEFAULT" )
[virtual]
```

Compute property by using OEP's.

#### **Parameters**

```
method - flavour of OEP model
```

Implements oepdev::OEPDevSolver.

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

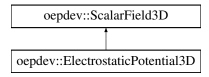
- oepdev/libutil/solver.h
- · oepdev/libutil/solver.cc

## 9.10 oepdev::ElectrostaticPotential3D Class Reference

Electrostatic potential of a molecule.

```
#include <space3d.h>
```

Inheritance diagram for oepdev::ElectrostaticPotential3D:



#### **Public Member Functions**

- ElectrostaticPotential3D (const int &np, const double &radius, const double &cx, const double &cy, const double &cz)
- **ElectrostaticPotential3D** (const int &np, const double &padding, psi::SharedWavefunction wfn, psi::Options &options)
- ElectrostaticPotential3D (const int &nx, const int &ny, const int &nz, const double &px, const double &px, const double &px, psi::SharedWavefunction wfn, psi::Options &options)
- virtual double compute\_xyz (const double &x, const double &y, const double &z)

Compute a value of scalar field at point (x, y, z)

virtual void print () const

Print information of the object to Psi4 output.

30 Class Documentation

#### **Additional Inherited Members**

#### 9.10.1 Detailed Description

Electrostatic potential of a molecule.

Computes the electrostatic potential of a molecule directly from the wavefunction. The electrostatic potential  $v(\mathbf{r})$  at point  $\mathbf{r}$  is computed from the following formula:

$$v(\mathbf{r}) = v_{\text{nuc}}(\mathbf{r}) + v_{\text{el}}(\mathbf{r})$$

where the nuclear and electronic contributions are defined accordingly as

$$v_{\text{nuc}}(\mathbf{r}) = \sum_{x} \frac{Z_{x}}{|\mathbf{r} - \mathbf{r}_{x}|}$$
$$v_{\text{el}}(\mathbf{r}) = \sum_{\mu\nu} \left\{ D_{\mu\nu}^{(\alpha)} + D_{\mu\nu}^{(\beta)} \right\} V_{\nu\mu}(\mathbf{r})$$

In the above equations,  $Z_x$  denotes the charge of xth nucleus,  $D_{\mu\nu}^{(\omega)}$  is the one-particle (relaxed) density matrix element in AO basis associated with the  $\omega$  electron spin, and  $V_{\mu\nu}(\mathbf{r})$  is the potential one-electron integral defined by

$$V_{\nu\mu}(\mathbf{r}) \equiv \int d\mathbf{r}' \phi_{\nu}^*(\mathbf{r}') \frac{1}{|\mathbf{r} - \mathbf{r}'|} \phi_{\mu}(\mathbf{r}')$$

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

- · oepdev/libutil/space3d.h
- · oepdev/libutil/space3d.cc

## 9.11 oepdev::ESPSolver Class Reference

Charges from Electrostatic Potential (ESP). A solver-type class.

#include <esp.h>

#### **Public Member Functions**

• ESPSolver (SharedScalarField3D field)

Construct from scalar field.

ESPSolver (SharedScalarField3D field, psi::SharedMatrix centres)

Construct from scalar field.

• virtual  $\sim$ ESPSolver ()

Destructor.

· virtual psi::SharedVector charges () const

Get the (fit) charges.

• virtual psi::SharedMatrix centres () const

Get the charge distribution centres.

virtual void compute ()

Perform fitting of effective charges.

#### **Protected Attributes**

const int nCentres

Number of fit centres.

• SharedScalarField3D field\_

Scalar field.

psi::SharedVector charges\_

Charges to be fit.

psi::SharedMatrix centres

Centres, at which fit charges will reside.

# 9.11.1 Detailed Description

Charges from Electrostatic Potential (ESP). A solver-type class.

Solves the least-squares problem to fit the generalized charges  $q_m$ , that reproduce the reference generalized potential  $v^{\text{ref}}(\mathbf{r})$  supplied by the ScalarField3D object:

$$\int d\mathbf{r}' \left[ v^{\rm ref}(\mathbf{r}') - \sum_m \frac{q_m}{|\mathbf{r}' - \mathbf{r}_m|} \right]^2 \to {\rm minimize}$$

The charges are subject to the following constraint:

$$\sum_{m}q_{m}=0$$

Method description.

M generalized charges is found by solving the matrix equation

$$\begin{pmatrix} \mathbf{A} & 0 \\ 0 & 1 \end{pmatrix}^{-1} \cdot \begin{pmatrix} \mathbf{b} \\ 0 \end{pmatrix} = \begin{pmatrix} \mathbf{q} \\ \lambda \end{pmatrix}$$

where the  ${\bf A}$  matrix of dimension  $M \times M$  and b} vector or length M are given as

$$A_{mn} = \sum_{i} rac{1}{r_{im}r_{in}} \ b_m = \sum_{i} rac{v^{ ext{ref}}(\mathbf{r}_m)}{r_{im}}$$

In the above equation, summations run over all sample points, at which reference potential is known.

#### 9.11.2 Constructor & Destructor Documentation

9.11.2.1 oepdev::ESPSolver::ESPSolver ( SharedScalarField3D field )

Construct from scalar field.

Assume that the centres are on atoms associated with the scalar field.

**Parameters** 

field - oepdev scalar field object

9.11.2.2 oepdev::ESPSolver::ESPSolver ( SharedScalarField3D field, psi::SharedMatrix centres )

Construct from scalar field.

Solve ESP equations for a custom set of charge distribution centres.

#### **Parameters**

field	- oepdev scalar field object
centres	- matrix with coordinates of charge distribution centres

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

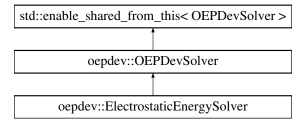
- · oepdev/libutil/esp.h
- · oepdev/libutil/esp.cc

# 9.12 oepdev::OEPDevSolver Class Reference

Solver of properties of molecular aggregates. Abstract base.

#include <solver.h>

Inheritance diagram for oepdev::OEPDevSolver:



# **Public Member Functions**

• OEPDevSolver (SharedWavefunctionUnion wfn union)

Take wavefunction union and initialize the Solver.

virtual ∼OEPDevSolver ()

Destroctor.

- virtual double compute\_oep\_based (const std::string &method="DEFAULT")=0
   Compute property by using OEP's.
- virtual double compute\_benchmark (const std::string &method="DEFAULT")=0
   Compute property by using benchmark method.

# **Static Public Member Functions**

- static std::shared\_ptr
  - < OEPDevSolver > build (const std::string &target, SharedWavefunctionUnion wfn\_union)

Build a solver of a particular property for given molecular cluster.

#### **Protected Attributes**

SharedWavefunctionUnion wfn union

Wavefunction union.

std::vector< std::string > methods\_oepBased\_

Names of all OEP-based methods implemented for a solver.

std::vector< std::string > methods\_benchmark\_

Names of all benchmark methods implemented for a solver.

# 9.12.1 Detailed Description

Solver of properties of molecular aggregates. Abstract base.

# 9.12.2 Constructor & Destructor Documentation

9.12.2.1 OEPDevSolver::OEPDevSolver ( SharedWavefunctionUnion wfn\_union )

Take wavefunction union and initialize the Solver.

#### **Parameters**

wfn_union	- wavefunction union of isolated molecular wavefunctions

#### 9.12.3 Member Function Documentation

9.12.3.1 std::shared\_ptr< OEPDevSolver > OEPDevSolver::build ( const std::string & target, SharedWavefunctionUnion wfn\_union ) [static]

Build a solver of a particular property for given molecular cluster.

#### **Parameters**

target	- target property
wfn_union	- wavefunction union of isolated molecular wavefunctions

9.12.3.2 double OEPDevSolver::compute\_benchmark( const std::string & method = "DEFAULT") [pure virtual]

Compute property by using benchmark method.

#### **Parameters**

method	- benchmark method

Implemented in oepdev::ElectrostaticEnergySolver.

9.12.3.3 double OEPDevSolver::compute\_oep\_based ( const std::string & method = "DEFAULT" ) [pure virtual]

Compute property by using OEP's.

## **Parameters**

method
--------

Implemented in oepdev::ElectrostaticEnergySolver.

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

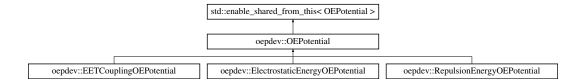
- · oepdev/libutil/solver.h
- · oepdev/libutil/solver.cc

# 9.13 oepdev::OEPotential Class Reference

Generalized One-Electron Potential: Abstract base.

#include <oep.h>

Inheritance diagram for oepdev::OEPotential:



# **Public Member Functions**

• OEPotential (SharedWavefunction wfn, Options & options)

ESP-based OEP object.

• OEPotential (SharedWavefunction wfn, SharedBasisSet auxiliary, Options & options)

DF-based OEP object.

virtual ∼OEPotential ()

Destructor.

· virtual void rotate (const Matrix &rotmat)

Rotate

virtual void translate (const Vector &trans)

Translate

virtual void superimpose (const Matrix &refGeometry, const std::vector< int > &supList, const std::vector< int > &reordList)

Superimpose.

• std::string name () const

Retrieve name of this OEP.

SharedMatrix matrix (const std::string &oepType) const

Retrieve matrix potential.

• SharedWavefunction wfn () const

Retrieve wavefunction object.

- void **set\_name** (const std::string &name)
- virtual void print\_header () const =0
- virtual void compute (const std::string &oepType)=0
- virtual void compute (void)
- virtual void write cube (const std::string &oepType, const std::string &fileName)
- virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, double &v)=0

# **Static Public Member Functions**

- · static std::shared\_ptr
  - < OEPotential > build (const std::string &category, SharedWavefunction wfn, Options &options)

Build ESP-based OEP object.

· static std::shared\_ptr

< OEPotential > build (const std::string &category, SharedWavefunction wfn, SharedBasisSet auxiliary, Options &options)

Build DF-based OEP object.

#### **Public Attributes**

· const bool is\_density\_fitted

Is this OEP density-fitted?

· const bool is\_esp\_based

Is this OEP ESP-based?

# **Protected Attributes**

· Options options\_

Psi4 options.

SharedWavefunction wfn

Wavefunction.

SharedBasisSet primary\_

Promary Basis set.

SharedBasisSet auxiliary\_

Auxiliary Basis set.

• std::string name\_

Name of this OEP;.

std::vector< std::string > oepTypes\_

Types of OEP's within the scope of this object.

std::map< std::string,</li>

SharedMatrix > oepMatrices

OEP's matrix forms for each OEP type.

· std::shared\_ptr

< psi::IntegralFactory > intsFactory\_

Integral factory.

std::shared\_ptr< psi::Matrix > potMat\_

Matrix of potential one-electron integrals.

· std::shared\_ptr

< psi::OneBodyAOInt > OEInt\_

One-electron integral shared pointer.

std::shared\_ptr< PotentialInt > potInt\_

One-electron potential shared pointer.

# 9.13.1 Detailed Description

Generalized One-Electron Potential: Abstract base.

Manages OEP's in matrix and 3D forms.

# 9.13.2 Constructor & Destructor Documentation

9.13.2.1 OEPotential::OEPotential ( SharedWavefunction wfn, Options & options )

ESP-based OEP object.

**Parameters** 

wfn	- wavefunction
options	- Psi4 options

9.13.2.2 OEPotential::OEPotential ( SharedWavefunction wfn, SharedBasisSet auxiliary, Options & options )

DF-based OEP object.

#### **Parameters**

ſ	wfn	- wavefunction
ſ	auxiliary	- basis set for density fitting of OEP's
Ī	options	- Psi4 options

# 9.13.3 Member Function Documentation

9.13.3.1 std::shared\_ptr< OEPotential > OEPotential::build ( const std::string & category, SharedWavefunction wfn, Options & options ) [static]

Build ESP-based OEP object.

#### **Parameters**

type	- OEP category
wfn	- wavefunction
options	- Psi4 options

9.13.3.2 std::shared\_ptr< OEPotential > OEPotential::build ( const std::string & category, SharedWavefunction wfn, SharedBasisSet auxiliary, Options & options ) [static]

Build DF-based OEP object.

#### **Parameters**

type	- OEP category
wfn	- wavefunction
auxiliary	- basis set for density fitting of OEP's
options	- Psi4 options

9.13.3.3 void OEPotential::compute\_3D ( const std::string & oepType, const double & x, const double & y, const double & z, double & v ) [pure virtual]

Compute value of potential in point x, y, z and save at v

Implemented in oepdev::EETCouplingOEPotential, oepdev::RepulsionEnergyOEPotential, and oepdev::ElectrostaticEnergyOEPotential.

9.13.3.4 void OEPotential::write\_cube ( const std::string & oepType, const std::string & fileName ) [virtual]

Write potential to a cube file

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

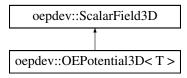
- · oepdev/liboep/oep.h
- oepdev/liboep/oep.cc

# 9.14 oepdev::OEPotential3D< T > Class Template Reference

Class template for OEP scalar fields.

#include <space3d.h>

Inheritance diagram for oepdev::OEPotential3D< T>:



#### **Public Member Functions**

- OEPotential3D (const int &np, const double &padding, std::shared\_ptr< T > oep, const std::string &oepType)

  Construct random spherical collection of scalar field of type T.
- OEPotential3D (const int &nx, const int &ny, const int &nz, const double &px, const double &py, const double &px, std::shared\_ptr< T > oep, const std::string &oepType, psi::Options &options)

Construct ordered 3D collection of scalar field of type T.

virtual ∼OEPotential3D ()

Destructor.

virtual double compute\_xyz (const double &x, const double &y, const double &z)

Compute a value of scalar field at point (x, y, z)

virtual void print () const

Print information of the object to Psi4 output.

#### **Protected Attributes**

std::shared\_ptr< T > oep\_

Shared pointer to the instance of class T

std::string oepType\_

Descriptor of the scalar field type stored in instance of  ${\it T}$ 

## **Additional Inherited Members**

# 9.14.1 Detailed Description

template < class T> class oepdev::OEPotential3D< T>

Class template for OEP scalar fields.

Used for special type of classes T that contain following public member functions:

with the descriptor of a certain scalar field type, x, y, z the points in 3D space in which the scalar field has to be computed and stored at v. Instances of T should store shared pointer to wavefunction object. List of classes T that are compatible with this class template and are currently implemented in oppdev is given below:

• oepdev::OEPotential abstract base (do not use derived classes as T)

Template parameters:

#### **Template Parameters**

T	the compatible class (e.g. oepdev::OEPotential)

# 9.14.2 Constructor & Destructor Documentation

9.14.2.1 template < class T > oepdev::OEPotential3D< T >::OEPotential3D ( const int & np, const double & padding, std::shared\_ptr< T > oep, const std::string & oepType )

Construct random spherical collection of scalar field of type T.

The points are drawn according to uniform distrinution in 3D space.

#### **Parameters**

np	- number of points to draw
padding	- spherical padding distance (au)
оер	- OEP object of type T
оерТуре	- type of OEP

9.14.2.2 template < class T > oepdev::OEPotential3D < T >::OEPotential3D ( const int & nx, const int & ny, const int & ny, const int & nz, const double & px, const double & px, std::shared\_ptr < T > oep, const std::string & oepType, psi::Options & options )

Construct ordered 3D collection of scalar field of type T.

The points are generated according to Gaussian cube file format.

#### **Parameters**

nx	- number of points along x direction
ny	- number of points along y direction
nz	- number of points along z direction
px	- padding distance along x direction
ру	- padding distance along y direction
pz	- padding distance along z direction
oep	- OEP object of type T
оерТуре	- type of OEP
options	- Psi4 options object

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

· oepdev/libutil/space3d.h

# 9.15 oepdev::Points3Dlterator::Point Struct Reference

#### **Public Attributes**

- double x
- double **v**
- double z
- int index

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

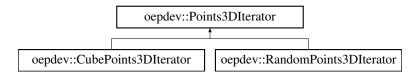
· oepdev/libutil/space3d.h

# 9.16 oepdev::Points3Dlterator Class Reference

Iterator over a collection of points in 3D space. Abstract base.

#include <space3d.h>

Inheritance diagram for oepdev::Points3Dlterator:



#### **Classes**

struct Point

#### **Public Member Functions**

• Points3DIterator (const int &np)

Plain constructor. Initializes the abstract features.

virtual ∼Points3DIterator ()

Destructor.

virtual bool is\_done ()

Check if iteration is finished.

• virtual void first ()=0

Initialize first iteration.

virtual void next ()=0

Step to next iteration.

- virtual double x () const
- virtual double y () const
- virtual double z () const
- · virtual int index () const

#### **Static Public Member Functions**

• static shared\_ptr

< Points3DIterator > build (const int &nx, const int &ny, const int &nz, const double &dx, const double &dx, const double &dx, const double &ox, const double &ox)

Build G09 Cube collection iterator.

· static shared ptr

< Points3DIterator > build (const int &np, const double &radius, const double &cx, const double &cy, const double &cz)

Build random collection iterator.

static shared ptr

< Points3DIterator > build (const int &np, const double &pad, psi::SharedMolecule mol)

Build random collection iterator.

# **Protected Attributes**

· const int np\_

Number of points.

• bool done\_

Status of the iterator.

int index

Current index.

Point current\_

# 9.16.1 Detailed Description

Iterator over a collection of points in 3D space. Abstract base.

Points3DIterators are constructed either as iterators over:

- · a random collections or
- an ordered (g09 cube-like) collections. Note: Always create instances by using static factory methods.

# 9.16.2 Constructor & Destructor Documentation

9.16.2.1 oepdev::Points3Dlterator::Points3Dlterator ( const int & np )

Plain constructor. Initializes the abstract features.

**Parameters** 

np	- number of points this iterator is constructed for

# 9.16.3 Member Function Documentation

9.16.3.1 std::shared\_ptr< Points3Dlterator > oepdev::Points3Dlterator::build ( const int & nx, const int & ny, const int & nz, const double & dx, const double & dx, const double & ox, const double & oy, const double & oz ) [static]

Build G09 Cube collection iterator.

The points are generated according to Gaussian cube file format.

# **Parameters**

nx	- number of points along x direction
ny	- number of points along y direction
nz	- number of points along z direction
dx	- spacing distance along x direction
dy	- spacing distance along y direction
dz	- spacing distance along y direction
OX	- coordinate x of cube origin
oy	- coordinate y of cube origin
OZ	- coordinate z of cube origin

9.16.3.2 std::shared\_ptr< Points3Dlterator > oepdev::Points3Dlterator::build ( const int & np, const double & radius, const double & cx, const double & cy, const double & cz ) [static]

Build random collection iterator.

The points are drawn according to uniform distrinution in 3D space.

#### **Parameters**

np	- number of points to draw
radius	- sphere radius inside which points are to be drawn
CX	- coordinate x of sphere's centre
су	- coordinate y of sphere's centre
CZ	- coordinate z of sphere's centre

9.16.3.3 shared\_ptr< Points3Dlterator > oepdev::Points3Dlterator::build ( const int & np, const double & pad, psi::SharedMolecule mol ) [static]

Build random collection iterator.

The points are drawn according to uniform distrinution in 3D space enclosing a molecule given. All drawn points lie outside the van der Waals volume.

#### **Parameters**

np	- number of points to draw
pad	- radius padding of a minimal sphere enclosing the molecule
mol	- Psi4 molecule object

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

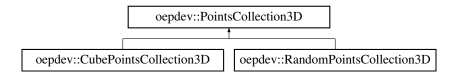
- · oepdev/libutil/space3d.h
- · oepdev/libutil/space3d.cc

# 9.17 oepdev::PointsCollection3D Class Reference

Collection of points in 3D space. Abstract base.

#include <space3d.h>

Inheritance diagram for oepdev::PointsCollection3D:



# **Public Types**

• enum Collection { Random, Cube }

Public descriptior of collection type.

# **Public Member Functions**

• PointsCollection3D (Collection collectionType, int &np)

Initialize abstract features.

- PointsCollection3D (Collection collectionType, const int &np)
- virtual ∼PointsCollection3D ()

Destructor.

· virtual int npoints () const

Get the number of points.

· virtual shared\_ptr

< Points3Dlterator > points\_iterator () const

Get the iterator over this collection of points.

virtual Collection get\_type () const

Get the collection type.

• virtual void print () const =0

Print the information to Psi4 output file.

#### **Static Public Member Functions**

· static shared ptr

< PointsCollection3D > build (const int &npoints, const double &radius, const double &cx=0.0, const double &cz=0.0)

Build random collection of points.

- · static shared\_ptr
  - < PointsCollection3D > build (const int &npoints, const double &padding, psi::SharedMolecule mol)

Build random collection of points.

- · static shared\_ptr
  - < PointsCollection3D > build (const int &nx, const int &ny, const int &nz, const double &px, const double &px, const double &px, psi::SharedBasisSet bs, psi::Options &options)

Build G09 Cube collection of points.

## **Protected Attributes**

const int np\_

Number of points.

• Collection collectionType\_

Collection type.

shared\_ptr< Points3DIterator > pointsIterator\_

iterator over points collection

#### 9.17.1 Detailed Description

Collection of points in 3D space. Abstract base.

Create random or ordered (g09 cube-like) collections of points in 3D space.

**Note:** Always create instances by using static factory methods.

# 9.17.2 Constructor & Destructor Documentation

9.17.2.1 oepdev::PointsCollection3D::PointsCollection3D ( Collection collectionType, int & np )

Initialize abstract features.

**Parameters** 

np	- number of points to be created

#### 9.17.3 Member Function Documentation

9.17.3.1 std::shared\_ptr< PointsCollection3D > oepdev::PointsCollection3D::build ( const int & *npoints*, const double & radius, const double & cx = 0.0, const double & cy = 0.0, const double & cz = 0.0) [static]

Build random collection of points.

Points uniformly span a sphere.

#### **Parameters**

npoints	- number of points to draw
radius	- sphere radius inside which points are to be drawn
CX	- coordinate x of sphere's centre
су	- coordinate y of sphere's centre
CZ	- coordinate z of sphere's centre

9.17.3.2 std::shared\_ptr< PointsCollection3D > oepdev::PointsCollection3D::build ( const int & npoints, const double & padding, psi::SharedMolecule mol ) [static]

Build random collection of points.

Points uniformly span space inside a sphere enclosing a molecule. exluding the van der Waals volume.

#### **Parameters**

np	- number of points to draw
padding	- radius padding of a minimal sphere enclosing the molecule
mol	- Psi4 molecule object

9.17.3.3 std::shared\_ptr< PointsCollection3D > oepdev::PointsCollection3D::build ( const int & nx, const int & ny, const int & nz, const double & px, const double & px, psi::SharedBasisSet bs, psi::Options & options ) [static]

Build G09 Cube collection of points.

The points span a parallelpiped according to Gaussian cube file format.

#### **Parameters**

nx	- number of points along x direction
ny	- number of points along y direction
nz	- number of points along z direction
px	- padding distance along x direction
ру	- padding distance along y direction
pz	- padding distance along z direction
bs	- Psi4 basis set object
options	- Psi4 options object

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

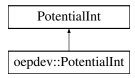
- · oepdev/libutil/space3d.h
- oepdev/libutil/space3d.cc

# 9.18 oepdev::PotentialInt Class Reference

Computes potential integrals.

#include <potential.h>

Inheritance diagram for oepdev::PotentialInt:



#### **Public Member Functions**

PotentialInt (std::vector< psi::SphericalTransform > &st, std::shared\_ptr< psi::BasisSet > bs1, std::shared\_ptr< psi::BasisSet > bs2, int deriv=0)

Constructor. Initialize identically like in psi::PotentilInt.

PotentialInt (std::vector< psi::SphericalTransform > &st, std::shared\_ptr< psi::BasisSet > bs1, std::shared\_ptr< psi::BasisSet > bs2, std::shared\_ptr< psi::Matrix > Qxyz, int deriv=0)

Constructor. Takes an arbitrary collection of charges.

• PotentialInt (std::vector< psi::SphericalTransform > &, std::shared\_ptr< psi::BasisSet >, std::shared\_ptr< psi::BasisSet >, const double &x, const double &z, const double &q=1.0, int deriv=0)

Constructor. Computes potential for one point x, y, z for a test particle of charge q.

void set\_charge\_field (const double &x, const double &y, const double &z, const double &q=1.0)
 Mutator. Set the charge field to be a x, y, z point of charge q.

#### 9.18.1 Detailed Description

Computes potential integrals.

# 9.18.2 Constructor & Destructor Documentation

9.18.2.1 oepdev::PotentialInt::PotentialInt ( std::vector< psi::SphericalTransform > & st, std::shared\_ptr< psi::BasisSet > bs1, std::shared\_ptr< psi::BasisSet > bs2, int deriv = 0 )

Constructor. Initialize identically like in psi::PotentilInt.

# **Parameters**

st	- Spherical transform object
bs1	- basis set for first space
bs2	- basis set for second space
deriv	- derivative level

9.18.2.2 oepdev::PotentialInt::PotentialInt (std::vector<psi::SphericalTransform > & st, std::shared\_ptr<psi::BasisSet > bs1, std::shared\_ptr<psi::BasisSet > bs2, std::shared\_ptr<psi::Matrix > Qxyz, int deriv = 0)

Constructor. Takes an arbitrary collection of charges.

## **Parameters**

st	- Spherical transform object

bs1	- basis set for first space
bs2	- basis set for second space
Qxyz	- matrix with charges and their positions
deriv	- derivative level

9.18.2.3 oepdev::PotentialInt::PotentialInt ( std::vector < psi::SphericalTransform > & st, std::shared\_ptr < psi::BasisSet > bs1, std::shared\_ptr < psi::BasisSet > bs2, const double & x, const double & y, const double & z, const double & q = 1.0, int deriv = 0)

Constructor. Computes potential for one point x, y, z for a test particle of charge q.

#### **Parameters**

st	- Spherical transform object
bs1	- basis set for first space
bs2	- basis set for second space
X	- x coordinate of q
У	- y coordinate of q
Z	- z coordinate of q
q	- value of the probe charge
deriv	- derivative level

#### 9.18.3 Member Function Documentation

9.18.3.1 void oepdev::PotentialInt::set\_charge\_field ( const double & x, const double & y, const double & z, const double & q = 1.0)

Mutator. Set the charge field to be a x, y, z point of charge q.

#### **Parameters**

Х	- x coordinate of q
у	- y coordinate of q
Z	- z coordinate of q
q	- value of the probe charge

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

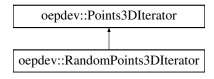
- · oepdev/libpsi/potential.h
- oepdev/libpsi/potential.cc

# 9.19 oepdev::RandomPoints3Dlterator Class Reference

Iterator over a collection of points in 3D space. Random collection.

#include <space3d.h>

Inheritance diagram for oepdev::RandomPoints3DIterator:



#### **Public Member Functions**

- RandomPoints3Dlterator (const int &np, const double &radius, const double &cx, const double &cy, const double &cz)
- RandomPoints3DIterator (const int &np, const double &pad, psi::SharedMolecule mol)
- virtual void first ()

Initialize first iteration.

• virtual void next ()

Step to next iteration.

#### **Protected Member Functions**

- virtual double random\_double ()
- virtual void draw\_random\_point ()
- virtual bool is\_in\_vdWsphere (double x, double y, double z) const

#### **Protected Attributes**

- double cx
- double cy\_
- double cz\_
- · double radius\_
- double r\_
- · double phi\_
- double theta\_
- double x
- double y\_
- double z\_
- psi::SharedMatrix excludeSpheres
- std::map< std::string, double > vdwRadius\_
- std::default\_random\_engine randomNumberGenerator\_
- std::uniform\_real\_distribution
  - $< {\sf double} > {\bf randomDistribution}\_$

#### **Additional Inherited Members**

# 9.19.1 Detailed Description

Iterator over a collection of points in 3D space. Random collection.

**Note:** Always create instances by using static factory method from Points3Dlterator. Do not use constructors of this class.

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

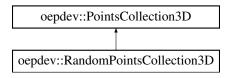
- · oepdev/libutil/space3d.h
- · oepdev/libutil/space3d.cc

# 9.20 oepdev::RandomPointsCollection3D Class Reference

Collection of random points in 3D space.

#include <space3d.h>

Inheritance diagram for oepdev::RandomPointsCollection3D:



#### **Public Member Functions**

- RandomPointsCollection3D (Collection collectionType, const int &npoints, const double &radius, const double &cx, const double &cy, const double &cz)
- RandomPointsCollection3D (Collection collectionType, const int &npoints, const double &padding, psi::-SharedMolecule mol)
- · virtual void print () const

Print the information to Psi4 output file.

#### **Additional Inherited Members**

# 9.20.1 Detailed Description

Collection of random points in 3D space.

**Note:** Do not use constructors of this class explicitly. Instead, use static factory methods of the superclass to create instances.

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

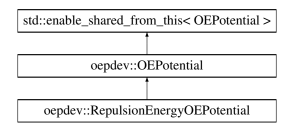
- · oepdev/libutil/space3d.h
- · oepdev/libutil/space3d.cc

# 9.21 oepdev::RepulsionEnergyOEPotential Class Reference

Generalized One-Electron Potential for Pauli repulsion energy calculations.

#include <oep.h>

Inheritance diagram for oepdev::RepulsionEnergyOEPotential:



**Public Member Functions** 

- RepulsionEnergyOEPotential (SharedWavefunction wfn, SharedBasisSet auxiliary, Options & options)
- RepulsionEnergyOEPotential (SharedWavefunction wfn, Options & options)
- virtual void compute (const std::string &oepType) override
- virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, double &v) override
- virtual void print\_header () const override

# **Additional Inherited Members**

# 9.21.1 Detailed Description

Generalized One-Electron Potential for Pauli repulsion energy calculations.

Contains the following OEP types:

#### 9.21.2 Member Function Documentation

9.21.2.1 void RepulsionEnergyOEPotential::compute\_3D ( const std::string & oepType, const double & x, const double & y, const double & v ) [override], [virtual]

Compute value of potential in point x, y, z and save at v

Implements oepdev::OEPotential.

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

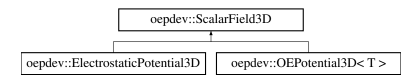
- · oepdev/liboep/oep.h
- · oepdev/liboep/oep.cc

# 9.22 oepdev::ScalarField3D Class Reference

Scalar field in 3D space. Abstract base.

#include <space3d.h>

Inheritance diagram for oepdev::ScalarField3D:



# **Public Member Functions**

- ScalarField3D (const int &np, const double &radius, const double &cx, const double &cy, const double &cz)
- ScalarField3D (const int &np, const double &pad, psi::SharedWavefunction wfn, psi::Options &options)
- ScalarField3D (const int &nx, const int &ny, const int &nz, const double &px, const double &py, const double &px, std::shared\_ptr< psi::Wavefunction > wfn, psi::Options &options)
- virtual ∼ScalarField3D ()

Destructor.

virtual int npoints () const

Get the number of points at which the scalar field is defined.

- · virtual std::shared ptr
  - < PointsCollection3D > points\_collection () const

Get the collection of points.

- · virtual std::shared ptr
  - < psi::Matrix > data () const

Get the data matrix in a form  $\{[x, y, z, f(x, y, z)]\}$ .

- virtual std::shared\_ptr
  - < psi::Wavefunction > wfn () const

Get the wavefunction.

virtual bool is\_computed () const

Get the information if data is already computed or not.

virtual void compute ()

Compute the scalar field in each point from the point collection.

• virtual double compute\_xyz (const double &x, const double &y, const double &z)=0

Compute a value of scalar field at point (x, y, z)

• virtual void write cube file (const std::string &name)

Write the cube file (only for Cube collections, otherwise does nothing)

virtual void print () const =0

Print information of the object to Psi4 output.

#### Static Public Member Functions

- static shared\_ptr< ScalarField3D > build (const std::string &type, const int &np, const double &radius, const double &cx, const double &cy, const double &cz)
- static shared\_ptr< ScalarField3D > **build** (const std::string &type, const int &np, const double &pad, psi::SharedWavefunction wfn, psi::Options &options)
- static shared\_ptr< ScalarField3D > build (const std::string &type, const int &nx, const int &ny, const int &nz, const double &px, const double &py, const double &px, psi::SharedWavefunction wfn, psi::Options &options)

# **Protected Attributes**

```
    std::shared ptr
```

< PointsCollection3D > pointsCollection\_

Collection of points at which the scalar field is to be computed.

std::shared\_ptr< psi::Matrix > data\_

The data matrix in a form  $\{[x, y, z, f(x, y, z)]\}$ .

- std::shared ptr
  - < psi::Wavefunction > wfn

Wavefunction.

psi::Matrix geom\_

Geometry of a molecule.

- · std::shared ptr
  - < psi::IntegralFactory > fact\_

Integral factory.

std::shared ptr< psi::Matrix > pot

Matrix of potential one-electron integrals.

- std::shared ptr
  - < psi::OneBodyAOInt > oneInt

One-electron integral shared pointer.

std::shared\_ptr< PotentialInt > potInt\_

One-electron potential shared pointer.

 std::shared\_ptr< psi::BasisSet > primary\_ Basis set.

int nbf

Number of basis functions.

bool isComputed\_

Has data already computed?

# 9.22.1 Detailed Description

Scalar field in 3D space. Abstract base.

Create scalar field defined at points distributed randomly or as an ordered g09 cube-like collection. Currently implemented scalar fields are:

- Electrostatic potential computes electrostatic potential (requires wavefunction)
- Template of generic classes compute custom scalar fields (requires generic object that is able to compute the field in 3D space)

Note: Always create instances by using static factory methods build.

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

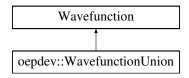
- oepdev/libutil/space3d.h
- · oepdev/libutil/space3d.cc

# 9.23 oepdev::WavefunctionUnion Class Reference

Union of two Wavefunction objects.

#include <wavefunction\_union.h>

Inheritance diagram for oepdev::WavefunctionUnion:



#### **Public Member Functions**

WavefunctionUnion (SharedWavefunction ref\_wfn, Options & options)

Constructor.

virtual ∼WavefunctionUnion ()

Destructor.

• virtual double compute\_energy ()

Compute Energy (now blank)

virtual double nuclear\_repulsion\_interaction\_energy ()

Compute Nuclear Repulsion Energy between unions.

· void localize\_orbitals ()

Localize Molecular Orbitals.

void transform\_integrals ()

Transform Integrals (2- and 4-index transformations)

- int **I\_nmo** (int n) const
- int I\_nso (int n) const
- int I\_nbf (int n) const
- int I ndocc (int n) const
- int I\_nvir (int n) const
- int I noffs ao (int n) const
- int **I\_nalpha** (int n) const
- int I\_nbeta (int n) const
- · double I\_energy (int n) const
- SharedMolecule I\_molecule (int n) const
- SharedBasisSet I primary (int n) const
- · SharedBasisSet I\_auxiliary (int n) const
- SharedWavefunction I\_wfn (int n) const
- SharedMOSpace I\_mospace (int n, const std::string &label) const
- SharedLocalizer I\_localizer (int n) const
- SharedIntegralTransform integrals (void) const
- · bool has\_localized\_orbitals (void) const
- · SharedBasisSet primary (void) const
- void print\_header (void)
- void print\_mo\_integrals (void)

#### **Protected Attributes**

int nlsolatedMolecules

Number of isolated molecules.

SharedWavefunction dimer\_wavefunction\_

The wavefunction for a dimer (electrons relaxed in the field of monomers)

· SharedIntegralTransform integrals\_

Integral transform object (2- and 4-index transformations)

bool hasLocalizedOrbitals

whether orbitals of the union were localized (or not)

std::vector< SharedMolecule > I\_molecule\_

List of molecules.

std::vector< SharedBasisSet > I primary

List of primary basis functions per molecule.

std::vector< SharedBasisSet > I\_auxiliary\_

List of auxiliary basis functions per molecule.

std::vector< SharedWavefunction > I\_wfn\_

List of original isolated wavefunctions (electrons unrelaxed)

std::vector< std::string > I\_name\_

List of names of isolated wavefunctions.

std::vector< int > l\_nbf\_

List of basis function numbers per molecule.

std::vector< int > I nmo

List of numbers of molecular orbitals (MO's) per molecule.

std::vector< int > l\_nso\_

List of numbers of SO's per molecule.

std::vector< int > I ndocc

List of numbers of doubly occupied orbitals per molecule.

std::vector< int > I nvir

List of numbers of virtual orbitals per molecule.

```
std::vector< int > l_noffs_ao_
```

List of basis set offsets per molecule.

std::vector< double > I energy

List of energies of isolated wavefunctions.

std::vector< double > l\_efzc\_

List of frozen-core energies per isolated wavefunction.

std::vector< bool > I density fitted

List of information per wfn whether it was obtained using DF or not.

std::vector< int > I nalpha

List of numbers of alpha electrons per isolated wavefunction.

std::vector< int > I nbeta

List of numbers of beta electrons per isolated wavefunction.

std::vector< int > I nfrzc

List of numbers of frozen-core orbitals per isolated molecule.

std::vector< SharedLocalizer > I localizer

List of orbital localizers.

std::vector< std::map< const</li>

std::string, SharedMOSpace > > I\_mospace\_

List of dictionaries of MO spaces.

#### 9.23.1 Detailed Description

Union of two Wavefunction objects.

The WavefunctionUnion is the union of two unperturbed Wavefunctions.

#### Notes:

- 1. Works only for C1 symmetry! Therefore this->nirrep() = 1.
- 2. Does not set reference\_wavefunction\_
- 3. Sets oeprop\_ for the union of uncoupled molecules
- 4. Performs Hadamard sums on H\_, Fa\_, Da\_, Ca\_ and S\_ based on uncoupled wavefunctions.
- 5. Since it is based on shallow copy of the original Wavefunction, it **changes** contents of this wavefunction. Reallocate and copy if you want to keep the original wavefunction.

# Warnings:

- 1. Gradients, Hessians and frequencies are not touched, hence they are wrong!
- 2. Lagrangian (if present) is not touched, hence its **wrong**!

The following variables are *shallow* copies of variables inside the Wavefunction object, that is created for the *whole* molecule cluster:

- basissets\_(DF/RI/F12/etc basis sets)\_
- basisset\_(ORBITAL basis set)
- sobasisset\_ (Primary basis set for SO integrals)
- AO2SO\_ (AO2SO conversion matrix (AO in rows, SO in cols)
- molecule\_(Molecule that this wavefunction is run on)
- options\_(Options object)

- psio\_(PSI file access variables)
- integral\_(Integral factory)
- factory\_ (Matrix factory for creating standard sized matrices)
- memory\_ (How much memory you have access to)
- nalpha\_, nbeta\_ (Total alpha and beta electrons)
- nfrzc\_ (Total frozen core orbitals)
- doccpi\_ (Number of doubly occupied per irrep)
- soccpi\_ (Number of singly occupied per irrep)
- frzcpi\_ (Number of frozen core per irrep)
- frzvpi\_ (Number of frozen virtuals per irrep)
- nalphapi\_ (Number of alpha electrons per irrep)
- nbetapi\_ (Number of beta electrons per irrep)
- nsopi\_ (Number of so per irrep)
- nmopi\_ (Number of mo per irrep)
- nso (Total number of SOs)
- nmo\_ (Total number of MOs)
- nirrep\_ (Number of irreps; must be equal to 1 due to symmetry reasons)
- same\_a\_b\_dens\_ and same\_a\_b\_orbs\_ The rest is altered so that the Wavefunction parameters reflect a cluster of non-interacting (uncoupled, isolated, unrelaxed) molecular electron densities.

# 9.23.2 Constructor & Destructor Documentation

9.23.2.1 oepdev::WavefunctionUnion::WavefunctionUnion ( SharedWavefunction ref wfn, Options & options )

#### Constructor.

Provide wavefunction with molecule containing at least 2 fragments.

#### **Parameters**

ref_wfn	- reference wavefunction
options	- Psi4 options

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

- · oepdev/libutil/wavefunction union.h
- · oepdev/libutil/wavefunction\_union.cc

# Index

AllAOIntegralsIterator	solve_scf, 17
oepdev::AllAOIntegralsIterator, 20	oepdev::AllAOIntegralsIterator, 19
AllAOShellCombinationsIterator	AllAOIntegralsIterator, 20
oepdev::AllAOShellCombinationsIterator, 21	index, 20
	oepdev::AllAOShellCombinationsIterator, 20
build	AllAOShellCombinationsIterator, 21
oepdev::OEPDevSolver, 33	compute_shell, 22
oepdev::OEPotential, 36	oepdev::CPHF, 22
oepdev::Points3Dlterator, 40, 42	CPHF, 23
oepdev::PointsCollection3D, 44	oepdev::CubePoints3DIterator, 24
CPHF	oepdev::CubePointsCollection3D, 25
oepdev::CPHF, 23	oepdev::DIISManager, 25
compute	compute, 26
oepdev::DIISManager, 26	DIISManager, 26
compute_3D	put, 26
oepdev::EETCouplingOEPotential, 27	update, 26
oepdev::ElectrostaticEnergyOEPotential, 28	oepdev::EETCouplingOEPotential, 26
oepdev::OEPotential, 36	compute_3D, 27
oepdev::RepulsionEnergyOEPotential, 49	oepdev::ESPSolver, 30
compute_benchmark	ESPSolver, 31
oepdev::ElectrostaticEnergySolver, 29	oepdev::ElectrostaticEnergyOEPotential, 27
oepdev::OEPDevSolver, 33	compute_3D, 28
compute_oep_based	oepdev::ElectrostaticEnergySolver, 28
oepdev::ElectrostaticEnergySolver, 29	compute_benchmark, 29
oepdev::OEPDevSolver, 33	compute_oep_based, 29
compute_shell	oepdev::ElectrostaticPotential3D, 29
oepdev::AllAOShellCombinationsIterator, 22	oepdev::OEPDevSolver, 32
create_superfunctional	build, 33
oepdev, 17	compute_benchmark, 33
	compute_oep_based, 33
DIISManager	OEPDevSolver, 33
oepdev::DIISManager, 26	oepdev::OEPotential, 33
ESPSolver	build, 36
	compute_3D, 36
oepdev::ESPSolver, 31 extract monomer	OEPotential, 35
<del>_</del>	write_cube, 36
oepdev, 17	oepdev::OEPotential3D
index	OEPotential3D, 38
oepdev::AllAOIntegralsIterator, 20	oepdev::OEPotential3D $<$ T $>$ , 36
	oepdev::Points3DIterator, 39
OEPDevSolver	build, 40, 42
oepdev::OEPDevSolver, 33	Points3DIterator, 40
OEPotential	oepdev::Points3DIterator::Point, 38
oepdev::OEPotential, 35	oepdev::PointsCollection3D, 42
OEPotential3D	build, 44
oepdev::OEPotential3D, 38	PointsCollection3D, 43
oepdev, 15	oepdev::PotentialInt, 44
create_superfunctional, 17	PotentialInt, 45, 46
extract monomer, 17	set charge field, 46

56 INDEX

```
oepdev::RandomPoints3DIterator, 46
oepdev::RandomPointsCollection3D, 48
oepdev::RepulsionEnergyOEPotential, 48
    compute_3D, 49
oepdev::ScalarField3D, 49
oepdev::WavefunctionUnion, 51
    WavefunctionUnion, 54
Points3DIterator
    oepdev::Points3DIterator, 40
PointsCollection3D
    oepdev::PointsCollection3D, 43
PotentialInt
    oepdev::PotentialInt, 45, 46
psi, 17
put
    oepdev::DIISManager, 26
set_charge_field
    oepdev::PotentialInt, 46
solve_scf
    oepdev, 17
update
    oepdev::DIISManager, 26
WavefunctionUnion
    oepdev::WavefunctionUnion, 54
write_cube
    oepdev::OEPotential, 36
```