## OEP-Dev 1.0.2-alpha

Novel methods for Quantum Chemistry of Extended Molecular Aggregates

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## **Contents**

1	Mair	n Page		1
2	Intro	ductio	n	3
	2.1	Resea	rch Project Methodology	4
	2.2	Expec	ted Impact on the Development of Science, Civilization and Society	4
	2.3	The O	EPDev Code	5
3	OEP	) Design	n.	7
	3.1	OEP C	Classes	7
		3.1.1	Structure of possible OEP-based expressions and their unification	7
4	List	of One-	-Electron Potentals	9
	4.1	Electro	ostatic Energy OEP's	9
	4.2	Pauli F	Repulsion OEP's	9
		4.2.1	First-order contribution in overlap matrix expansion	10
		4.2.2	Second-order contribution in overlap matrix expansion	10
	4.3	Charge	e-Transfer Energy OEP's	10
	4.4	Excito	nic Energy Transfer OEP's	10
		4.4.1	ET contributions.	10
		4.4.2	HT contributions.	11
		4.4.3	CT contributions.	11
	4.5	Full H	Interaction OEP's	11
5	Den	sity-fitt	ing Specialized for OEP's	13
	5.1	Fitting	in Complete Space	13
	5.2	Fitting	in Incomplete Space	14
6	Impl	emente	ed Models	17
	6.1	Target	Properties	17

iv CONTENTS

10	Mod	ule Inde	ay.					33
9	Lice	nse						31
			8.4.1.2	Abstract Base	 	 	 	 29
			8.4.1.1	Implementing OEP Types	 	 	 	 28
		8.4.1	Drafting a	n OEP Subclass	 	 	 	 27
	8.4	Develo	ping OEP's	5	 	 	 	 27
			8.3.3.1	OEPDevSolver	 	 	 	 26
		8.3.3	OEPDev	Solver Module	 	 	 	 26
			8.3.2.3	GenEffFrag	 	 	 	 26
			8.3.2.2	GenEffParFactory	 	 	 	 26
			8.3.2.1	GenEffPar	 	 	 	 26
		8.3.2	GEFP Mo	odule	 	 	 	 26
			8.3.1.2	GeneralizedDensityFit .	 	 	 	 26
			8.3.1.1	OEPPotential	 	 	 	 26
		8.3.1	OEP Mod	lule	 	 	 	 26
	8.3	OEPD	ev Classes	: Overview	 	 	 	 26
		8.2.2						25
		8.2.1		tine				25
	8.2			ructure				24
		8.1.2		ion				24
	· ·	8.1.1		Psi4				23
•	8.1				 	 	 	 23
8	Adv	anced L	Isage					23
	7.7	Use O	oject-Orien	ted Programming	 	 	 	 22
	7.6	Track t	iming wher	evaluating the code	 	 	 	 21
	7.5	Namin	g convention	ons	 	 	 	 21
	7.4	Docum	enting the	code	 	 	 	 20
	7.3	Enviro	nmental va	riables	 	 	 	 20
	7.2	Heade	r files in lib	raries	 	 	 	 19
	7.1	Main ro	outine and	libraries	 	 	 	 19
7	Con	tributin	g to OEP-I	Dev				19
	6.2	Target,	Benchmar	rk and Competing Models	 	 	 	 17

CONTENTS

	10.1	Modules	33
11	Nam	espace Index	35
		Namespace List	35
12		archical Index	37
	12.1	Class Hierarchy	37
13	Class	s Index	39
		Class List	39
14	File I		43
	14.1	File List	43
15	Mod	ule Documentation	45
	15.1	The Generalized One-Electron Potentials Library	45
		15.1.1 Detailed Description	45
	15.2	The OEPDev Solver Library	46
		15.2.1 Detailed Description	46
	15.3	The Generalized Effective Fragment Potentials Library	47
		15.3.1 Detailed Description	48
	15.4	The Integral Package Library	49
		15.4.1 Detailed Description	50
		15.4.2 Hermite Operators	51
		15.4.2.1 Polynomial Expansions as Hermite Series	51
		15.4.3 One-Body Integrals over Hermite Functions	52
		15.4.4 Two-Body Integrals over Hermite Functions	53
		15.4.5 The R(N,L,M) Coefficients	53
		15.4.6 Function Documentation	54
		15.4.6.1 d_N_n1_n2()	54
		15.4.6.2 make_mdh_D1_coeff()	54
		15.4.6.3 make_mdh_D2_coeff()	55
		15.4.6.4 make_mdh_D2_coeff_explicit_recursion()	55
		15.4.6.5 make_mdh_D3_coeff()	56
		15.4.6.6 make_mdh_R_coeff()	57
	15.5	The Three-Dimensional Vector Fields Library	59
		15.5.1 Detailed Description	60

vi CONTENTS

		15.5.2	Function	Documen	tation .					 	 	 			60
			15.5.2.1	OEPoten	tial3D()	[1/2]				 	 	 			60
			15.5.2.2	OEPoten	tial3D()	[2/2]				 	 	 			61
	15.6	The De	ensity Fund	ctional The	eory Libr	ary .				 	 	 			62
	15.7	The OE	EPDev Util	ities						 	 	 			63
		15.7.1	Detailed	Descriptio	n					 	 	 			65
		15.7.2	Function	Documen	tation .					 	 	 			65
			15.7.2.1	average_	moment	()				 	 	 			65
			15.7.2.2	create_sı	Jperfunc	tional()				 	 	 			65
			15.7.2.3	extract_m	nonomer	·()				 	 	 			66
			15.7.2.4	solve_scf	()					 	 	 			66
	15.8	The OE	EPDev Tes	ting Platfo	orm Libra	ary				 	 	 			68
		15.8.1	Detailed	Descriptio	n					 	 	 			68
16	Nom	00000	Documei	station											69
10		•	Namespa		nnoo										69
	10.1		Detailed												74
	16.2		nespace F	•											7 <del>4</del> 74
	10.2	•	Detailed												74
			Function	•											75
		10.2.2	16.2.2.1	oepdev()											75 75
				read_opti											75
			10.2.2.2	Toda_opti	0110()				•	 	 	 	•	•	, 0
17	Clas	s Docui	mentation	1											77
	17.1	oepdev	:::ABCD S	truct Refe	rence .					 	 	 			77
		17.1.1	Detailed	Descriptio	n					 	 	 			77
	17.2	oepdev	:::AbInitioF	PolarGEFa	ctory Cla	ass Re	feren	ce .		 	 	 			77
		17.2.1	Detailed	Descriptio	n					 	 	 			78
	17.3	oepdev	::AllAOInt	egralsItera	ator_2 Cla	ass Re	eferen	ce .		 	 	 			79
		17.3.1	Detailed	Descriptio	n					 	 	 			79
		17.3.2	Construc	tor & Dest	ructor D	ocume	ntatio	n		 	 	 			79
			17.3.2.1	AllAOInte	gralsIte	rator_2	() [1/	2] .		 	 	 			80
			17.3.2.2	AllAOInte	gralsIte	rator_2	<b>()</b> [2/	2] .		 	 	 			80
		17.3.3	Member	Function [	Documer	ntation				 	 	 			80
			17.3.3.1	index()						 	 	 			80
	17.4	oepdev	::AllAOInt	egralsItera	ator_4 Cla	ass Re	eferen	ce .		 	 	 			80

CONTENTS

17.4.1 Detailed Description	81
17.4.2 Constructor & Destructor Documentation	81
17.4.2.1 AllAOIntegralsIterator_4() [1/2]	82
17.4.2.2 AllAOIntegralsIterator_4() [2/2]	82
17.4.3 Member Function Documentation	82
17.4.3.1 index()	82
17.5 oepdev::AllAOShellCombinationsIterator_2 Class Reference	82
17.5.1 Detailed Description	83
17.5.2 Constructor & Destructor Documentation	83
17.5.2.1 AllAOShellCombinationsIterator_2() [1/5]	84
17.5.2.2 AllAOShellCombinationsIterator_2() [2/5]	84
17.5.2.3 AllAOShellCombinationsIterator_2() [3/5]	84
17.5.2.4 AllAOShellCombinationsIterator_2() [4/5]	84
17.5.2.5 AllAOShellCombinationsIterator_2() [5/5]	84
17.5.3 Member Function Documentation	85
17.5.3.1 compute_shell()	85
17.6 oepdev::AllAOShellCombinationsIterator_4 Class Reference	85
17.6.1 Detailed Description	86
17.6.2 Constructor & Destructor Documentation	86
17.6.2.1 AllAOShellCombinationsIterator_4() [1/5]	86
17.6.2.2 AllAOShellCombinationsIterator_4() [2/5]	87
17.6.2.3 AllAOShellCombinationsIterator_4() [3/5]	87
17.6.2.4 AllAOShellCombinationsIterator_4() [4/5]	87
17.6.2.5 AllAOShellCombinationsIterator_4() [5/5]	87
17.6.3 Member Function Documentation	87
17.6.3.1 compute_shell()	88
17.7 oepdev::AOIntegralsIterator Class Reference	88
17.7.1 Detailed Description	89
17.7.2 Member Function Documentation	89
<b>17.7.2.1</b> build() [1/2]	89
<b>17.7.2.2</b> build() [2/2]	90
17.8 oepdev::CAMM Class Reference	90
17.8.1 Detailed Description	90
17.9 oepdev::ChargeTransferEnergyOEPotential Class Reference	91
17.9.1 Detailed Description	92

VIII CONTENTS

17.10oepdev::ChargeTransferEnergySolver Class Reference	92
17.10.1 Detailed Description	93
17.10.2 Member Function Documentation	95
17.10.2.1 compute_benchmark()	95
17.10.2.2 compute_oep_based()	95
17.11oepdev::CPHF Class Reference	96
17.11.1 Detailed Description	98
17.11.2 Constructor & Destructor Documentation	98
17.11.2.1 CPHF()	99
17.12oepdev::CubePoints3DIterator Class Reference	99
17.12.1 Detailed Description	00
17.13oepdev::CubePointsCollection3D Class Reference	00
17.13.1 Detailed Description	01
17.14oepdev::DIISManager Class Reference	01
17.14.1 Detailed Description	01
17.14.2 Constructor & Destructor Documentation	01
17.14.2.1 DIISManager()	01
17.14.3 Member Function Documentation	02
17.14.3.1 compute()	02
17.14.3.2 put()	02
17.14.3.3 update()	02
17.15oepdev::DMTPole Class Reference	03
17.15.1 Detailed Description	07
17.15.2 Constructor & Destructor Documentation	07
17.15.2.1 DMTPole()	07
17.15.3 Member Function Documentation	07
17.15.3.1 build()	80
17.15.3.2 energy()	80
17.15.3.3 potential()	09
17.15.3.4 recenter()	09
17.15.4 Friends And Related Function Documentation	10
17.15.4.1 MultipoleConvergence	10
17.16oepdev::DoubleGeneralizedDensityFit Class Reference	10
17.16.1 Detailed Description	11
17.16.2 Determination of the OEP matrix	11

CONTENTS

17.16.2.1 Theory behind the double GDF scheme
17.16.3 Member Function Documentation
17.16.3.1 compute()
17.17oepdev::EETCouplingOEPotential Class Reference
17.17.1 Detailed Description
17.18oepdev::ElectrostaticEnergyOEPotential Class Reference
17.18.1 Detailed Description
17.19oepdev::ElectrostaticEnergySolver Class Reference
17.19.1 Detailed Description
17.19.2 Member Function Documentation
17.19.2.1 compute_benchmark()
17.19.2.2 compute_oep_based()
17.20oepdev::ElectrostaticPotential3D Class Reference
17.20.1 Detailed Description
17.21oepdev::ERI_1_1 Class Reference
17.21.1 Detailed Description
17.21.2 Implementation
17.22oepdev::ERI_2_2 Class Reference
17.22.1 Detailed Description
17.22.2 Implementation
17.23oepdev::ERI_3_1 Class Reference
17.23.1 Detailed Description
17.23.2 Implementation
17.24oepdev::ESPSolver Class Reference
17.24.1 Detailed Description
17.24.2 Constructor & Destructor Documentation
17.24.2.1 ESPSolver() [1/2]
17.24.2.2 ESPSolver() [2/2]
17.25oepdev::FFAbInitioPolarGEFactory Class Reference
17.25.1 Detailed Description
17.26oepdev::Field3D Class Reference
17.26.1 Detailed Description
17.26.2 Constructor & Destructor Documentation
17.26.2.1 Field3D()
17.26.3 Member Function Documentation

CONTENTS

17.26.3.1 build() [1/2]
17.26.3.2 build() [2/2]
17.27oepdev::Fourier9 Struct Reference
17.27.1 Detailed Description
17.28oepdev::GenEffFrag Class Reference
17.28.1 Detailed Description
17.28.2 Member Function Documentation
17.28.2.1 susceptibility() [1/3]
17.28.2.2 susceptibility() [2/3]
<b>17.28.2.3 susceptibility()</b> [3/3]
17.29oepdev::GenEffPar Class Reference
17.29.1 Detailed Description
17.29.2 Member Function Documentation
17.29.2.1 allocate()
17.29.2.2 compute_density_matrix() [1/4]
17.29.2.3 compute_density_matrix() [2/4]
17.29.2.4 compute_density_matrix() [3/4]
17.29.2.5 compute_density_matrix() [4/4]
17.29.2.6 set_susceptibility()
17.29.2.7 susceptibility() [1/3]
17.29.2.8 susceptibility() [2/3]
17.29.2.9 susceptibility() [3/3]
17.30oepdev::GenEffParFactory Class Reference
17.30.1 Detailed Description
17.30.2 Member Function Documentation
17.30.2.1 build()
17.31oepdev::GeneralizedDensityFit Class Reference
17.31.1 Detailed Description
17.31.2 Member Function Documentation
17.31.2.1 build() [1/2]
<b>17.31.2.2 build()</b> [2/2]
17.31.2.3 compute()
17.32oepdev::GeneralizedPolarGEFactory Class Reference
17.32.1 Detailed Description
17.33oepdev::IntegralFactory Class Reference

CONTENTS xi

17.33.1 Detailed Description	155
17.34oepdev::LinearGradientNonUniformEFieldPolarGEFactory Class Reference	155
17.34.1 Detailed Description	156
17.35oepdev::LinearNonUniformEFieldPolarGEFactory Class Reference	156
17.35.1 Detailed Description	157
17.36oepdev::LinearUniformEFieldPolarGEFactory Class Reference	157
17.36.1 Detailed Description	158
17.37oepdev::MultipoleConvergence Class Reference	158
17.37.1 Detailed Description	159
17.37.2 Member Enumeration Documentation	160
17.37.2.1 ConvergenceLevel	160
17.37.2.2 Property	160
17.37.3 Constructor & Destructor Documentation	160
17.37.3.1 MultipoleConvergence()	161
17.37.4 Member Function Documentation	161
17.37.4.1 compute()	161
17.37.4.2 level()	161
17.38oepdev::NonUniformEFieldPolarGEFactory Class Reference	162
17.38.1 Detailed Description	162
17.39oepdev::OEPDevSolver Class Reference	162
17.39.1 Detailed Description	163
17.39.2 Constructor & Destructor Documentation	164
17.39.2.1 OEPDevSolver()	164
17.39.3 Member Function Documentation	164
17.39.3.1 build()	164
17.39.3.2 compute_benchmark()	165
17.39.3.3 compute_oep_based()	165
17.40oepdev::OEPotential Class Reference	165
17.40.1 Detailed Description	168
17.40.2 Constructor & Destructor Documentation	168
<b>17.40.2.1 OEPotential()</b> [1/2]	168
<b>17.40.2.2 OEPotential()</b> [2/2]	168
17.40.3 Member Function Documentation	169
<b>17.40.3.1 build()</b> [1/2]	169
<b>17.40.3.2 build()</b> [2/2]	169

xii CONTENTS

17.40.3.3 make_oeps3d()	169
17.41oepdev::OEPotential3D $<$ T $>$ Class Template Reference	170
17.41.1 Detailed Description	171
17.42oepdev::OEPType Struct Reference	171
17.43oepdev::PerturbCharges Struct Reference	172
17.43.1 Detailed Description	172
17.44oepdev::Points3Dlterator::Point Struct Reference	172
17.45oepdev::Points3DIterator Class Reference	173
17.45.1 Detailed Description	174
17.45.2 Constructor & Destructor Documentation	174
17.45.2.1 Points3DIterator()	174
17.45.3 Member Function Documentation	175
<b>17.45.3.1 build()</b> [1/3]	175
<b>17.45.3.2 build()</b> [2/3]	175
<b>17.45.3.3 build()</b> [3/3]	176
17.46oepdev::PointsCollection3D Class Reference	176
17.46.1 Detailed Description	177
17.46.2 Constructor & Destructor Documentation	178
17.46.2.1 PointsCollection3D()	178
17.46.3 Member Function Documentation	178
<b>17.46.3.1 build()</b> [1/3]	178
<b>17.46.3.2 build()</b> [2/3]	178
<b>17.46.3.3 build()</b> [3/3]	179
17.47oepdev::PolarGEFactory Class Reference	179
17.47.1 Detailed Description	181
17.48oepdev::PotentialInt Class Reference	181
17.48.1 Constructor & Destructor Documentation	181
17.48.1.1 PotentialInt() [1/3]	182
<b>17.48.1.2 PotentialInt()</b> [2/3]	182
<b>17.48.1.3 PotentialInt()</b> [3/3]	182
17.48.2 Member Function Documentation	183
17.48.2.1 set_charge_field()	183
$17.49 oep dev:: Quadratic Gradient Non Uniform EField Polar GEF actory\ Class\ Reference\ .$	183
17.49.1 Detailed Description	184
17.50oepdev::QuadraticNonUniformEFieldPolarGEFactory Class Reference	185

CONTENTS xiii

17.50.1 Detailed Description	185
17.51oepdev::QuadraticUniformEFieldPolarGEFactory Class Reference	186
17.51.1 Detailed Description	186
17.52oepdev::RandomPoints3DIterator Class Reference	187
17.52.1 Detailed Description	188
17.53oepdev::RandomPointsCollection3D Class Reference	188
17.53.1 Detailed Description	189
17.54oepdev::RepulsionEnergyOEPotential Class Reference	189
17.54.1 Detailed Description	189
17.55oepdev::RepulsionEnergySolver Class Reference	190
17.55.1 Detailed Description	190
17.55.2 Member Function Documentation	195
17.55.2.1 compute_benchmark()	195
17.55.2.2 compute_oep_based()	195
17.56oepdev::RHFPerturbed Class Reference	196
17.56.1 Detailed Description	197
17.57oepdev::ShellCombinationsIterator Class Reference	198
17.57.1 Detailed Description	200
17.57.2 Constructor & Destructor Documentation	200
17.57.2.1 ShellCombinationsIterator()	200
17.57.3 Member Function Documentation	200
17.57.3.1 ao_iterator()	200
<b>17.57.3.2 build()</b> [1/2]	201
<b>17.57.3.3 build()</b> [2/2]	201
17.57.3.4 compute_shell()	202
17.58oepdev::SingleGeneralizedDensityFit Class Reference	202
17.58.1 Detailed Description	202
17.58.2 Determination of the OEP matrix	203
17.58.3 Member Function Documentation	203
17.58.3.1 compute()	203
17.59oepdev::GeneralizedPolarGEFactory::StatisticalSet Struct Reference	203
17.60oepdev::test::Test Class Reference	204
17.61oepdev::TwoBodyAOInt Class Reference	206
17.61.1 Member Function Documentation	206
<b>17.61.1.1 compute()</b> [1/2]	206

XIV CONTENTS

	<b>17.61.1.2 compute()</b> [2/2]	207
17	7.62oepdev::TwoElectronInt Class Reference	207
	17.62.1 Detailed Description	209
	17.62.2 Member Function Documentation	209
	17.62.2.1 compute_shell()	209
17	7.63oepdev::UniformEFieldPolarGEFactory Class Reference	209
	17.63.1 Detailed Description	210
17	7.64oepdev::UnitaryOptimizer Class Reference	210
	17.64.1 Detailed Description	214
	17.64.2 Constructor & Destructor Documentation	215
	17.64.2.1 UnitaryOptimizer() [1/3]	215
	<b>17.64.2.2 UnitaryOptimizer()</b> [2/3]	216
	<b>17.64.2.3 UnitaryOptimizer()</b> [3/3]	216
17	7.65oepdev::UnitaryOptimizer_4_2 Class Reference	216
	17.65.1 Detailed Description	219
	17.65.2 Constructor & Destructor Documentation	221
	17.65.2.1 UnitaryOptimizer_4_2() [1/2]	221
	17.65.2.2 UnitaryOptimizer_4_2() [2/2]	221
17	7.66oepdev::UnitaryTransformedMOPolarGEFactory Class Reference	222
	17.66.1 Detailed Description	222
17	7.67oepdev::WavefunctionUnion Class Reference	223
	17.67.1 Detailed Description	226
	17.67.2 Constructor & Destructor Documentation	228
	17.67.2.1 WavefunctionUnion() [1/2]	228
	<b>17.67.2.2 WavefunctionUnion()</b> [2/2]	228
	17.67.3 Member Function Documentation	229
	17.67.3.1 Ca₋subset()	229
	17.67.3.2 Cb_subset()	229
18 Fi	le Documentation	231
18	3.1 include/oepdev_files.h File Reference	231
18	3.2 include/oepdev_options.h File Reference	231
18	3.3 main.cc File Reference	232
18	3.4 oepdev/lib3d/dmtp.h File Reference	232
18	B.5 oepdev/lib3d/esp.h File Reference	233

CONTENTS xv

	18.6 oepdev/libgefp/gefp.h File Reference	234
	18.7 oepdev/libints/eri.h File Reference	235
	18.8 oepdev/libints/recurr.h File Reference	236
	18.9 oepdev/liboep/oep.h File Reference	237
	18.10oepdev/liboep/oep_gdf.h File Reference	238
	18.11oepdev/libpsi/integral.h File Reference	239
	18.12oepdev/libpsi/potential.h File Reference	239
	18.13oepdev/libsolver/solver.h File Reference	240
	18.14oepdev/libtest/test.h File Reference	240
	18.15oepdev/libutil/diis.h File Reference	241
	18.16oepdev/libutil/integrals_iter.h File Reference	242
	18.17oepdev/libutil/scf_perturb.h File Reference	243
	18.18oepdev/libutil/unitary_optimizer.h File Reference	243
	18.19oepdev/libutil/util.h File Reference	244
	18.20oepdev/libutil/wavefunction_union.h File Reference	245
19	Example Documentation	247
	19.1 example_cphf.cc	247
	19.2 example_integrals_iter.cc	247
	19.3 example_scf_perturb.cc	248
Inc	lex	249

## **Main Page**

### **OEP-Dev**

Generalized One-Electron Potentials: Development Platform.

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#### 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 charge-transfer interaction energy (Project I)
- 3. the polarization of electronic density (Project III)
- 4. the excitation energy transfer couplings (Project I)

against reference solutions (exact or other approximations).

#### Places to go:

- OEP-Dev Code
- Current Issues

This wikipages will be updated soon.

#### References

[1] B. Błasiak, "One-Particle Density Matrix Polarization Susceptibility Tensors", *J. Chem. Phys.* **149**, 164115 (2018)

2 Main Page

### Introduction

Exploring biological phenomena at molecular scale is oftentimes indispensable to develop new drugs and intelligent materials.

Most of relevant system properties are affected by intermolecular interactions with nearby environment such as solvent or closely bound electronic chromophores. Studying such molecular aggregates requires rigorous and accurate quantum chemistry methods, the cost of which grows very fast with the number of electrons. Despite many methodologies have been devised to describe energetic and dynamical properties of **extended molecular systems** efficiently and accurately, there exist particularly difficult cases in which modelling is still challenging:

- · describing electronic transitions in solution or
- when coupled with other electronic transition via resonance energy transfer,
- performing molecular dynamics at very high level of theory including dynamic electron correlation,
- vibrational frequency calculations of particular normal mode in condensed phases

and so on. The reason behind (sometimes prohibitively) high costs of fully *ab initio* calculations in the above areas is the complexity of mathematical models often based on wave functions rather then (conceptually more straightforward) electronic densities. On the other hand, it has been pointed out before that the one-electron density distributions are of particular importance in chemistry. It can be thus utilized as a means of developing a general model that re-expresses the physics of intermolecular interactions in terms of effective one-electron functions that are easier to handle in practice.

This Project will focus on finding a unified way to simplify various equations of Quantum Chemistry of extended molecular systems, i.e., molecular aggregates such as interacting chromophores and molecules solvated by water and other solvents. Indeed, one of the important difficulties encountered in Quantum Chemistry of large systems is the need of evaluation of special kind of numbers known as *electron repulsion integrals*, or in short, ERI's. In a typical calculation, the amount of ERI's can be as high as tens or even hundreds of millions (!) that unfortunately prevents from application of conventional methods when the number of particles in question is too large. In the Project, the complicated expressions involving ERI's shall be greatly simplified to reduce the computational costs as much as possible while introducing no or minor approximations to the original theories.

4 Introduction

### 2.1 Research Project Methodology

In this Project the new theoretical protocol based on the one-electron effective potentials (OEP's) is developed. The main principle is to rewrite arbitrary sum of functions f of electron repulsion integrals (ERI's) by defining OEP's according to the following general prescription:

$$\begin{split} &\sum_{f} f\left[\left(\phi_{i}^{A}\phi_{j}^{A}||\phi_{k}^{B}\phi_{l}^{B}\right)\right] = \left(\phi_{i}^{A}|v_{kl}^{B}|\phi_{j}^{A}\right) \rightarrow \text{ point charge or density fitting} \\ &\sum_{f} f\left[\left(\phi_{i}^{A}\phi_{j}^{B}||\phi_{k}^{B}\phi_{l}^{B}\right)\right] = \left(\phi_{i}^{A}|v_{kl}^{B}|\phi_{j}^{B}\right) \rightarrow \text{ density fitting,} \end{split}$$

where A and B denote different molecules and  $\phi_i$  is the i-th molecular orbital or basis function. Here,  $v_{kl}^B$  denotes the List of One-Electron Potentals ab initio "OEP matrix element". The technique described above will be applied to simplify expressions for

- short-range excitation energy transfer couplings between chlorophyll subunits of reaction centres in photosynthesis
- · Pauli interaction repulsion energy
- · charge-transfer interaction energy
- electric field-induced charge density polarization of molecules.

The above developments might be used in fragment-based *ab initio* molecular dynamics protocols of new generation.

# 2.2 Expected Impact on the Development of Science, Civilization and Society

The proposed OEP's are expected to significantly develop the fragment-based methods that are widely used in physical chemistry and modelling of biologically important systems. Owing to universality of OEP's, they could find applications in many branches of chemical science: non-empirical\* molecular dynamics, short-range resonance energy transfer in photosynthesis, electronic and vibrational solvatochromism, multidimensional spectroscopy and so on. In particular:

- the OEP-based models of Pauli repulsion energy and charge-transfer (CT) energy could be used to improve the computational performance of the second generation effective fragment potential method (EFP2). At present, the CT term is very time consuming and due to this reasons it is not used in most of applications of EFP2 to perform molecular dynamics simulations.
- the OEP-based model of EET couplings could significantly improve modelling of energy transfer in the light harvesting complexes. At present, short-range phenomena (Dexter mechanisms of EET) are very difficult to efficiently and quantitatively asses when performing statistical averaging and applying to large molecular aggregates. Such Dexter effects could be computed by using OEP's in much more efficient manner without loosing high accuracy of parent TDFI-TI method.

2.3 The OEPDev Code 5

 the density matrix polarization (DMS) tensors could be used in new generation fragmentbased ab initio molecular dynamics protocols that rigorously take into consideration electron correlation effects.

Therefore, it is strongly believed that the OEP's could have an indirect impact on the design of novel drugs and materials for industry.

### 2.3 The OEPDev Code

To pursue the above challenges in the field of computational quantum chemistry of extended molecular aggregates, the OEPDev platform is developed. Accurate and efficient *ab initio* models based on OEP's are implemented in the OEPDev code, along with the state-of-the-art benchmark and competiting methods. Written entirely in C++, OEPDev is a plugin to Psi4 quantum chemistry package. Therefore, compilation and running the OEPDev code is straightforward and follows the API interface similar to the one used in Psi4 with just a few specific programing conventions. The detailed discussion about using the OEPDev code can be found in advanced usage section.

6 Introduction

## **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:

Туре	Matrix Element	Comment
Type 1	$\left(I \hat{v}^A J ight)$	$I \in A, J \in B$
Type 2	$\left(J \hat{v}^A L ight)$	$J,L\in B$

8 OEP Design.

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
$\left(\mu \hat{v}^{A[\mu]} \sigma ight)$	$\sum_{\iota \in A} v_{\mu \iota}^A S_{\iota \sigma}$	$\sum_{lpha\in A}q_lpha^{A[\mu]}V_{\mu\sigma}^{(lpha)}$
$\boxed{\left(i \hat{v}^{A[i]} j\right)}$	$\sum_{\iota \in A} v_{i\iota}^A S_{\iota j}$	$\sum_{lpha \in A} q_{lpha}^{A[i]} V_{ij}^{(lpha)}$
$\left(j \hat{v}^{A[i]} l ight)$	$\sum_{\iota\kappa\in A} S_{j\iota} v_{\iota\kappa}^{A[i]} S_{\kappa l}$	$\sum_{lpha\in A}q_lpha^{A[i]}V_{jl}^{(lpha)}$

In the formulae above, the OEP-part (stored by OEP instances) and the Solver-part (to be computed by the Solver) are separated. 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.

Another possibility is to use psi::Matrix objects, instead of psi4::Tensor objects, possibly putting them into a std::vector container in case there is more than two axes.

### **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.

Note

Add here a table with all the OEP types along with their symbols used in the OEPDev code (e.g., Murrell.etal-S1 etc).

### 4.1 Electrostatic Energy OEP's

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

3D form:

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

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 Charge-Transfer Energy OEP's

To be added here!

### 4.4 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.

#### 4.4.1 ET contributions.

3D forms:

$$v(\mathbf{r})_{1}^{A[\mu]} = -C_{\mu L}^{*} \sum_{x \in A} \frac{Z_{x}}{|\mathbf{r} - \mathbf{r}_{x}|} + \sum_{v \kappa \in A} \left\{ C_{\mu L}^{*} D_{v \kappa} - \frac{1}{2} C_{v L}^{*} D_{\mu \kappa} \right\} \int d\mathbf{r}' \frac{\phi_{v}^{*}(\mathbf{r}') \phi_{\kappa}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}$$

$$v(\mathbf{r})_{2}^{A[\mu]} = C_{\kappa H} \sum_{v \kappa \in A} \left\{ 2C_{v L}^{*} C_{\mu H}^{*} - C_{v H}^{*} C_{\mu L}^{*} \right\} \int d\mathbf{r}' \frac{\phi_{v}^{*}(\mathbf{r}') \phi_{\kappa}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}$$

$$v(\mathbf{r})_{3}^{A[\mu]} = v(\mathbf{r})_{1}^{A[\mu]} + v(\mathbf{r})_{1}^{A[\mu]}$$

Matrix forms:

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

#### 4.4.2 HT contributions.

Do be derived.

### 4.4.3 CT contributions.

To be derived.

### 4.5 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.

List of	One-Electron	Potentals

12

## **Density-fitting Specialized for OEP's**

To get the ab-initio representation of a OEP, one can use a procedure similar to the typical density fitting or resolution of identity, both of which are nowadays widely used to compute electron-repulsion integrals (ERI's) more efficiently.

### 5.1 Fitting in Complete Space

An arbitrary one-electron potential of molecule A acting on any state vector associated with molecule A can be expanded in an auxiliary space centered on A as

$$v|i) = \sum_{\xi \eta} v|\xi) [\mathbf{S}^{-1}]_{\xi \eta} (\eta|i)$$

under the necessary assumption that the auxiliary basis set is *complete*. In a special case when the basis set is orthogonal (e.g., molecular orbitals) the above relation simplifies to

$$v|i) = \sum_{\xi} v|\xi)(\xi|i)$$

It can be easily shown that the above general and exact expansion can be obtained by performing a density fitting in the complete space. We expand the LHS of the first equation on this page in a series of the auxiliary basis functions scaled by the undetermined expansion coefficients:

$$v|i) = \sum_{\xi} G_{i\xi}|\xi)$$

which we shall refer here as to the matrix form of the OEP operator. By constructing the least-squares objective function

$$Z[\{G_{\xi}^{(i)}\}] = \int d\mathbf{r}_1 \left[ v(\mathbf{r}_1)\phi_i(\mathbf{r}_1) - \sum_{\xi} G_{\xi}^{(i)}\phi_{\xi}(\mathbf{r}_1) \right]^2$$

and requiring that

$$rac{\partial Z[\{G_{\xi}^{(i)}\}]}{\partial G_{\mu}^{(i)}}=0$$
 for all  $\mu$ 

we find the coefficients  $G_{\xi}^{(i)}$  to be

$$\mathbf{G}^{(i)} = \mathbf{v}^{(i)} \cdot \mathbf{S}^{-1}$$

where

$$v_{\eta}^{(i)} = (\eta | vi)$$
  
 $S_{\eta \xi} = (\eta | \xi)$ 

or explitictly

$$G_{i\xi} = \sum_{m{\eta}} [\mathbf{S}^{-1}]_{\xi m{\eta}} (m{\eta} | v | i)$$

identical to what we obtained from application of the resolution of identity in space spanned by non-orthogonal complete set of basis vectors.

Since matrix elements of an OEP operator in auxiliary space can be computed in the same way as the matrix elements with any other basis function, one can formally write the following identity

$$(X|v|i) = \sum_{\xi \eta} S_{X\xi}[\mathbf{S}^{-1}]_{\xi \eta}(\eta|v|i)$$

where X is an arbitrary orbital. When the other orbital does not belong to molecule A but to the (changing) environment, it is straightforward to compute the resulting matrix element, which is simply given as

$$(j_{\in B}|v^A|i_{\in A}) = \sum_{\xi} S_{j\xi} G_{i\xi}$$

where *i* denotes the other (environmental) basis function.

In the above equation, the OEP-part (fragment parameters for molecule *A* only) and the Solver-part (subject to be computed by solver on the fly) are separated. This then forms a basis for fragment-based approach to solve Quantum Chemistry problems related to the extended molecular aggregates.

### 5.2 Fitting in Incomplete Space

Density fitting scheme from previous section has practical disadvantage of a nearly-complete basis set being usually very large (spanned by large amount of basis set vectors). Any non-complete basis set won't work in the previous example. Since most of basis sets used in quantum chemistry do not form a complete set, it is beneficial to design a modified scheme in which it is possible to obtain the **effective** matrix elements of the OEP operator in a **incomplete** auxiliary space. This can be achieved by minimizing the following objective function

$$Z[\{G_{\xi}^{(i)}\}] = \iint d\mathbf{r}_1 d\mathbf{r}_2 \frac{\left[v(\mathbf{r}_1)\phi_i(\mathbf{r}_1) - \sum_{\xi} G_{\xi}^{(i)} \phi_{\xi}(\mathbf{r}_1)\right] \left[v(\mathbf{r}_2)\phi_i(\mathbf{r}_2) - \sum_{\xi} G_{\eta}^{(i)} \phi_{\eta}(\mathbf{r}_1)\right]}{|\mathbf{r}_1 - \mathbf{r}_2|}$$

Thus requesting that

$$rac{\partial Z[\{G_{\xi}^{(i)}\}]}{\partial G_{u}^{(i)}}=0$$
 for all  $\mu$ 

we find the coefficients  $G_{\xi}^{(i)}$  to be

$$\mathbf{G}^{(i)} = \mathbf{b}^{(i)} \cdot \mathbf{A}^{-1}$$

where

$$b_{\eta}^{(i)} = (\eta || vi)$$
  
 $A_{\eta \xi} = (\eta || \xi)$ 

The symbol || is to denote the operator  $r_{12}^{-1}$  and double integration over  $\mathbf{r}_1$  and  $\mathbf{r}_2$ . Thus, it is clear that in order to use this generalized density fitting scheme one must to compute two-centre electron repulsion integrals (implemented in oepdev::ERI\_1\_1) as well as four-centre asymmetric electron repulsion integrals of the type  $(\alpha\beta\gamma||\eta)$  (implemented in oepdev::ERI\_3\_1).

Density-fitting Specialized for OEP's

16

## **Implemented Models**

### **6.1 Target Properties**

Detailed list of models which is to be implemented in the OEPDev project is given below:

**Table 1.** Models subject to be implemented and analyzed within oep-dev.

Pauli energy	Induction energy	EET Coupling
EFP2-Pauli	EFP2-Induced Dipoles	TrCAMM
Murrel et al.'s theory	Density Susceptibility	OEP-ET/HT
OEP-Murrel et al.'s		TDFI-TI
		FED
Exact (Stone's)	Exact (incl. CT)	Exact (ESD)

### 6.2 Target, Benchmark and Competing Models

The target models introduced in the Project shall be tested against the following benchmarks and compared with the following state-of-the-art models:

**Table 2.** Target models vs benchmarks and competitor models.

Target Model	Benchmarks	Competing Model
OEP-Murrel et al.'s	Murrel et al.'s	EFP2-Pauli
	Exact (Stone's)	
OEP-ET/HT + TrCAMM	Exact (ESD)	TDFI-TI
	FED	FED
	TDFI-TI	
Density Susceptibility	Exact (incl. CT)	EFP2-Induced Dipoles

## **Contributing to OEP-Dev**

OepDev is a plugin to Psi4.

Therefore it should follow the programming etiquette of Psi4. Also, oep-dev has additional programming tips to make the code more versatile and easy in further development. Here, I emphasise on most important aspects regarding the **programming rules**.

#### 7.1 Main routine and libraries

Oep-dev has only *one* source file in the plugin base directory, i.e., main.cc. This is the main driver routine that handles the functionality of the whole OEP testing platform: specifies options for Psi4 input file and implements test routines based on the options. Include files directly related to main.cc are stored in the include directory, where only header files are present. Options are specified in include/oepdev\_options.h whereas macros and defines in include/oepdev\_files.h. Other sources are stored in MODULE/libNAME\* directories where NAME is the name of the library with sources and header files, whereas MODULE is the directory of the oep-dev module.

#### Things to remember:

- 1. **No other sources in base directory.** It is not permitted to place any new source or other files in the plugin base directory (i.e., where main.cc resides).
- 2. Sources in library directories. Any additional source code has to be placed in oepdev/libNAME\* directory (either existing one or a new one; in the latter case remember to add the new \*.cc files to CMakeLists.txt in the plugin base directory.
- 3. **Miscellanea in special directories.** If you want to add additional documentation, put it in the doc directory. If you want to add graphics, put it in the images directory.

### 7.2 Header files in libraries

Header files are handy in obtaining a quick glimpse of the functionality within certain library. Each library directory should contain at least one header file in oep-dev. However, header files can be problematic if not managed properly.

Things to remember:

1. **Header preprocessor variable**. Define the preprocessor variable specyfying the existence of include of the particular header file. The format of such is

```
#ifndef MODULE_LIBRARY_HEADER_h
#define MODULE_LIBRARY_HEADER_h
// rest of your code goes here
#endif // MODULE_LIBRARY_HEADER_h
```

Last line is the **end** of the header file. The preprocessor variables represents the directory tree <code>oepdev/MODULE/LIBRARY/HEADER.h</code> structure (where <code>oepdev</code> is the base plugin directory). MODULE is the plugin module name (e.g. <code>oepdev</code>, the name of the module directory) <code>LIBRARY</code> is the name of the library (e.g. <code>libutil</code>, should be the same as library directory name) <code>HEADER</code> is the name of the header in library directory (e.g. <code>diis</code> for <code>diis.h</code> header file)

2. **Set module namespace**. To prevent naming clashes with other modules and with Psi4 it is important to operate in separate namespace (e.g. for a module).

```
namespace MODULE {
// your code goes here
} // EndNameSpace MODULE
```

For instance, all classes and functions in <code>oepdev</code> module are implemented within the namespace of the same label. Considering addition of other local namespaces within a module can also be useful in certain cases.

### 7.3 Environmental variables

Defining the set of intrinsic environmental variables can help in code management and conditional compilation. The oep-dev environmental variables are defined in include/oepdev\_files.h
file. Remember also about psi4 environmental variables defined in psi4/psifiles.h
header. As a rule, the oep-dev environmental variable should have the following format:

```
OEPDEV_XXXX
```

where XXXX is the descriptive name of variable.

### 7.4 Documenting the code

Code has to be documented (at best at a time it is being created). The place for documentation is always in header files. Additional documentation can be also placed in source files. Leaving a chunk of code for a production run without documentation is unacceptable.

Use Doxygen style for documentation all the time. Remember that it supports markdown which can make the documentation even more clear and easy to understand. Additionally you can create a nice <code>.rst</code> documentation file for Sphinx program. If you are coding equations, always include formulae in the documentation!

Things to remember:

- 1. **Descriptions of classes, structures, global functions, etc**. Each programming object should have a description.
- Documentation for function arguments and return object. Usage of functions and class methods should be explained by providing the description of all arguments (use \param and \return Doxygen keywords).
- 3. **One-line description of class member variables**. Any class member variable should be preceded by a one-liner documentation (starting from ///).
- 4. **Do not be afraid of long names in the code**. Self-documenting code is a bless!

### 7.5 Naming conventions

Naming is important because it helps to create more readable and clear self-documented code. Some loose suggestions:

- 1. Do not be afraid of long names in the code, but avoid redundancy. Examples of good and bad names: good name: get\_density\_matrix; bad name: get\_matrix. Unless there is only one type of matrix a particular objects can store, matrix is not a good name for a getter method. good name: class Wavefunction, bad name: class WFN good name: int numberOfErrorVectors, bad name: int nvec, bad name: the\_number\_of\_error\_vectors good name: class EFPotential, probably bad name: class EffectiveFragmentPotential. The latter might be understood by some people as a class that inherits from EffectiveFragment class. If it is not the case, compromise between abbreviation and long description is OK.
- 2. **Short names are OK in special situations**. In cases meaning of a particular variable is obvious and it is frequently used in the code locally, it can be named shortly. Examples are: i when iterating no number of occupied orbitals, nv number of virtual orbitals, etc.
- 3. Clumped names for variables and dashed names for functions. Try to distinguish between variable name like sizeOfOEPTypeList and a method name get\_matrix() (neither size\_of\_OEP\_type\_list, nor getMatrix()). This is little bit cosmetics, but helps in managing the code when it grows.
- 4. Class names start from capital letter. However, avoid only capital letters in class names, unless it is obvious. Avoid also dashes in class names (they are reserved for global functions and class methods). Examples: good name: DIISManager, bad name: DIIS. good name: EETCouplingSolver, bad name: EETSolver, very bad: EET.

### 7.6 Track timing when evaluating the code

It is useful to track time elapsed for performing a particular task by a computer. For this, use psi::timer\_on and psi::timer\_off functions defined in psi4/libqt/qt.h. Psi4 always generates the report file timer.dat that contains all the defined timings. For example,

```
#include "psi/libqt/qt.h"
psi::timer_on("OEP E(Paul) Murrell-etal S1 ");
// Your code goes here
psi::timer_off("OEP E(Paul) Murrell-etal S1 ");
```

To maintain the printout in a neat form, the timing associated with the OEPDev code should be generated via misc/python/timing.py utility script.

### 7.7 Use Object-Oriented Programming

Try to organise your creations in objects having special relationships and data structures. Encapsulation helps in producing self-maintaining code and is much easier to use. Use:

- · factory design for creating objects
- container design for designing data structures
- **polymorphysm** when dealing with various flavours of one particular feature in the data structure

*Note:* In Psi4, factories are frequently implemented as static methods of the base classes, for example psi::BasisSet::build static method. It can be followed when building object factories in oep-dev too.

## **Advanced Usage**

This section is addressed for advanced users.

Make sure you have first read the introduction before proceeding.

#### 8.1 Installation

#### 8.1.1 Preparing Psi4

OEPDev is a Psi4 plugin. It requires

- Psi4, at least 1.2 version (git commit 9d4a61c). Has to be modified (see below)
- Eigen3, any newer version

Note

Before compiling, make sure EFP is enabled in CMakeLists.txt (now it is not used in OEPDev but maybe in the future it would).

Recently, Psi4 introduced API visibility management. Only certain Psi4 classes and functions are *exposed* in the core.so library, that is further linked to Psi4 plugin shared library. Due to this reason, not all Psi4 functionalities can be directly used from outside Psi4. In order to access local API of Psi4 (also used in the OEPDev code) slight modification of Psi4 code and concomitant rebuild is necessary.

In order to expose local API used by OEPDev and hidden within Psi4 1.2, two types of small modifications are necessary:

- M1: add PSI\_API macro after required class or function declaration in header file
- M2: add #include "psi4/pragma.h" line at the include section of an appropriate header file

Modification M1 is obligatory for all affected files whereas modification M2 needs to be done only in headers that do not have "psi4/pragma.h" included explicitly or implicitly. The list of Psi4 header files along with the respective changes that need to be done are listed in the table below:

24 Advanced Usage

Psi4 Header File	Psi4 Class	Required Changes
libfunctional/superfunctiona	1 <b>Sh</b> perfunctional	M1
libscf_solver/hf.h	HF	M1
libscf_solver/rhf.h	RHF	M1
libcubeprop/csg.h	CubicScalarGrid	M1
libmints/onebody.h	OneBodyAOInt	M1
libmints/potential.h	PotentialInt	M1
libmints/multipoles.h	MultupoleInt	M1
libmints/multipolesymmetry.h	MultipoleSymmetry	M1
libmints/fjt.h	Taylor_Fjt	M1
libmints/fjt.h	Fjt	M1
libmints/oeprop.h	OEProp	M1, M2
libmints/gshell.h	GaussianShell	M1, M2

To quickly apply these modifications, use the patch files stored in misc/patch directory. Please makes sure to use a proper patch for a chosen Psi4 version.

#### 8.1.2 Compiltation

After all the above changes have been done in Psi4 (followed by its rebuild) compile the OEPDev code by running <code>compile</code> script. Make sure Eigen3 path is set to environment variable <code>EIGEN3\_INCLUDE\_DIR</code> (instructions will appear on the screen). After compilation is successful, run <code>ctest</code> to check if the code works fine.

#### Note

It may happen, that during code development there will be symbol lookup error when importing <code>oepdev.so</code> (in such case OEPDev compiles without error but Python cannot import the module <code>oepdev</code>). In such circumstance, probably there some local Psi4 feature that is needed in OEPDev is not exposed by <code>PSI\_API</code> macro. To fix this, run <code>c++filt[name]</code> where <code>[name]</code> is the mangled undefined symbol. This will show you which Psi4 class or function is not exposed and requires <code>PSI\_API</code> (change M1 and perhaps M2 too). Such change requires Psi4 rebuild and recompilation of OEPDev code. In any case, please contact me and report new undefined symbol (<code>blasiak.bartosz@gmail.com</code>).

#### 8.2 OEPDev Code Structure

As a plugin to Psi4, OEPDev consists of the main.cc file with the plugin main routine, include/oepdev\_options.h specifying the options of the plugin, include/oepdev\_files.h defining all global macros and environmental variables, as well as the oepdev directory. The latter contains the actual OEPDev code that is divided into several subdirectories called modules.

#### 8.2.1 Main Routine

Before the actual OEPDev calculations are started, the wavefunction of the input molecular aggregate is computed by Psi4. See the plugin driver script pymodule.py for more details on how the calculation environment is initialized. Subsequently, one out of four types of target operations can be performed by the program:

- 1. OEP\_BUILD Compute the OEP effective parameters for one molecule.
- 2. DMATPOL Compute the generalized density matrix susceptibility tensors (DMS's) for one molecule.
- 3. SOLVER Perform calculations for a molecular aggregate. As for now, only dimers are handled.
- 4. TEST Perform the testing routine.

The first two modes are single molecule calculations. OEP\_BUILD uses the oepdev::OEPotential::bus static factory to create OEP objects whereas DMATPOL uses the oepdev::GenEffParFactory::build static factory to greate generalized effective fragment parameters (GEFP's) for polarization.

#### Note

In the future, OEP\_BUILD will be handled also by oepdev::GenEffParFactory::build since OEP parameters are part of the GEFP's.

SOLVER requires at least molecular dimer and the oepdev::WavefunctionUnion object (being the Hartree product of the unperturbed monomer wavefunctions) is constructed at the beginning, which is then passed to the oepdev::OEPDevSolver::build static factory. TEST can refer to single- or multiple-molecule calculations, whereby each of the testing routines is listed in the cmake/CTestTestfile.cmake.in file.

#### 8.2.2 Modules

The source code is distributed into directories called modules:

- liboep
- libgefp
- libsolver
- libints
- libpsi
- lib3d
- libutil
- libtest

See Modules for a detailed description of each of the modules.

26 Advanced Usage

#### 8.3 OEPDev Classes: Overview

#### 8.3.1 OEP Module

The OEP module located in oepdev/liboep consists of the following abstract bases:

- oepdev::OEPotential implementing the OEP,
- oepdev::GeneralizedDensityFit implementing the GDF technique.

Each of the bases contains static factory method called build that creates instances of chosen subclasses. The module contains also a structure oepdev::OEPType which is a container storing all the data associated with a particular OEP: type name, dimensions, OEP coefficients and whether is density-fitted or not.

#### 8.3.1.1 OEPPotential

It is a container and computer class of OEP. Among others, the most important public method is oepdev::OEPotential::compute which computes all the OEP's (by iterating over all possible OEP types within a chosen OEP subclass or category). OEP's can be extracted by oepdev::OEPotential::oep method, for instance. From protected attributes, each OEPotential instance stores blocks of the LCAO-MO matrices associated with the occupied (cOcc\_) and virtual (cVir\_) MO's. It also contains the pointers to the primary, auxiliary and intermediate basis sets (primary\_, auxiliary\_ and intermediate\_, accordingly). Usage example:

```
#include "oepdev/liboep/oep.h"
oep = oepdev::OEPotential::build("ELECTROSTATIC ENERGY", wfn, options);
oep->compute();
oep->write_cube("V", "oep_cube_file");
```

So far, four OEPotential subclasses are implemented, from which <code>oepdev::ElectrostaticEnergyOEPotential</code> and <code>oepdev::RepulsionEnergyOEPotential</code> are fully operative, while the rest is under development.

- 8.3.1.2 GeneralizedDensityFit
- 8.3.2 GEFP Module
- 8.3.2.1 GenEffPar
- 8.3.2.2 GenEffParFactory
- 8.3.2.3 GenEffFrag
- 8.3.3 OEPDev Solver Module
- 8.3.3.1 OEPDevSolver

### 8.4 Developing OEP's

OEP's are implemented in a suitable subclass of the oepdev::OEPotential base. Due to the fact that OEP's can be density-based or ESP-based, the classes oepdev::GeneralizedDensityFit as well as oepdev::ESPSolver are usually necessary in the implementations. Handling the one-electron integrals (OEI's) and the two-electron integrals (ERI's) in AO basis is implemented in oepdev::IntegralFactory. In particular, potential integrals evaluated at arbitrary centres can be accessed by using the oepdev::PotentialInt instances. Useful iterators for looping over AO ERI's the oepdev::ShellCombinationsIterator and oepdev::AOIntegralsIterator classes. Transformations of OEI's to MO basis can be easily achieved by transforming AO integral matrices by cocc\_ and cvir\_members of OEPotential instances, e.g., by using the psi::Matrix::doublet or psi::Matrix::triplet static methods. Transformations of ERI's to MO basis can be performed by using the psi4/libtrans/integraltransform.h library.

It is recommended that the implementation of all the new OEP's follows the following steps:

- 1. **Write the class framework.** This includes choosing a proper name of a OEPotential subclass, sketching the constructors and a destructor, and all the necessary methods.
- 2. **Implement OEP types.** Each type of OEP is implemented, including the 3D vector field in case ESP-based OEP's are of use.
- 3. **Update base factory method**. Add appropriate entries in the oepdev::OEPotential::build static factory method.

Below, we shall go through each of these steps separately and discuss them in detail.

#### 8.4.1 Drafting an OEP Subclass

This stage is the design of the overall framework of OEP subclass. The name should end with <code>OEPotential</code> to maintain the convention used so far. The template for the header file definition can be depicted as follows:

```
class SampleOEPotential : public OEPotential
 public:
    // Purely ESP-based OEP's
    SampleOEPotential(SharedWavefunction wfn, Options& options);
    // GDF-based OEP's
   SampleOEPotential (SharedWavefunction wfn, SharedBasisSet auxiliary, SharedBasisSet intermediate,
     Options& options);
    // Necessary destructor
    virtual ~SampleOEPotential();
    // Necessary computer
    virtual void compute (const std::string& oepType) override;
    // Necessary computer
    virtual void compute_3D(const std::string& oepType,
                           const double& x, const double& y, const double& z, std::shared_ptr<psi::Vector>
      & v) override;
    // Necessary printer
```

28 Advanced Usage

```
virtual void print_header() const override;

private:
   // Set defaults - good practice
   void common_init();

   // Auxilary computers - exemplary
   double compute_3D_sample_V(const double& x, const double& y, const double& z);
};
```

The constructors need to call the abstract base constructor and then specialized initializations. It is a good practice to put the specialized common initializers in a separate private method common\_init (which is a convention in Psi4 and is adopted also in OEPDev). For instance, the exemplary constructor is show below:

```
SampleOEPotential::SampleOEPotential(SharedWavefunction wfn,
                                     SharedBasisSet auxiliary, SharedBasisSet intermediate, Options&
     options)
: OEPotential(wfn, auxiliary, intermediate, options)
   common_init();
}
void SampleOEPotential::common_init()
   int n1 = wfn_->Ca_subset("AO", "OCC")->ncol();
   int n2 = auxiliary_->nbf();
  int n3 = wfn_->molecule()->natom();
  SharedMatrix mat1 = std::make_shared<psi::Matrix>("G(S^{-1})", n2, n1);
  SharedMatrix mat_2 = std::make_shared<psi::Matrix>("G(S^{-2})", n3, n1);
  OEPType type_1 = {"Murrell-etal.S1", true , n1, mat_1};
  OEPType type_2 = {"Otto-Ladik.S2" , false, n1, mat_2};
  oepTypes_[type_1.name] = type_1;
  oepTypes_[type_2.name] = type_2;
```

Note that the <code>oepdev::OEPotential::oepTypes\_</code> attribute, which is a <code>std::map</code> of structures <code>oepdev::OEPType</code>, is initialized here. All the OEP types need to be stated in the constructors. Destructors usually call nothing, unless dynamically allocated memory is also of use.

It is also a good practice to already sketch the compute method here by adding certain private computers, like in the example below:

#### 8.4.1.1 Implementing OEP Types

Implementation of the inner body of compute method requires populating the members of oepTypes\_ with data. This means, that for each OEP type there has to be a specific implementation of OEP parameters. GDF-based OEP's need to create the psi::Matrix with OEP

parameters and put them into <code>oepTypes.</code> In the case of ESP-based OEP's <code>compute\_3D</code> method has to be additionally implemented before <code>compute</code> is fully functional. To implement <code>compute\_3D</code>, <code>oepdev::OEPotential::make\_oeps3d</code> method is of high relevance: it creates <code>oepdev::OEPotential3D<T></code> instances, where <code>T</code> is the OEP subclass. These instances are <code>oepdev::Field3D</code> objects that define OEP's in 3D Euclidean space. For example,

```
void SampleOEPotential::compute_otto_ladik_s2()
      // Switch on timer
     psi::timer_on("OEP
                          E(Paul) Otto-Ladik S2
      // Create 3D field, automated through 'make_oeps3d'. Requires 'compute_3D' implementation.
      std::shared_ptr<OEPotential3D<OEPotential>> oeps3d = this->make_oeps3d("Otto-Ladik.S2");
      oeps3d->compute();
      // Perform ESP fit to get OEP effective charges
      ESPSolver esp(oeps3d);
      esp.set_charge_sums(0.5);
      esp.compute();
      // Put the OEP coefficients into 'oepTypes_'
      for (int i=0; i<esp.charges()->nrow(); ++i) {
           for (int o=0; o<oepTypes_["Otto-Ladik.S2"].n; ++o) {</pre>
                oepTypes_["Otto-Ladik.S2"].matrix->set(i, o, esp.charges()->get(i, o));
      }
      // Switch off timer
      psi::timer_off("OEP
                            E(Paul) Otto-Ladik S2
// Necessary implementation for 'make_oeps3d' to work
void SampleOEPotential::compute.3D(const std::string& oepType, const double& x, const double& y, const
     double& z, std::shared_ptr<psi::Vector>& v)
   // Loop over all possibilities for OEP types and exclude illegal names
   if (oepType == "Otto-Ladik.S2") {
       // this computes the actual values of OEP = v(x,y,z) and stores it in 'vec_otto_ladik_s2_'
       this->compute_3D_otto_ladik_s2(x, y, z);
       // Assign final value to the buffer vector
       for (int o = 0; o < oepTypes_["Otto-Ladik.S2"].n; ++o) v->set(o, vec_otto_ladik_s2_[o]);
   else if (oepType == "Murrell-etal.S1" ) \{/* Even if it is not ESP-based OEP, this line is necessary */\}
   else {
      throw psi::PSIEXCEPTION("OEPDEV: Error. Incorrect OEP type specified!\n"); // Safety
}
```

Note that make\_oeps3d is not overridable and is fully defined in the base. Do not call oepdev::OEPotential3D constructors in the OEPotential subclass (it can be done only from the level of the abstract base where all the pointers are dynamically converted to an appropriate data type due to polymorphism)!

#### 8.4.1.2 Abstract Base



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32 License

# **Module Index**

### 10.1 Modules

1.1					1 1
Here	าร ล	list	ot al	။ mဂ	dules

The Generalized One-Electron Potentials Library	45
The OEPDev Solver Library	46
The Generalized Effective Fragment Potentials Library	47
The Integral Package Library	49
The Three-Dimensional Vector Fields Library	59
The Density Functional Theory Library	62
The OEPDev Utilities	63
The OEPDev Testing Platform Library	68

34 **Module Index** 

# Namespace Index

## 11.1 Namespace List

Here	ic a	liet o	f all	documented	namespaces	with	hrief	descriptions
11010	is a	iiot u	ıaıı	aocumented	Halliespaces	VVILII	ושוט	uescriptions

oepdev																
	OEPDev module namespace	)														69
psi																
	Psi4 package namespace															74



## **Hierarchical Index**

## 12.1 Class Hierarchy

This inheritance list is sorted roughly, but not completely, alphabetically:	
oepdev::ABCD	
oepdev::AOIntegralsIterator	88
oepdev::AllAOIntegralsIterator_2	. 79
oepdev::AllAOIntegralsIterator_4	. 80
oepdev::CPHF	96
CubicScalarGrid	
oepdev::CubePointsCollection3D	. 100
oepdev::DIISManager	101
enable_shared_from_this	
oepdev::DMTPole	
oepdev::CAMM	
oepdev::OEPDevSolver	
oepdev::ChargeTransferEnergySolver	
oepdev::ElectrostaticEnergySolver	
oepdev::RepulsionEnergySolver	
oepdev::OEPotential	
oepdev::ChargeTransferEnergyOEPotential	
oepdev::EETCouplingOEPotential	
oepdev::RepulsionEnergyOEPotential	
oepdev::ESPSolver	
oepdev::Field3D	
oepdev::ElectrostaticPotential3D	
oepdev::OEPotential3D< T >	
oepdev::Fourier9	
oepdev::GenEffFrag	
oepdev::GenEffPar	
oepdev::GenEffParFactory	
oepdev::PolarGEFactory	
oopaoviii olai alli aoloi y	. 175

38 Hierarchical Index

oepdev::AbInitioPolarGEFactory	. 77
oepdev::UnitaryTransformedMOPolarGEFactory	. 222
oepdev::FFAbInitioPolarGEFactory	. 127
oepdev::GeneralizedPolarGEFactory	. 148
oepdev::NonUniformEFieldPolarGEFactory	. 162
oepdev::LinearGradientNonUniformEFieldPolarGEFactory	
oepdev::LinearNonUniformEFieldPolarGEFactory	
oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory	
oepdev::QuadraticNonUniformEFieldPolarGEFactory	
oepdev::UniformEFieldPolarGEFactory	
oepdev::LinearUniformEFieldPolarGEFactory	
oepdev::QuadraticUniformEFieldPolarGEFactory	
oepdev::GeneralizedDensityFit	
oepdev::DoubleGeneralizedDensityFit	. 110
oepdev::SingleGeneralizedDensityFit	
IntegralFactory	
oepdev::IntegralFactory	154
oepdev::MultipoleConvergence	
oepdev::OEPType	
oepdev::PerturbCharges	
oepdev::Points3DIterator::Point	
oepdev::Points3DIterator	
oepdev::CubePoints3DIterator	
oepdev::RandomPoints3DIterator	
oepdev::PointsCollection3D	176
oepdev::CubePointsCollection3D	100
oepdev::RandomPointsCollection3D	
PotentialInt	
oepdev::PotentialInt	. 181
RHF	
oepdev::RHFPerturbed	. 196
oepdev::ShellCombinationsIterator	
oepdev::AllAOShellCombinationsIterator_2	. 82
oepdev::AllAOShellCombinationsIterator_4	
oepdev::GeneralizedPolarGEFactory::StatisticalSet	
oepdev::test::Test	
TwoBodyAOInt	201
oepdev::TwoBodyAOInt	206
oepdev::TwoElectronInt	
oepdev::ERI_1_1	
oepdev::ERI_2_2	
oepdev::ERI_3_1	
oepdev::UnitaryOptimizer	
oepdev::UnitaryOptimizer_4_2	
Wavefunction	210
oepdev::WavefunctionUnion	223

# **Class Index**

### 13.1 Class List

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

oepdev::ABCD	
Simple structure to hold the Fourier series expansion coefficients	77
oepdev::AbInitioPolarGEFactory	
Polarization GEFP Factory from First Principles. Hartree-Fock Approximation	77
oepdev::AllAOIntegralsIterator_2	
Loop over all possible ERI within a particular shell doublet	79
oepdev::AllAOIntegralsIterator_4	
Loop over all possible ERI within a particular shell quartet	80
oepdev::AllAOShellCombinationsIterator_2	
Loop over all possible ERI shells in a shell doublet	82
oepdev::AllAOShellCombinationsIterator_4	
Loop over all possible ERI shells in a shell quartet	85
oepdev::AOIntegralsIterator	
Iterator for AO Integrals. Abstract Base	88
oepdev::CAMM	
Cumulative Atomic Multipole Moments	90
oepdev::ChargeTransferEnergyOEPotential	
Generalized One-Electron Potential for Charge-Transfer Interaction Energy	91
oepdev::ChargeTransferEnergySolver	
Compute the Charge-Transfer interaction energy between unperturbed wave-	
functions	92
oepdev::CPHF	
CPHF solver class	96
oepdev::CubePoints3DIterator	
Iterator over a collection of points in 3D space. g09 Cube-like order	99
oepdev::CubePointsCollection3D	
G09 cube-like ordered collection of points in 3D space	00
oepdev::DIISManager	
DIIS manager	01

40 Class Index

oepdev::DMTPole
Distributed Multipole Analysis Container and Computer. Abstract Base 103
oepdev::DoubleGeneralizedDensityFit
Generalized Density Fitting Scheme - Double Fit
oepdev::EETCouplingOEPotential
Generalized One-Electron Potential for EET coupling calculations
oepdev::ElectrostaticEnergyOEPotential
Generalized One-Electron Potential for Electrostatic Energy
oepdev::ElectrostaticEnergySolver
Compute the Coulombic interaction energy between unperturbed wavefunc-
tions
oepdev::ElectrostaticPotential3D
Electrostatic potential of a molecule
oepdev::ERI_1_1
2-centre ERI of the form (a O(2) b) where O(2) = $1/r12$
oepdev::ERI_2_2
4-centre ERI of the form (ab $ O(2) $ cd) where $O(2) = 1/r12 \dots 121$
oepdev::ERI_3_1
4-centre ERI of the form (abc $ O(2) d$ ) where $O(2) = 1/r12 \dots 123$
oepdev::ESPSolver
Charges from Electrostatic Potential (ESP). A solver-type class
oepdev::FFAbInitioPolarGEFactory
Polarization GEFP Factory from First Principles: Finite-Difference Model. Ar-
bitrary level of theory
oepdev::Field3D
General Vector Dield in 3D Space. Abstract base
oepdev::Fourier9
Simple structure to hold the Fourier series expansion coefficients for $N=4$ 132
oepdev::GenEffFrag
Generalized Effective Fragment. Container Class
oepdev::GenEffPar
Generalized Effective Fragment Parameters. Container Class
oepdev::GenEffParFactory
Generalized Effective Fragment Factory. Abstract Base
oepdev::GeneralizedDensityFit
Generalized Density Fitting Scheme. Abstract Base
oepdev::GeneralizedPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization 148
oepdev::IntegralFactory
Extended IntegralFactory for computing integrals
oepdev::LinearGradientNonUniformEFieldPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization 155
oepdev::LinearNonUniformEFieldPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization 156
oepdev::LinearUniformEFieldPolarGEFactory  Polarization GEFP Factory with Least-Squares Parameterization
oepdev::MultipoleConvergence
Multipole Convergence

13.1 Class List

oepdev::NonUniformEFieldPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization 162
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 3D fields
oepdev::OEPType
Container to handle the type of One-Electron Potentials
oepdev::PerturbCharges
Structure to hold perturbing charges
oepdev::Points3DIterator::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::PolarGEFactory
Polarization GEFP Factory. Abstract Base
oepdev::PotentialInt
Computes potential integrals
oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization
oepdev::QuadraticNonUniformEFieldPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization 185
oepdev::QuadraticUniformEFieldPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization 186
oepdev::RandomPoints3DIterator
Iterator over a collection of points in 3D space. Random collection 187
oepdev::RandomPointsCollection3D
Collection of random points in 3D space
oepdev::RepulsionEnergyOEPotential
Generalized One-Electron Potential for Pauli Repulsion Energy
oepdev::RepulsionEnergySolver
Compute the Pauli-Repulsion interaction energy between unperturbed wave-
functions
oepdev::RHFPerturbed
RHF theory under electrostatic perturbation
oepdev::ShellCombinationsIterator
Iterator for Shell Combinations. Abstract Base
oepdev::SingleGeneralizedDensityFit
Generalized Density Fitting Scheme - Single Fit
oepdev::GeneralizedPolarGEFactory::StatisticalSet
A structure to handle statistical data
oepdev::test::Test
Manages test routines
oepdev::TwoBodyAOInt

42 Class Index

oepdev::TwoElectronInt
General Two Electron Integral
oepdev::UniformEFieldPolarGEFactory
Polarization GEFP Factory with Least-Squares Parameterization 209
oepdev::UnitaryOptimizer
Find the optimim unitary matrix of quadratic matrix equation
oepdev::UnitaryOptimizer_4_2
Find the optimim unitary matrix for quartic-quadratic matrix equation with trace 216
oepdev::UnitaryTransformedMOPolarGEFactory
Polarization GEFP Factory with Least-Squares Scaling of MO Space 222
oepdev::WavefunctionUnion
Union of two Wavefunction objects

## File Index

### 14.1 File List

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

main.cc
include/oepdev_files.h
include/oepdev_options.h
include/doxygen/oepdev_manual.h??
include/doxygen/oepdev_modules.h
include/doxygen/oepdev_namespaces.h??
oepdev/lib3d/dmtp.h
oepdev/lib3d/esp.h
oepdev/lib3d/ <b>space3d.h</b>
oepdev/libgefp/gefp.h
oepdev/libints/eri.h
oepdev/libints/recurr.h
oepdev/liboep/oep.h
oepdev/liboep/oep_gdf.h
oepdev/libpsi/integral.h
oepdev/libpsi/potential.h
oepdev/libsolver/solver.h
oepdev/libtest/test.h
oepdev/libutil/cphf.h
oepdev/libutil/diis.h
oepdev/libutil/integrals_iter.h
oepdev/libutil/scf_perturb.h
oepdev/libutil/unitary_optimizer.h
oepdev/libutil/util.h
oepdev/libutil/wavefunction_union.h

File Index 44

### **Module Documentation**

### 15.1 The Generalized One-Electron Potentials Library

Implements the goal of this project: The Generalized One-Electron Potentials (OEP's). You will find here OEP's for computation of Pauli repulsion energy, charge-transfer energy and others. The routines for the generalized density fitting are also implemented here. Located at oepdev/liboep.

#### Classes

struct oepdev::OEPType

Container to handle the type of One-Electron Potentials.

· class oepdev::OEPotential

Generalized One-Electron Potential: Abstract base.

class oepdev::ElectrostaticEnergyOEPotential

Generalized One-Electron Potential for Electrostatic Energy.

class oepdev::RepulsionEnergyOEPotential

Generalized One-Electron Potential for Pauli Repulsion Energy.

class oepdev::ChargeTransferEnergyOEPotential

Generalized One-Electron Potential for Charge-Transfer Interaction Energy.

class oepdev::EETCouplingOEPotential

Generalized One-Electron Potential for EET coupling calculations.

class oepdev::GeneralizedDensityFit

Generalized Density Fitting Scheme. Abstract Base.

class oepdev::SingleGeneralizedDensityFit

Generalized Density Fitting Scheme - Single Fit.

· class oepdev::DoubleGeneralizedDensityFit

Generalized Density Fitting Scheme - Double Fit.

#### 15.1.1 Detailed Description

### 15.2 The OEPDev Solver Library

Implementations of various solvers for molecular properties as a functions of unperturbed monomeric wavefunctions. This is the place all target OEP-based models are implemented and compared with benchmark and competitor models. Located at oepdev/libsolver.

#### **Classes**

- class oepdev::OEPDevSolver
  - Solver of properties of molecular aggregates. Abstract base.
- class oepdev::ElectrostaticEnergySolver
  - Compute the Coulombic interaction energy between unperturbed wavefunctions.
- class oepdev::RepulsionEnergySolver
  - Compute the Pauli-Repulsion interaction energy between unperturbed wavefunctions.
- class oepdev::ChargeTransferEnergySolver
  - Compute the Charge-Transfer interaction energy between unperturbed wavefunctions.

#### 15.2.1 Detailed Description

### 15.3 The Generalized Effective Fragment Potentials Library

Implements the GEFP method, the far goal of the OEPDev project. Here you will find the containers for GEFP parameters, the density matrix susceptibility tensors and GEFP solvers. Located at oepdev/libgefp.

#### Classes

· class oepdev::GenEffPar

Generalized Effective Fragment Parameters. Container Class.

class oepdev::GenEffFrag

Generalized Effective Fragment. Container Class.

class oepdev::GenEffParFactory

Generalized Effective Fragment Factory. Abstract Base.

class oepdev::PolarGEFactory

Polarization GEFP Factory. Abstract Base.

class oepdev::AbInitioPolarGEFactory

Polarization GEFP Factory from First Principles. Hartree-Fock Approximation.

class oepdev::FFAbInitioPolarGEFactory

Polarization GEFP Factory from First Principles: Finite-Difference Model. Arbitrary level of theory.

class oepdev::GeneralizedPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::UniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::NonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::LinearUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::QuadraticUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::LinearNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::QuadraticNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::LinearGradientNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class oepdev::UnitaryTransformedMOPolarGEFactory

Polarization GEFP Factory with Least-Squares Scaling of MO Space.

15.3.1 Detailed Description

### 15.4 The Integral Package Library

Implementations of various two-, three- or four-centre two-body electron repulsion integrals via utilizing the McMurchie-Davidson recurrence scheme. Located at <code>oepdev/libints</code> and <code>oepdev/libpsi</code>.

#### **Classes**

class oepdev::TwoElectronInt

General Two Electron Integral.

class oepdev::ERI\_1\_1

2-centre ERI of the form (a|O(2)|b) where O(2) = 1/r12.

class oepdev::ERI\_2\_2

4-centre ERI of the form (ab|O(2)|cd) where O(2) = 1/r12.

class oepdev::ERI\_3\_1

4-centre ERI of the form (abc|O(2)|d) where O(2) = 1/r12.

- class oepdev::TwoBodyAOInt
- class oepdev::IntegralFactory

Extended IntegralFactory for computing integrals.

· class oepdev::PotentialInt

Computes potential integrals.

#### **Macros**

#define D1\_INDEX(x, i, n) ((81\*(x))+(9\*(i))+(n))

Get the index of McMurchie-Davidson-Hermite D1 coefficient stored in the mdh\_buffer\_, that is attributed to the x Cartesian coordinate from angular momentum i of function 1, and the Hermite index n.

#define D2\_INDEX(x, i, j, n) ((1377\*(x))+(153\*(i))+(17\*(j))+(n))

Get the index of McMurchie-Davidson-Hermite D2 coefficient stored in the mdh\_buffer\_, that is attributed to the x Cartesian coordinate from angular momenta i, j of function 1 and 2, and the Hermite index n.

#define D3\_INDEX(x, i, j, k, n) ((18225\*(x))+(2025\*(i))+(225\*(j))+(25\*(k))+(n))

Get the index of McMurchie-Davidson-Hermite D3 coefficient stored in the mdh\_buffer\_, that is attributed to the x Cartesian coordinate from angular momenta i, j and k of function 1, 2 and 3, and the Hermite index n.

#define R\_INDEX(n, I, m, j) ((14739\*(n))+(867\*(I))+(51\*(m))+(j))

Get the index of McMurchie-Davidson R coefficient stored in the  $mdh\_buffer\_R\_$  from angular momenta n, I and m and the Boys index j.

#### **Functions**

- double oepdev::d\_N\_n1\_n2 (int N, int n1, int n2, double PA, double PB, double aP)
   Compute McMurchie-Davidson-Hermite (MDH) coefficient for binomial expansion.
- void oepdev::make\_mdh\_D1\_coeff (int n1, double aPd, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for monomial expansion.

void oepdev::make\_mdh\_D2\_coeff (int n1, int n2, double aPd, double \*PA, double \*PB, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for binomial expansion.

void oepdev::make\_mdh\_D3\_coeff (int n1, int n2, int n3, double aPd, double \*PA, double \*PB, double \*PC, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for trinomial expansion.

void oepdev::make\_mdh\_D2\_coeff\_explicit\_recursion (int n1, int n2, double aP, double \*PA, double \*PB, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for binomial expansion by explicit recursion. This function makes the same changes to buffers as oepdev::make\_mdh\_D2\_coeff, but implements it through explicit recursion by calls to oepdev::d\_N\_n1\_n2. Therefore, it is slightly slower. Here for debugging purposes.

void oepdev::make\_mdh\_R\_coeff (int N, int L, int M, double alpha, double a, double b, double c, double \*F, double \*buffer)

Compute the McMurchie-Davidson R coefficients.

#### 15.4.1 Detailed Description

Here, we define the primitive Gaussian type functions (GTO's)

$$\phi_{i}(\mathbf{r}) \equiv x_{A}^{n_{1}} y_{A}^{l_{1}} z_{A}^{m_{1}} e^{-\alpha_{1} r_{A}^{2}}$$

$$\phi_{j}(\mathbf{r}) \equiv x_{B}^{n_{2}} y_{B}^{l_{2}} z_{B}^{m_{2}} e^{-\alpha_{2} r_{B}^{2}}$$

$$\phi_{k}(\mathbf{r}) \equiv x_{C}^{n_{3}} y_{C}^{l_{3}} z_{C}^{m_{3}} e^{-\alpha_{3} r_{C}^{2}}$$

in which  $\mathbf{r}_A \equiv \mathbf{r} - \mathbf{A}$  and so on.  $\mathbf{A}$  is the centre of the GTO,  $\alpha_1$  its exponent, whereas  $n_1, l_1, m_1$  the Cartesian angular momenta, with the total angular momentum  $\theta_1 = n_1 + l_1 + m_1$ .

In OEPDev implementations, the following definition shall be in use:

$$\mathbf{P} \equiv \frac{\alpha_1 \mathbf{A} + \alpha_2 \mathbf{B}}{\alpha_1 + \alpha_2}$$

$$\mathbf{Q} \equiv \frac{\alpha_3 \mathbf{C} + \alpha_4 \mathbf{D}}{\alpha_3 + \alpha_4}$$

$$\mathbf{R} \equiv \frac{\alpha_1 \mathbf{A} + \alpha_2 \mathbf{B} + \alpha_3 \mathbf{C}}{\alpha_1 + \alpha_2 + \alpha_3}$$

$$\alpha_P \equiv \alpha_1 + \alpha_2$$

$$\alpha_Q \equiv \alpha_3 + \alpha_4$$

$$\alpha_R \equiv \alpha_1 + \alpha_2 + \alpha_3$$

The unnormalized products of primitive GTO's are denoted here as

$$[ij] \equiv \phi_i(\mathbf{r})\phi_j(\mathbf{r})$$
  
 $[ijk] \equiv \phi_i(\mathbf{r})\phi_j(\mathbf{r})\phi_k(\mathbf{r})$ 

#### 15.4.2 Hermite Operators

It is convenient to define

$$\Lambda_j(x_P; \alpha_P) \equiv \left(\frac{\partial}{\partial P_x}\right)^j = \alpha_P^{j/2} H_j(\sqrt{\alpha_P} x_P)$$

where  $H_j(x)$  is the Hermite polynomial of order j evaluated at x. Introduction of the above Hermite operator can be used by invoking the recurrence relationship due to Hermite polynomial properties:

$$x_A \Lambda_j(x_P; \alpha_P) = j\Lambda_{j-1} + |\mathbf{P} - \mathbf{A}|_x \Lambda_j + \frac{1}{2\alpha_P} \Lambda_{j+1}$$

This can be directly used to derive very useful McMurchie-Davidson-Hermite coefficients as expansion coefficients of the polynomial expansions.

#### 15.4.2.1 Polynomial Expansions as Hermite Series

By using the previous relation, it is possible to express the following expansions in Hermite series:

$$x_A^{n_1} = \sum_{N=0}^{n_1} d_N^{n_1} \Lambda_N(x_A; \alpha_A)$$

$$x_A^{n_1} x_B^{n_2} = \sum_{N=0}^{n_1+n_2} d_N^{n_1 n_2} \Lambda_N(x_P; \alpha_P)$$

$$x_A^{n_1} x_B^{n_2} x_C^{n_3} = \sum_{N=0}^{n_1+n_2+n_3} d_N^{n_1 n_2 n_3} \Lambda_N(x_R; \alpha_R)$$

The recurrence relationships can be easily found and they read

$$d_N^{n_1+1} = \frac{1}{2\alpha_A} d_{N-1}^{n_1} + (N+1)d_{N+1}^{n_1}$$

as well as

$$\begin{split} d_N^{n_1+1,n_2} &= \frac{1}{2\alpha_P} d_{N-1}^{n_1n_2} + |\mathbf{P} - \mathbf{A}|_x d_N^{n_1n_2} + (N+1) d_{N+1}^{n_1n_2} \\ d_N^{n_1,n_2+1} &= \frac{1}{2\alpha_P} d_{N-1}^{n_1n_2} + |\mathbf{P} - \mathbf{B}|_x d_N^{n_1n_2} + (N+1) d_{N+1}^{n_1n_2} \end{split}$$

and

$$d_{N}^{n_{1}+1,n_{2},n_{3}} = \frac{1}{2\alpha_{R}} d_{N-1}^{n_{1}n_{2}n_{3}} + |\mathbf{R} - \mathbf{A}|_{x} d_{N}^{n_{1}n_{2}n_{3}} + (N+1) d_{N+1}^{n_{1}n_{2}n_{3}}$$

$$d_{N}^{n_{1},n_{2}+1,n_{3}} = \frac{1}{2\alpha_{R}} d_{N-1}^{n_{1}n_{2}n_{3}} + |\mathbf{R} - \mathbf{B}|_{x} d_{N}^{n_{1}n_{2}n_{3}} + (N+1) d_{N+1}^{n_{1}n_{2}n_{3}}$$

$$d_{N}^{n_{1},n_{2},n_{3}+1} = \frac{1}{2\alpha_{R}} d_{N-1}^{n_{1}n_{2}n_{3}} + |\mathbf{R} - \mathbf{C}|_{x} d_{N}^{n_{1}n_{2}n_{3}} + (N+1) d_{N+1}^{n_{1}n_{2}n_{3}}$$

respectively. The first elements are given by

$$d_0^0 = 1$$
$$d_0^{00} = 1$$
$$d_0^{000} = 1$$

By using the above formalisms, it is strightforward to express the doublet of primitive GTO's as

$$[ij] = E_{ij} \sum_{N=0}^{n_1+n_2} \sum_{L=0}^{l_1+l_2} \sum_{M=0}^{m_1+m_2} d_N^{n_1n_2} d_L^{l_1l_2} d_M^{m_1m_2} \Lambda_N(x_P) \Lambda_L(y_P) \Lambda_M(z_P) e^{-\alpha_P r_P^2}$$

Analogously, the triplet of primitive GTO's is given by

$$[ijk] = E_{ijk} \sum_{N=0}^{n_1 + n_2 + n_3} \sum_{L=0}^{l_1 + l_2 + l_3} \sum_{M=0}^{m_1 + m_2 + m_3} d_N^{n_1 n_2 n_3} d_L^{l_1 l_2 l_3} d_M^{m_1 m_2 m_3} \Lambda_N(x_R) \Lambda_L(y_R) \Lambda_M(z_R) e^{-\alpha_R r_R^2}$$

The multiplicative constants are given by

$$E_{ij}(\alpha_1, \alpha_2) = \exp\left[-\frac{\alpha_1 \alpha_2}{\alpha_1 + \alpha_2} |\mathbf{A} - \mathbf{B}|^2\right]$$

$$E_{ijk}(\alpha_1, \alpha_2, \alpha_3) = \exp\left[-\frac{\alpha_1 \alpha_2}{\alpha_1 + \alpha_2} |\mathbf{A} - \mathbf{B}|^2\right] \exp\left[-\frac{(\alpha_1 + \alpha_2) \alpha_3}{\alpha_1 + \alpha_2 + \alpha_3} |\mathbf{P} - \mathbf{C}|^2\right]$$

#### 15.4.3 One-Body Integrals over Hermite Functions

The fundamental Hermite integrals that appear during computations of any kind of one-body integrals over GTO's are as follows

$$[NLM|\Theta(1)] \equiv \int d\mathbf{r}_1 \Theta(\mathbf{r}_1) \Lambda_N(x_{1P}; \alpha_P) \Lambda_L(y_{1P}; \alpha_P) \Lambda_M(z_{1P}; \alpha_P) e^{-\alpha_P r_{1P}^2}$$

It immediately follows that the overlap, dipole, quadrupole and potential integrals are given as

$$[NLM|1] = \delta_{N0}\delta_{L0}\delta_{M0} \left(\frac{\pi}{\alpha_P}\right)^{3/2}$$

$$[NLM|x_C] = [\delta_{N1} + |\mathbf{PC}|_x \delta_{N0}] \,\delta_{L0}\delta_{M0} \left(\frac{\pi}{\alpha_P}\right)^{3/2}$$

$$[NLM|x_C^2] = \left[2\delta_{N2} + 2|\mathbf{PC}|_x \delta_{N1} + \left(|\mathbf{PC}|_x^2 + \frac{1}{2\alpha_P}\right)\delta_{N0}\right] \delta_{L0}\delta_{M0} \left(\frac{\pi}{\alpha_P}\right)^{3/2}$$

$$[NLM|x_C y_C] = (\delta_{N1} + |\mathbf{PC}|_x \delta_{N0}) \left(\delta_{L1} + |\mathbf{PC}|_y \delta_{L0}\right) \delta_{M0} \left(\frac{\pi}{\alpha_P}\right)^{3/2}$$

$$[NLM|r_C^{-1}] = \frac{2\pi}{\alpha_P} R_{NLM}$$

The coefficients  $R_{NLM}$  are discussed in separate section below.

#### 15.4.4 Two-Body Integrals over Hermite Functions

The fundamental Hermite integrals that appear during computations of any kind of two-electron integrals over GTO's are as follows

$$[N_{1}L_{2}M_{2}|N_{2}L_{2}M_{2}] \equiv \iint d\mathbf{r}_{1}d\mathbf{r}_{2}\Lambda_{N_{1}}(x_{1P};\alpha_{P})\Lambda_{L_{1}}(y_{1P};\alpha_{P})\Lambda_{M_{1}}(z_{1P};\alpha_{P})\Lambda_{N_{2}}(x_{2Q};\alpha_{Q})\Lambda_{L_{2}}(y_{2Q};\alpha_{Q})\Lambda_{M_{2}}(z_{2Q};\alpha_{Q})\Lambda_{M_{2}}($$

The above formula dramatically reduces to the following

$$[N_1L_2M_2|N_2L_2M_2] = \lambda (-)^{N_2+L_2+M_2}R_{N_1+N_2,L_1+L_2,M_1+M_2}$$

with

$$\lambda \equiv rac{2\pi^{5/2}}{lpha_Plpha_Q\sqrt{lpha_P+lpha_Q}}$$

To compute the  $R_{N1+N2,L1+L2,M1+M2}$  coefficients, the parameter T is given by

$$T = \frac{\alpha_P \alpha_Q}{\alpha_P + \alpha_Q} |\mathbf{P} - \mathbf{Q}|^2$$

#### 15.4.5 The R(N,L,M) Coefficients

The R coefficients are defined as

$$R_{NLM} \equiv \left(\frac{\partial}{\partial a}\right)^N \left(\frac{\partial}{\partial b}\right)^L \left(\frac{\partial}{\partial c}\right)^M \int_0^1 e^{-Tu^2} du$$

with

$$T \equiv \alpha \left( a^2 + b^2 + c^2 \right)$$

By extending the above definition to more general

$$R_{NLMj} \equiv \left(-\sqrt{\alpha}\right)^{N+L+M} (-2\alpha)^j \int_0^1 u^{N+L+M+2j} H_N(au\sqrt{\alpha}) H_L(bu\sqrt{\alpha}) H_M(cu\sqrt{\alpha}) e^{-Tu^2} du$$

one can see that

$$R_{000i} = (-2\alpha)^j F_i(T)$$

The Boys function is here given by

$$F_j(T) \equiv \int_0^1 u^{2j} e^{-Tu^2} du$$

and its efficient implementation can be discussed elsewhere. In Psi4, psi::Taylor\_Fjt class is used for this purpose.

Now, it is possible to show that the following recursion relationships are true:

$$R_{0,0,M+1,j} = cR_{0,0,M,j+1} + MR_{0,0,M-1,j+1}$$

$$R_{0,L+1,M,j} = bR_{0,L,M,j+1} + LR_{0,L-1,M,j+1}$$

$$R_{N+1,L,M,j} = aR_{N,L,M,j+1} + NR_{N-1,L,M,j+1}$$

This scheme is implemented in OEPDev.

#### 15.4.6 Function Documentation

#### 15.4.6.1 d\_N\_n1\_n2()

#### **Parameters**

Ν	- increment in the summation of MDH series
n1	- angular momentum of first function
n2	- angular momentum of second function
PA	- cartesian component of P-A distance
PB	- cartesian component of P-B distance
аP	- free parameter of MDH expansion

#### **Returns**

the McMurchie-Davidson-Hermite coefficient

#### 15.4.6.2 make\_mdh\_D1\_coeff()

```
void oepdev::make_mdh_D1_coeff (
    int n1,
    double aPd,
    double * buffer )
```

#### **Parameters**

n1	- angular momentum of first function
aPd	- parameter equal to 0.500/Pa where Pa is exponent
buffer	- the McMurchie-Davidson-Hermite 3-dimensional array (raveled to vector):
	axis 0: dimension 3 (x, y or z Cartesian component)
	axis 1: dimension n1+1 (0 to n1)
	• axis 2: dimension n1+1 (0 to n1)

#### See also

#### D1\_INDEX

#### 15.4.6.3 make\_mdh\_D2\_coeff()

```
void oepdev::make_mdh_D2_coeff (
    int n1,
    int n2,
    double aPd,
    double * PA,
    double * PB,
    double * buffer )
```

#### **Parameters**

n1	- angular momentum of first function
n2	- angular momentum of second function
aPd	- parameter equal to 0.500/Pa where Pa is exponent
PA	- cartesian components of P-A distance
PB	- cartesian components of P-B distance
buffer	- the McMurchie-Davidson-Hermite 4-dimensional array (raveled to vector):
	axis 0: dimension 3 (x, y or z Cartesian component)
	axis 1: dimension n1+1 (0 to n1)
	axis 2: dimension n2+1 (0 to n2)
	<ul> <li>axis 3: dimension n1+n2+1 (0 to n1+n2)</li> </ul>

#### See also

#### D2\_INDEX

#### 15.4.6.4 make\_mdh\_D2\_coeff\_explicit\_recursion()

```
void oepdev::make_mdh_D2_coeff_explicit_recursion (
    int n1,
    int n2,
    double aP,
    double * PA,
    double * PB,
    double * buffer )
```

#### **Parameters**

n1	- angular momentum of first function
n2	- angular momentum of second function
aPd	- parameter equal to 0.500/Pa where Pa is exponent
PA	- cartesian components of P-A distance
PB	- cartesian components of P-B distance
buffer	- the McMurchie-Davidson-Hermite 4-dimensional array (raveled to vector):
	axis 0: dimension 3 (x, y or z Cartesian component)
	axis 1: dimension n1+1 (0 to n1)
	axis 2: dimension n2+1 (0 to n2)
	• axis 3: dimension n1+n2+1 (0 to n1+n2)

#### See also

#### D2\_INDEX

#### 15.4.6.5 make\_mdh\_D3\_coeff()

```
void oepdev::make_mdh_D3_coeff (
    int n1,
    int n2,
    int n3,
    double aPd,
    double * PA,
    double * PB,
    double * PC,
    double * buffer )
```

#### **Parameters**

n1	- angular momentum of first function
n2	- angular momentum of second function
n3	- angular momentum of third function
aPd	- parameter equal to 0.500/Pa where Pa is exponent
PA	- cartesian components of P-A distance
PB	- cartesian components of P-B distance
PC	- cartesian components of P-C distance

# **Parameters**

```
the McMurchie-Davidson-Hermite 5-dimensional array (raveled to vector):
axis 0: dimension 3 (x, y or z Cartesian component)
axis 1: dimension n1+1 (0 to n1)
axis 2: dimension n2+1 (0 to n2)
axis 3: dimension n3+1 (0 to n3)
axis 4: dimension n1+n2+n3+1 (0 to n1+n2+n3)
```

### See also

### D3\_INDEX

# 15.4.6.6 make\_mdh\_R\_coeff()

```
void oepdev::make_mdh_R_coeff (
    int N,
    int L,
    int M,
    double alpha,
    double a,
    double b,
    double * F,
    double * buffer )
```

N	- increment in the summation of MDH series along x direction
L	- increment in the summation of MDH series along y direction
М	- increment in the summation of MDH series along z direction
alpha	- alpha parameter of R coefficient
а	- x component of PQ vector of R coefficient
b	- y component of PQ vector of R coefficient
С	- z component of PQ vector of R coefficient
F	- array of Boys function values for given alpha and PQ

buffer	- the McMurchie-Davidson 4-dimensional array (raveled to vector):
	axis 0: dimension N+1
	axis 1: dimension L+1
	axis 2: dimension M+1
	• axis 3: dimension N+L+M+1 (j-th element)

# 15.5 The Three-Dimensional Vector Fields Library

Handles all sorts of scalar distributions in 3D Euclidean space, such as general vector potentials defined at particular collection of points. In this Module, you will also find handling both random and ordered points collections in a form of a G09 cube, as well as handling G09 Cube files. You will also find solvers used to fit the generalized multipole moments of a generalized density distribution, such as the electrostatic potential (ESP) fitting method. Located at oepdev/lib3d.

### Classes

• class oepdev::MultipoleConvergence

Multipole Convergence.

• class oepdev::DMTPole

Distributed Multipole Analysis Container and Computer. Abstract Base.

class oepdev::CAMM

Cumulative Atomic Multipole Moments.

class oepdev::ESPSolver

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

· class oepdev::Points3Dlterator

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

class oepdev::CubePoints3DIterator

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

class oepdev::RandomPoints3DIterator

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

class oepdev::PointsCollection3D

Collection of points in 3D space. Abstract base.

class oepdev::RandomPointsCollection3D

Collection of random points in 3D space.

• class oepdev::CubePointsCollection3D

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

class oepdev::Field3D

General Vector Dield in 3D Space. Abstract base.

· class oepdev::ElectrostaticPotential3D

Electrostatic potential of a molecule.

class oepdev::OEPotential3D< T >

Class template for OEP 3D fields.

# **Typedefs**

using oepdev::SharedField3D = std::shared\_ptr< oepdev::Field3D >

### **Functions**

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

Construct random spherical collection of 3D field of type T.

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

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

- virtual oepdev::OEPotential3D
   T>::~OEPotential3D ()
- virtual void oepdev::OEPotential3D< T >::print () const
   Print information of the object to Psi4 output.
- virtual std::shared\_ptr< psi::Vector > oepdev::OEPotential3D< T >::compute\_xyz (const double &x, const double &z)

Compute a value of 3D field at point (x, y, z)

# 15.5.1 Detailed Description

### 15.5.2 Function Documentation

# 15.5.2.1 **OEPotential3D()** [1/2]

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

ndim	- dimensionality of 3D field (1: scalar field, >2: vector field)
np	- number of points to draw
padding	- spherical padding distance (au)
оер	- OEP object of type T
оерТуре	- type of OEP

# **15.5.2.2 OEPotential3D()** [2/2]

The points are generated according to Gaussian cube file format.

ndim	- dimensionality of 3D field (1: scalar field, >2: vector field)
nx	- number of points along x direction
ny	- number of points along y direction
nz	- number of points along z direction
рх	- padding distance along x direction
ру	- padding distance along y direction
pz	- padding distance along z direction
оер	- OEP object of type T
оерТуре	- type of OEP
options	- Psi4 options object

# 15.6 The Density Functional Theory Library

Implements the OEPDev ab initio DFT methods. Located at oepdev/libdft. Currently, this library is empty.

# 15.7 The OEPDev Utilities

Contains utility functions such as printing OEPDev preambule to the output file, class for wavefunction union, DIIS converger, CPHF Solver, SCF solver for external electrostatic perturbations, and others. You will also find here various iterators to go through orbital shells while computing ERI, or iterators over ERI itself. Located at oepdev/libutil.

#### Classes

· class oepdev::CPHF

CPHF solver class.

class oepdev::DIISManager

DIIS manager.

· class oepdev::ShellCombinationsIterator

Iterator for Shell Combinations. Abstract Base.

class oepdev::AOIntegralsIterator

Iterator for AO Integrals. Abstract Base.

class oepdev::AllAOShellCombinationsIterator\_4

Loop over all possible ERI shells in a shell quartet.

class oepdev::AllAOShellCombinationsIterator\_2

Loop over all possible ERI shells in a shell doublet.

class oepdev::AllAOIntegralsIterator\_4

Loop over all possible ERI within a particular shell quartet.

class oepdev::AllAOIntegralsIterator\_2

Loop over all possible ERI within a particular shell doublet.

struct oepdev::PerturbCharges

Structure to hold perturbing charges.

class oepdev::RHFPerturbed

RHF theory under electrostatic perturbation.

struct oepdev::ABCD

Simple structure to hold the Fourier series expansion coefficients.

struct oepdev::Fourier9

Simple structure to hold the Fourier series expansion coefficients for N=4.

class oepdev::UnitaryOptimizer

Find the optimim unitary matrix of quadratic matrix equation.

class oepdev::UnitaryOptimizer\_4\_2

Find the optimim unitary matrix for quartic-quadratic matrix equation with trace.

class oepdev::WavefunctionUnion

Union of two Wavefunction objects.

### **Macros**

#define OEPDEV\_USE\_PSI4\_DIIS\_MANAGER 0

Use DIIS from Psi4 (1) or OEPDev (0)?

#define OEPDEV\_MAX\_AM 8

L\_max.

#define OEPDEV\_N\_MAX\_AM 17

2L\_max+1

#define OEPDEV\_CRIT\_ERI 1e-9

ERI criterion for E12, E34, E123 and lambda\*EXY coefficients.

#define OEPDEV\_SIZE\_BUFFER\_R 250563

Size of R buffer (OEPDEV\_N\_MAX\_AM\*OEPDEV\_N\_MAX\_AM\*OEPDEV\_N\_MAX\_AM\*OEPDEV\_N\_MAX\_AM\*3)

#define OEPDEV\_SIZE\_BUFFER\_D2 3264

Size of D2 buffer (3\*(OEPDEV\_MAX\_AM+1)\*(OEPDEV\_MAX\_AM+1)\*OEPDEV\_N\_MAX\_AM)

# **Typedefs**

- using oepdev::SharedShellsIterator = std::shared\_ptr< ShellCombinationsIterator >
   Iterator over shells as shared pointer.
- using oepdev::SharedAOIntsIterator = std::shared\_ptr< AOIntegralsIterator >
   Iterator over AO integrals as shared pointer.

# **Functions**

PSI\_API void oepdev::preambule (void)

Print preambule for module OEPDEV.

template<typename... Args>

std::string oepdev::string\_sprintf (const char \*format, Args... args)

Format string output. Example: std::string text = oepdev::string\_sprinff("Test %3d, %13.5f", 5, -10.5425);.

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

Set up DFT functional.

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

Extract molecule from dimer.

PSI\_API std::shared\_ptr< Wavefunction > oepdev::solve\_scf (std::shared\_ptr< Molecule > molecule, std::shared\_ptr< BasisSet > primary, std::shared\_ptr< BasisSet > auxiliary, std::shared\_ptr< SuperFunctional > functional, Options & options, std::shared\_ptr< PSIO > psio, bool compute\_mints=false)

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

PSI\_API double oepdev::average\_moment (std::shared\_ptr< psi::Vector > moment)

Compute the scalar magnitude of multipole moment.

# 15.7.1 Detailed Description

### 15.7.2 Function Documentation

### 15.7.2.1 average\_moment()

### **Parameters**

mor	ment	- multipole moment vector with unique matrix elements. Now supported only for	
		dipole and quadrupole.	

#### Returns

- the average multipole moment value.

The magnitudes of multipole moments are defined here as follows:

• The dipole moment magnitude is just a norm

$$|\mu| \equiv \sqrt{\mu_x^2 + \mu_y^2 + \mu_z^2}$$

• The quadrupole moment magnitude refers to the traceless moment in Buckingham convention

$$|\Theta| \equiv \sqrt{\Theta_{zz}^2 + \frac{1}{3} \left(\Theta_{xx} - \Theta_{yy}\right)^2 + \frac{4}{3} \left(\Theta_{xy}^2 + \Theta_{xz}^2 + \Theta_{yz}^2\right)}$$

In the above equation, the quadrupole moment elements refer to its traceless form.

### 15.7.2.2 create\_superfunctional()

Now it accepts only pure HF functional.

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

### Returns

psi::SharedSuperFunctional object with functional.

# **Examples:**

example\_scf\_perturb.cc.

### 15.7.2.3 extract\_monomer()

### **Parameters**

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

#### **Returns**

psi::SharedMolecule object with indicated monomer

# 15.7.2.4 solve\_scf()

molecule	psi::SharedMolecule object with molecule
primary	shared primary basis set
auxiliary	shared auxiliary basis set
functional	DFT functional
options	psi::Options object
psio	psi::PSIO object
compute_mints	Compute integrals (write IWL TOC entry - necessary when transforming integrals)

psi::SharedWavefunction SCF wavefunction of the molecule

# 15.8 The OEPDev Testing Platform Library

Testing platform at C++ level of code. You should add more tests here when developing new functionalities, theories or models. Located at oepdev/libtest.

# **Classes**

• class oepdev::test::Test

Manages test routines.

# 15.8.1 Detailed Description

# **Chapter 16**

# **Namespace Documentation**

# 16.1 oepdev Namespace Reference

OEPDev module namespace.

### Classes

struct ABCD

Simple structure to hold the Fourier series expansion coefficients.

class AbInitioPolarGEFactory

Polarization GEFP Factory from First Principles. Hartree-Fock Approximation.

class AllAOIntegralsIterator\_2

Loop over all possible ERI within a particular shell doublet.

class AllAOIntegralsIterator\_4

Loop over all possible ERI within a particular shell quartet.

class AllAOShellCombinationsIterator\_2

Loop over all possible ERI shells in a shell doublet.

class AllAOShellCombinationsIterator\_4

Loop over all possible ERI shells in a shell quartet.

class AOIntegralsIterator

Iterator for AO Integrals. Abstract Base.

class CAMM

Cumulative Atomic Multipole Moments.

class ChargeTransferEnergyOEPotential

Generalized One-Electron Potential for Charge-Transfer Interaction Energy.

class ChargeTransferEnergySolver

Compute the Charge-Transfer interaction energy between unperturbed wavefunctions.

class CPHF

CPHF solver class.

class CubePoints3DIterator

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

class CubePointsCollection3D

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

class DIISManager

DIIS manager.

class DMTPole

Distributed Multipole Analysis Container and Computer. Abstract Base.

· class DoubleGeneralizedDensityFit

Generalized Density Fitting Scheme - Double Fit.

class EETCouplingOEPotential

Generalized One-Electron Potential for EET coupling calculations.

class ElectrostaticEnergyOEPotential

Generalized One-Electron Potential for Electrostatic Energy.

class ElectrostaticEnergySolver

Compute the Coulombic interaction energy between unperturbed wavefunctions.

class ElectrostaticPotential3D

Electrostatic potential of a molecule.

class ERI\_1\_1

2-centre ERI of the form (a|O(2)|b) where O(2) = 1/r12.

• class ERI\_2\_2

4-centre ERI of the form (ab|O(2)|cd) where O(2) = 1/r12.

class ERI\_3\_1

4-centre ERI of the form (abc|O(2)|d) where O(2) = 1/r12.

class ESPSolver

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

class FFAbInitioPolarGEFactory

Polarization GEFP Factory from First Principles: Finite-Difference Model. Arbitrary level of theory.

class Field3D

General Vector Dield in 3D Space. Abstract base.

struct Fourier9

Simple structure to hold the Fourier series expansion coefficients for N=4.

class GenEffFrag

Generalized Effective Fragment. Container Class.

class GenEffPar

Generalized Effective Fragment Parameters. Container Class.

class GenEffParFactory

Generalized Effective Fragment Factory. Abstract Base.

class GeneralizedDensityFit

Generalized Density Fitting Scheme. Abstract Base.

class GeneralizedPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class IntegralFactory

Extended IntegralFactory for computing integrals.

class LinearGradientNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class LinearNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class LinearUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

· class MultipoleConvergence

Multipole Convergence.

class NonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class OEPDevSolver

Solver of properties of molecular aggregates. Abstract base.

class OEPotential

Generalized One-Electron Potential: Abstract base.

class OEPotential3D

Class template for OEP 3D fields.

struct OEPType

Container to handle the type of One-Electron Potentials.

struct PerturbCharges

Structure to hold perturbing charges.

class Points3DIterator

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

class PointsCollection3D

Collection of points in 3D space. Abstract base.

class PolarGEFactory

Polarization GEFP Factory. Abstract Base.

class PotentialInt

Computes potential integrals.

class QuadraticGradientNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class QuadraticNonUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class QuadraticUniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class RandomPoints3DIterator

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

class RandomPointsCollection3D

Collection of random points in 3D space.

class RepulsionEnergyOEPotential

Generalized One-Electron Potential for Pauli Repulsion Energy.

class RepulsionEnergySolver

Compute the Pauli-Repulsion interaction energy between unperturbed wavefunctions.

class RHFPerturbed

RHF theory under electrostatic perturbation.

· class ShellCombinationsIterator

Iterator for Shell Combinations. Abstract Base.

class SingleGeneralizedDensityFit

Generalized Density Fitting Scheme - Single Fit.

- class TwoBodyAOInt
- class TwoElectronInt

General Two Electron Integral.

class UniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

class UnitaryOptimizer

Find the optimim unitary matrix of quadratic matrix equation.

class UnitaryOptimizer\_4\_2

Find the optimim unitary matrix for quartic-quadratic matrix equation with trace.

class UnitaryTransformedMOPolarGEFactory

Polarization GEFP Factory with Least-Squares Scaling of MO Space.

class WavefunctionUnion

Union of two Wavefunction objects.

# **Typedefs**

- using SharedField3D = std::shared\_ptr< oepdev::Field3D >
- using SharedWavefunction = std::shared\_ptr< Wavefunction >
- using SharedBasisSet = std::shared\_ptr< BasisSet >
- using SharedMatrix = std::shared\_ptr< Matrix >
- using SharedVector = std::shared\_ptr< Vector >
- using SharedDMTPole = std::shared\_ptr< DMTPole >
- using SharedWavefunctionUnion = std::shared\_ptr< WavefunctionUnion >
- using SharedOEPotential = std::shared\_ptr< OEPotential >
- using SharedIntegralFactory = std::shared\_ptr< IntegralFactory >
- using SharedTwoBodyAOInt = std::shared\_ptr< TwoBodyAOInt >
- using SharedShellsIterator = std::shared\_ptr< ShellCombinationsIterator >

Iterator over shells as shared pointer.

- using SharedAOIntsIterator = std::shared\_ptr< AOIntegralsIterator >
   Iterator over AO integrals as shared pointer.
- 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**

- double d\_N\_n1\_n2 (int N, int n1, int n2, double PA, double PB, double aP)
   Compute McMurchie-Davidson-Hermite (MDH) coefficient for binomial expansion.
- void make\_mdh\_D2\_coeff\_explicit\_recursion (int n1, int n2, double aP, double \*PA, double \*PB, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for binomial expansion by explicit recursion. This function makes the same changes to buffers as oepdev::make\_mdh\_D2\_coeff, but implements it through explicit recursion by calls to oepdev::d\_N\_n1\_n2. Therefore, it is slightly slower. Here for debugging purposes.

void make\_mdh\_D1\_coeff (int n1, double aPd, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for monomial expansion.

void make\_mdh\_D2\_coeff (int n1, int n2, double aPd, double \*PA, double \*PB, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for binomial expansion.

void make\_mdh\_D3\_coeff (int n1, int n2, int n3, double aPd, double \*PA, double \*PB, double \*PC, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for trinomial expansion.

void make\_mdh\_R\_coeff (int N, int L, int M, double alpha, double a, double b, double c, double \*F, double \*buffer)

Compute the McMurchie-Davidson R coefficients.

- constexpr std::complex< double > operator""\_i (unsigned long long d)
- constexpr std::complex< double > operator""\_i (long double d)
- PSI\_API void preambule (void)

Print preambule for module OEPDEV.

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

Set up DFT functional.

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

Extract molecule from dimer.

PSI\_API std::shared\_ptr< Wavefunction > solve\_scf (std::shared\_ptr< Molecule > molecule, std::shared\_ptr< BasisSet > primary, std::shared\_ptr< BasisSet > auxiliary, std::shared\_ptr< SuperFunctional > functional, Options & options, std::shared\_ptr< PSIO > psio, bool compute\_mints=false)

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

PSI\_API double average\_moment (std::shared\_ptr< psi::Vector > moment)

Compute the scalar magnitude of multipole moment.

template<typename... Args>
 std::string string\_sprintf (const char \*format, Args... args)

Format string output. Example: std::string text = oepdev::string\_sprinff("Test %3d, %13.5f", 5, -10.5425);.

# 16.1.1 Detailed Description

Contains all the functionalities for the development of the Generalized One-Electroc Potentials (OEP's).

# 16.2 psi Namespace Reference

Psi4 package namespace.

# **Typedefs**

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

# **Functions**

- PSI\_API int read\_options (std::string name, Options & options)
   Options for the OEPDev plugin.
- void export\_dmtp (py::module &)
- void export\_cphf (py::module &)
- void export\_solver (py::module &)
- PSI\_API SharedWavefunction oepdev (SharedWavefunction ref\_wfn, Options & options)
   Main routine of the OEPDev plugin.
- PYBIND11\_MODULE (oepdev, m)

# 16.2.1 Detailed Description

Contains all Psi4 functionalities.

# 16.2.2 Function Documentation

### 16.2.2.1 oepdev()

Created with intention to test various models of the interaction energy between two molecules, described by the Hartree-Fock-Roothaan-Hall theory or the configuration interaction with singles theory.

In particular, the plugin tests the models of:

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

against benchmarks (exact or reference solutions). The list of implemented models can be found in Implemented Models .

# **Parameters**

ref_wfn	shared wavefunction of a dimer
options	psi::Options object

#### **Returns**

psi::SharedWavefunction (either ref\_wfn or wavefunction union)

# 16.2.2.2 read\_options()

name	name of driver function
options	psi::Options object

Returns

true

# **Chapter 17**

# **Class Documentation**

# 17.1 oepdev::ABCD Struct Reference

Simple structure to hold the Fourier series expansion coefficients.

```
#include <unitary_optimizer.h>
```

# **Public Attributes**

- double A
- · double B
- · double C
- double **D**

# 17.1.1 Detailed Description

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

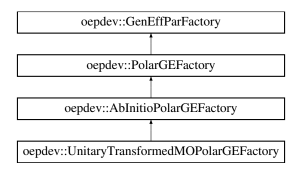
• oepdev/libutil/unitary\_optimizer.h

# 17.2 oepdev::AbInitioPolarGEFactory Class Reference

Polarization GEFP Factory from First Principles. Hartree-Fock Approximation.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::AbInitioPolarGEFactory:



### **Public Member Functions**

- AbInitioPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
- virtual std::shared\_ptr< GenEffPar > compute (void)
   Compute the density matrix susceptibility tensors.

# **Additional Inherited Members**

# 17.2.1 Detailed Description

Implements creation of the density matrix susceptibility tensors for which X=1. Guarantees the idempotency of the density matrix up to first-order in LCAO-MO variation. The density matrix susceptibility tensor is represented by:

$$\delta D_{\alpha\beta} = \sum_{i} \mathbf{B}_{\alpha\beta}^{(i;1)} \cdot \mathbf{F}(\mathbf{r}_i)$$

where  $\mathbf{B}_{\alpha\beta}^{(i;1)}$  is the density matrix dipole polarizability defined for the distributed LMO site at  $\mathbf{r}_i$ . Its explicit form is given by

$$\mathbf{B}_{\alpha\beta}^{(i;1)} = C_{\alpha i}^{(0)} \mathbf{b}_{\beta}^{(i;1)} C_{\beta i}^{(0)} \mathbf{b}_{\alpha}^{(i;1)} - \sum_{\gamma} \left( D_{\alpha\gamma}^{(0)} C_{\beta i}^{(0)} + D_{\beta\gamma}^{(0)} C_{\alpha i}^{(0)} \right) \mathbf{b}_{\gamma}^{(i;1)}$$

where the susceptibility of the LCAO-MO coefficient is given by

$$b_{\alpha;w}^{(i;1)} = \frac{1}{4} \sum_{u}^{x,y,z} \left[\alpha_i\right]_{uw} \left[ \left[\mathbf{L}_i\right]_{\text{Left}}^{-1} \right]_{u;\alpha}$$

for w = x, y, z. The auxiliary tensor  $\mathbb{L}$  is defined as

$$\mathbb{L} = \mathbf{C}^{(0)\mathrm{T}} \cdot \mathbb{M} \cdot \left( \mathbf{1} - \mathbf{D}^{(0)} \right)$$

where  $\mathbb M$  is the dipole integral vector of matrices in AO representation. The left inverse of the i-th element is defined as

$$[\mathbf{L}_i]_{\mathrm{Left}}^{-1} \equiv \left[\mathbf{L}_i^{\mathrm{T}} \cdot \mathbf{L}_i\right]^{-1} \cdot \mathbf{L}_i^{\mathrm{T}}$$

Note that  $L_i \equiv [\mathbb{L}]_i$  is a  $n \times 3$  matrix, whereas its left inverse is a  $3 \times n$  matrix with n being the size of the AO basis set.

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

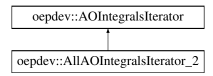
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_abinitio.cc

# 17.3 oepdev::AllAOIntegralsIterator\_2 Class Reference

Loop over all possible ERI within a particular shell doublet.

#include <integrals\_iter.h>

Inheritance diagram for oepdev::AllAOIntegralsIterator\_2:



### **Public Member Functions**

AllAOIntegralsIterator\_2 (const ShellCombinationsIterator \*shellIter)

Construct by shell iterator (const object)

AllAOIntegralsIterator\_2 (std::shared\_ptr< ShellCombinationsIterator > shellIter)

Construct by shell iterator (pointed by shared pointer)

void first ()

First iteration.

void next ()

Next iteration.

• int i () const

Grab the current integral i index.

int j () const

Grab the current integral j index.

• int index () const

### **Additional Inherited Members**

# 17.3.1 Detailed Description

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

See also

AllAOShellCombinationsIterator\_2

### 17.3.2 Constructor & Destructor Documentation

# 17.3.2.1 AllAOIntegralsIterator\_2() [1/2]

#### **Parameters**

80

```
shellIter - shell iterator object
```

### 17.3.2.2 AllAOIntegralsIterator\_2() [2/2]

### **Parameters**

```
shellIter - shell iterator object
```

# 17.3.3 Member Function Documentation

### 17.3.3.1 index()

Grab the current index of integral value stored in the buffer

Implements oepdev::AOIntegralsIterator.

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

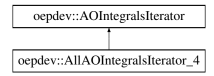
- oepdev/libutil/integrals\_iter.h
- oepdev/libutil/integrals\_iter.cc

# 17.4 oepdev::AllAOIntegralsIterator\_4 Class Reference

Loop over all possible ERI within a particular shell quartet.

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

Inheritance diagram for oepdev::AllAOIntegralsIterator\_4:



### **Public Member Functions**

AllAOIntegralsIterator\_4 (const ShellCombinationsIterator \*shellIter)

Construct by shell iterator (const object)

AllAOIntegralsIterator\_4 (std::shared\_ptr< ShellCombinationsIterator > shellIter)

Construct by shell iterator (pointed by shared pointer)

void first ()

First iteration.

void next ()

Next iteration.

• 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

# **Additional Inherited Members**

# 17.4.1 Detailed Description

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

See also

AllAOShellCombinationsIterator\_4

# 17.4.2 Constructor & Destructor Documentation

# 17.4.2.1 AllAOIntegralsIterator\_4() [1/2]

#### **Parameters**

```
shellIter - shell iterator object
```

### 17.4.2.2 AllAOIntegralsIterator\_4() [2/2]

### **Parameters**

```
shellIter - shell iterator object
```

# 17.4.3 Member Function Documentation

### 17.4.3.1 index()

Grab the current index of integral value stored in the buffer

Implements oepdev::AOIntegralsIterator.

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

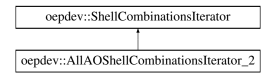
- oepdev/libutil/integrals\_iter.h
- oepdev/libutil/integrals\_iter.cc

# 17.5 oepdev::AllAOShellCombinationsIterator\_2 Class Reference

Loop over all possible ERI shells in a shell doublet.

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

Inheritance diagram for oepdev::AllAOShellCombinationsIterator\_2:



### **Public Member Functions**

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

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

- AllAOShellCombinationsIterator\_2 (std::shared\_ptr< IntegralFactory > integrals)
  - Construct by providing integral factory.
- AllAOShellCombinationsIterator\_2 (const IntegralFactory &integrals)
- AllAOShellCombinationsIterator\_2 (std::shared\_ptr< psi::IntegralFactory > integrals)

Construct by providing integral factory.

- AllAOShellCombinationsIterator\_2 (const psi::IntegralFactory &integrals)
- void first ()

First iteration.

void next ()

Next iteration.

- void compute\_shell (std::shared\_ptr< oepdev::TwoBodyAOInt > tei) const
   Compute ERI's for the current shell. The eris are stored in the buffer of the argument object.
- void compute\_shell (std::shared\_ptr< psi::TwoBodyAOInt > tei) const
- int P () const

Grab the current shell P index.

• int Q () const

Grab the current shell Q index.

# **Additional Inherited Members**

# 17.5.1 Detailed Description

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

### 17.5.2 Constructor & Destructor Documentation

# 17.5.2.1 AllAOShellCombinationsIterator\_2() [1/5]

#### **Parameters**

bs_1	- basis set of axis 1
bs_2	- basis set of axis 2

# 17.5.2.2 AllAOShellCombinationsIterator\_2() [2/5]

### **Parameters**

```
integrals - OepDev integral factory object
```

# 17.5.2.3 AllAOShellCombinationsIterator\_2() [3/5]

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

# 17.5.2.4 AllAOShellCombinationsIterator\_2() [4/5]

### **Parameters**

```
integrals - Psi4 integral factory object
```

# 17.5.2.5 AllAOShellCombinationsIterator\_2() [5/5]

```
AllAOShellCombinationsIterator_2::AllAOShellCombinationsIterator_2 (
```

```
const psi::IntegralFactory & integrals )
```

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

### 17.5.3 Member Function Documentation

### 17.5.3.1 compute\_shell()

### **Parameters**

```
tei - two electron AO integral
```

Implements oepdev::ShellCombinationsIterator.

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

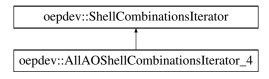
- oepdev/libutil/integrals\_iter.h
- oepdev/libutil/integrals\_iter.cc

# 17.6 oepdev::AllAOShellCombinationsIterator\_4 Class Reference

Loop over all possible ERI shells in a shell quartet.

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

Inheritance diagram for oepdev::AllAOShellCombinationsIterator\_4:



### **Public Member Functions**

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

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

AllAOShellCombinationsIterator\_4 (std::shared\_ptr< IntegralFactory > integrals)

Construct by providing integral factory.

- AllAOShellCombinationsIterator\_4 (const IntegralFactory &integrals)
- AllAOShellCombinationsIterator\_4 (std::shared\_ptr< psi::IntegralFactory > integrals)

Construct by providing integral factory.

- AllAOShellCombinationsIterator\_4 (const psi::IntegralFactory &integrals)
- void first ()

Do the first iteration.

void next ()

Do the next iteration.

- void compute\_shell (std::shared\_ptr< oepdev::TwoBodyAOInt > tei) const
- void compute\_shell (std::shared\_ptr< psi ::TwoBodyAOInt > tei) const
- 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.

# **Additional Inherited Members**

# 17.6.1 Detailed Description

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

### 17.6.2 Constructor & Destructor Documentation

# 17.6.2.1 AllAOShellCombinationsIterator\_4() [1/5]

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

### 17.6.2.2 AllAOShellCombinationsIterator\_4() [2/5]

### **Parameters**

### 17.6.2.3 AllAOShellCombinationsIterator\_4() [3/5]

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

### 17.6.2.4 AllAOShellCombinationsIterator\_4() [4/5]

#### **Parameters**

```
integrals - OepDev integral factory object
```

### 17.6.2.5 AllAOShellCombinationsIterator\_4() [5/5]

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

### 17.6.3 Member Function Documentation

### 17.6.3.1 compute\_shell()

Compute integrals in a current shell. Works both for oepdev::TwoBodyAOInt and psi::TwoBodyAOInt

### **Parameters**

```
tei - two body integral object
```

Implements oepdev::ShellCombinationsIterator.

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

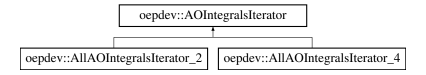
- oepdev/libutil/integrals\_iter.h
- oepdev/libutil/integrals\_iter.cc

# 17.7 oepdev::AOIntegralsIterator Class Reference

Iterator for AO Integrals. Abstract Base.

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

Inheritance diagram for oepdev::AOIntegralsIterator:



# **Public Member Functions**

AOIntegralsIterator ()

Base Constructor.

virtual ∼AOIntegralsIterator ()

Base Destructor.

virtual void first (void)=0

Do the first iteration.

virtual void next (void)=0

Do the next iteration.

virtual int i (void) const

Grab i-th index.

virtual int j (void) const

Grab j-th index.

virtual int k (void) const

Grab k-th index.

virtual int I (void) const

Grab I-th index.

virtual int index (void) const =0

Grab index in the integral buffer.

virtual bool is\_done (void)

Returns the status of an iterator.

### **Static Public Member Functions**

- static std::shared\_ptr< AOIntegralsIterator > build (const ShellCombinationsIterator \*shellIter, std::string mode="ALL")
- static std::shared\_ptr< AOIntegralsIterator > build (std::shared\_ptr< ShellCombinationsIterator > shellIter, std::string mode="ALL")

### **Protected Attributes**

· bool done

The status of an iterator.

# 17.7.1 Detailed Description

# 17.7.2 Member Function Documentation

Build AO integrals iterator from current state of iterator over shells

#### **Parameters**

```
shellIter - iterator over shells - either "ALL" or "UNIQUE" (iterate over all or unique integrals)
```

### Returns

iterator over AO integrals

# **17.7.2.2** build() [2/2]

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

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

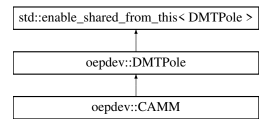
- oepdev/libutil/integrals\_iter.h
- · oepdev/libutil/integrals\_iter.cc

# 17.8 oepdev::CAMM Class Reference

Cumulative Atomic Multipole Moments.

```
#include <dmtp.h>
```

Inheritance diagram for oepdev::CAMM:



# **Public Member Functions**

CAMM (psi::SharedWavefunction wfn, int n)

Construct CAMM DMTPole object.

virtual void compute (psi::SharedMatrix D, bool transition, int n)

Compute DMTP's from the one-particle density matrix.

### **Additional Inherited Members**

# 17.8.1 Detailed Description

Cumulative atomic multipole representation of the molecular charge distribution. Method of Sokalski and Poirier. Ref.: W. A. Sokalski and R. A. Poirier, *Chem. Phys. Lett.*, 98(1) **1983** 

# Methodology.

The distributed multipole moments are computed in the following way:

- first the atomic additive multipole moments (AAMM's) with origins set to the global coordinate system origin are computed. AO basis set partitioning is used to dostribute the AAMM's onto the atomic centres.
- subsequently, the AAMM's origins are moved to the corresponding atomic site.

The computation of the AAMM's is performed according to the following prescription:

$$M_{uw...z}^{(A)}(\mathbf{0}) = \sum_{lpha \in A} \sum_{eta \in ext{allAO's}} D_{lphaeta}^{ ext{OED}} ra{lpha} \mathscr{M}_{uw...z}(\mathbf{0}) \ket{eta}$$

where  $M_{uw...z}^{(A)}$  denotes the (uw...z)-th component of the multipole centered at atomic site A, the symbol  $\mathcal{M}(\mathbf{0})$  is the associated quantum mechanical operator and  $D_{\alpha\beta}^{\mathrm{OED}}$  is the (generalized) one-particle density matrx element in AO basis (Greek indices).

Recentering of the multipole moments is described in the documentation of oepdev::DMTPole::recenter.

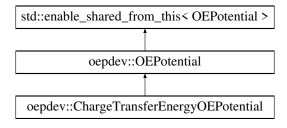
The documentation for this class was generated from the following files:

- oepdev/lib3d/dmtp.h
- oepdev/lib3d/dmtp\_camm.cc

# 17.9 oepdev::ChargeTransferEnergyOEPotential Class Reference

Generalized One-Electron Potential for Charge-Transfer Interaction Energy.

Inheritance diagram for oepdev::ChargeTransferEnergyOEPotential:



# **Public Member Functions**

- ChargeTransferEnergyOEPotential (SharedWavefunction wfn, SharedBasisSet auxiliary, SharedBasisSet intermediate, Options & options)
- ChargeTransferEnergyOEPotential (SharedWavefunction wfn, Options & options)
- virtual void compute (const std::string &oepType) override

Compute matrix forms of all OEP's within a specified OEP type.

virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, std::shared\_ptr< psi::Vector > &v) override

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

virtual void print\_header () const override

### **Additional Inherited Members**

# 17.9.1 Detailed Description

Contains the following OEP types:

- Otto-Ladik.V1 DF-based term
- Otto-Ladik.V2 ESP-based term
- Otto-Ladik.V3 ESP-based term

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

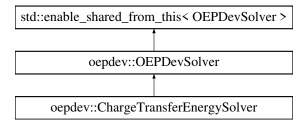
- oepdev/liboep/oep.h
- oepdev/liboep/oep\_energy\_ct.cc

# 17.10 oepdev::ChargeTransferEnergySolver Class Reference

Compute the Charge-Transfer interaction energy between unperturbed wavefunctions.

```
#include <solver.h>
```

Inheritance diagram for oepdev::ChargeTransferEnergySolver:



# **Public Member Functions**

- ChargeTransferEnergySolver (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**

## 17.10.1 Detailed Description

The implemented methods are shown below

Table 17.14: Methods available in the Solver

Keyword	Method Description	
	Benchmark Methods	
OTTO_LADIK	*Default*. CT energy at HF level from Otto and Ladik (1975).	
EFP2	CT energy at HF level from EFP2 model.	
OEP-Based Methods		
OTTO_LADIK	*Default*. OEP-based Otto-Ladik expressions.	

In order to construct this solver, **always** use the <code>OEPDevSolver::build</code> static factory method.

Below the detailed description of the implemented equations is given for each of the above provided methods. In the formulae across, it is assumed that the orbitals are real. The Coulomb notation for electron repulsion integrals (ERI's) is adopted; i.e,

$$(ac|bd) = \iint d\mathbf{r}_1 d\mathbf{r}_2 \phi_a(\mathbf{r}_1) \phi_c(\mathbf{r}_1) \frac{1}{r_{12}} \phi_b(\mathbf{r}_2) \phi_d(\mathbf{r}_2)$$

Greek subscripts denote basis set orbitals whereas Italic subscripts denote the occupied molecular orbitals.

The CT energy between molecules A and B is given by

$$E^{\text{CT}} = E^{\text{A}^{+}\text{B}^{-}} + E^{\text{A}^{-}\text{B}^{+}}$$

#### **Benchmark Methods**

CT energy at HF level by Otto and Ladik (1975).

For a closed-shell system, CT energy equation of Otto and Ladik becomes

$$E^{\mathrm{A^{+}B^{-}}} \approx 2 \sum_{i \in A}^{\mathrm{Occ_{A}}} \sum_{n \in B}^{\mathrm{Vir_{B}}} \frac{V_{in}^{2}}{\varepsilon_{i} - \varepsilon_{n}}$$

where

$$\begin{split} V_{in} &= V_{in}^{B} + 2\sum_{j \in B}^{\text{Occ}_{B}} (in|jj) - \sum_{k \in A}^{\text{Occ}_{A}} S_{kn} \left\{ V_{ik}^{B} + 2\sum_{j \in B}^{\text{Occ}_{B}} (ik|jj) \right\} \\ &- \sum_{j \in B}^{\text{Occ}_{B}} \left[ S_{ij} \left\{ V_{nj}^{A} + 2\sum_{k \in A}^{\text{Occ}_{A}} (1 - \delta_{ik})(nj|kk) \right\} - (nj|ij) \right] + \sum_{k \in A}^{\text{Occ}_{A}} \sum_{j \in B}^{\text{Occ}_{A}} S_{kj} (1 - \delta_{ik})(ik|nj) \end{split}$$

and analogously the twin term.

CT energy at HF level by EFP2.

In EFP2 method, CT energy is given as

$$E^{\mathrm{A^+B^-}} \approx 2 \sum_{i \in A}^{\mathrm{Occ_A}} \sum_{n \in B}^{\mathrm{Vir_B}} \frac{V_{in}^2}{F_{ii} - T_{nn}}$$

where

$$V_{in}^2 = \frac{V_{in}^B - \sum_{m \in A}^{\text{All}_A} V_{im} S_{mn}^B}{1 - \sum_{m \in A}^{\text{All}_A} S_{mn}^2} \left\{ V_{in}^B - \sum_{m \in A}^{\text{All}_A} V_{im}^B S_{mn} + \sum_{j \in B}^{\text{Occ}_B} S_{ij} \left( T_{nj} - \sum_{m \in A}^{\text{All}_A} S_{nm} T_{mj} \right) \right\}$$

and analogously the twin term.

## **OEP-Based Methods**

**OEP-Based Otto-Ladik's theory** 

After introducing OEP's, the original Otto-Ladik's theory is reformulated *without* approximation as

$$E^{\mathrm{A^+B^-}} pprox 2 \sum_{i \in A}^{\mathrm{Occ_A}} \sum_{n \in B}^{\mathrm{Vir_B}} rac{\left(V_{in}^{\mathrm{DF}} + V_{in}^{\mathrm{ESP,A}} + V_{in}^{\mathrm{ESP,B}}
ight)^2}{arepsilon_i - arepsilon_n}$$

where

$$V_{in}^{ ext{DF}} = \sum_{\eta \in B}^{ ext{Aux}_B} S_{i\eta} G_{\eta n}^B$$
 $V_{in}^{ ext{ESP,A}} = \sum_{k \in A}^{ ext{Occ}_A} \sum_{j \in B}^{ ext{Occ}_B} S_{kj} \sum_{x \in A} V_{nj}^{(x)} q_{ik}^{(x)}$ 
 $V_{in}^{ ext{ESP,B}} = -\sum_{k \in A}^{ ext{Occ}_A} S_{kn} V_{ik}^B$ 

The OEP matrix for density fitted part is given by

$$G_{\eta n}^{B} = \sum_{\eta' \in B}^{\text{Aux}_{\text{B}}} [\mathbf{S}^{-1}]_{\eta \eta'} \left\{ V_{\eta' n}^{B} + \sum_{j \in B}^{\text{Occ}_{\text{B}}} \left[ 2(\eta' n | jj) - (\eta' j | nj) \right] \right\}$$

The OEP ESP-A charges are fit to reproduce the OEP potential

$$v_{ik}^{A}(\mathbf{r}) \equiv (1 - \delta_{ik}) \int \frac{\phi_{i}(\mathbf{r}')\phi_{k}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}' - \delta_{ik} \left( \sum_{x \in A} \frac{-Z_{x}}{|\mathbf{r} - \mathbf{r}_{x}|} + 2 \sum_{k \in A}^{\text{Occ}_{A}} \int \frac{\phi_{k}(\mathbf{r}')\phi_{k}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}' - 2 \int \frac{\phi_{i}(\mathbf{r}')\phi_{i}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}' \right)$$

so that

$$v_{ik}^{A}(\mathbf{r}) \cong \sum_{x \in A} \frac{q_{ik}^{(x)}}{|\mathbf{r} - \mathbf{r}_x|}$$

The OEP ESP-B charges are fit to reproduce the electrostatic potential of molecule *B* (they are standard ESP charges).

#### 17.10.2 Member Function Documentation

## 17.10.2.1 compute\_benchmark()

Each solver object has one DEFAULT benchmark method

#### **Parameters**

```
method - benchmark method
```

Implements oepdev::OEPDevSolver.

#### 17.10.2.2 compute\_oep\_based()

Each solver object has one DEFAULT OEP-based method.

#### **Parameters**

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

Implements oepdev::OEPDevSolver.

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

- oepdev/libsolver/solver.h
- oepdev/libsolver/solver\_energy\_ct.cc

## 17.11 oepdev::CPHF Class Reference

#### **CPHF** solver class.

```
#include <cphf.h>
```

#### **Public Member Functions**

CPHF (SharedWavefunction ref\_wfn, Options &options)

Constructor.

∼CPHF ()

Desctructor.

· void compute (void)

run the calculations

void print (void) const

print to output file

• int nocc (void) const

get the number of occupied orbitals

- std::shared\_ptr< Wavefunction > wfn (void) const grab the wavefunction
- · Options & options (void) const

grab the Psi4 options

- std::shared\_ptr< Matrix > polarizability (void) const retrieve the molecular (total) polarizability
- std::shared\_ptr< Matrix > polarizability (int i) const retrieve the i-th orbital-associated polarizability
- std::shared\_ptr< Matrix > polarizability (int i, int j) const
   retrieve the charge-transfer polarizability associated with orbitals i and j
- std::shared\_ptr< Matrix > X (int x) const

retrieve the X operator O-V perturbation matrix in AO basis for x-th component

std::vector< std::shared\_ptr< Matrix >> X (void) const

retrieve the X operator O-V perturbation matrix in AO basis for all three Cartesian components

std::shared\_ptr< Matrix > X\_mo (int x) const

retrieve the X operator O-V perturbation matrix in MO basis for x-th component

- std::vector < std::shared\_ptr < Matrix > > X\_mo (void) const
   retrieve the X operator O-V perturbation matrix in MO basis for all three Cartesian components
- std::shared\_ptr< Matrix > F\_mo (int x) const
- retrieve the F operator O-V perturbation matrix in MO basis for x-th component
   std::vector< std::shared\_ptr< Matrix >> F\_mo (void) const
- retrieve the F operator O-V perturbation matrix in MO basis for all three Cartesian components
- std::shared\_ptr< Matrix > T (void) const

retrieve the transformation from old to new MO's

- std::shared\_ptr< Matrix > Cocc (void) const retrieve the Cocc
- std::shared\_ptr< Matrix > Cvir (void) const retrieve the Cvir
- std::shared\_ptr< Vector > Imo\_centroid (int i) const retrieve the i-th orbital (LMO) centroid
- std::shared\_ptr< Localizer > localizer (void) const retrieve the orbital localizer

#### **Protected Attributes**

- std::shared\_ptr< psi::Wavefunction > \_wfn
   Wavefunction object.
- std::shared\_ptr< Localizer > \_localizer
   Orbital localizer.
- 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 \_conv

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 > \_molecularPolarizability

Total (molecular) polarizability tensor.

• std::vector < std::shared\_ptr < Vector > > \_orbitalCentroids

LMO centroids.

std::vector < std::shared\_ptr < Matrix > > \_orbitalPolarizabilities
 orbital-associated polarizability tensors

- std::vector < std::vector < std::shared\_ptr < Matrix > > \_orbitalChargeTransferPolarizabilities
   orbital-orbital charge-transfer polarizability tensors
- std::vector< std::shared\_ptr< Matrix >> \_X\_OV\_ao\_matrices

Perturbation X Operator O-> V matrices in AO basis.

std::vector< std::shared\_ptr< Matrix >> \_X\_OV\_mo\_matrices

Perturbation X Operator O-> V matrices in MO basis.

std::vector< std::shared\_ptr< Matrix >> \_F\_OV\_mo\_matrices

Electric Field Operator O-> V matrices in MO basis.

std::shared\_ptr< psi::Matrix > \_T

Transformation from old to new MO's.

## 17.11.1 Detailed Description

Solves CPHF equations (now only for RHF wavefunction). Computes molecular and polarizabilities associated with the localized molecular orbitals (LMO).

Note

#### Useful options:

- CPHF\_CONVER convergence of CPHF. Default: 1e-8 (au)
- CPHF\_CONVER maximum number of iterations. Default: 50
- CPHF\_DIIS wheather use DIIS or not. Default: true
- CPHF\_DIIS\_DIM dimension of iterative subspace. Default: 3
- CPHF\_LOCALIZE localize the molecular orbitals? Default: true
- CPHF\_LOCALIZER set orbital localization method. Available: BOYS and PIPEK MEZEY. Default: BOYS

#### 17.11.2 Constructor & Destructor Documentation

## 17.11.2.1 CPHF()

#### **Parameters**

ref_wfn	reference HF wavefunction
options	set of Psi4 options

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

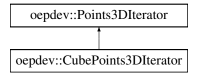
- · oepdev/libutil/cphf.h
- · oepdev/libutil/cphf.cc

# 17.12 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**

- CubePoints3DIterator (const int &nx, const int &ny, const int &nz, const double &dx, const double &dy, const double &dx, const double &ox, const double &oy, const double &oz)
- · 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**

## 17.12.1 Detailed Description

**Note:** Always create instances by using static factory method from Points3DIterator. Do not use constructor of this class.

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

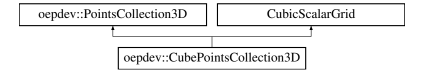
- · oepdev/lib3d/space3d.h
- oepdev/lib3d/space3d.cc

# 17.13 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 &py, const double &pz, 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, const int &col=0)

#### **Additional Inherited Members**

## 17.13.1 Detailed Description

**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/lib3d/space3d.h
- · oepdev/lib3d/space3d.cc

# 17.14 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)

## 17.14.1 Detailed Description

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.

#### 17.14.2 Constructor & Destructor Documentation

#### 17.14.2.1 DIISManager()

```
int na,
int nb )
```

#### Constructor.

#### **Parameters**

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

## 17.14.3 Member Function Documentation

```
17.14.3.1 compute()
```

Perform DIIS interpolation.

## 17.14.3.2 put()

Put the current solution to the DIIS manager.

#### **Parameters**

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

## 17.14.3.3 update()

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

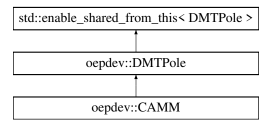
oepdev/libutil/diis.cc

## 17.15 oepdev::DMTPole Class Reference

Distributed Multipole Analysis Container and Computer. Abstract Base.

```
#include <dmtp.h>
```

Inheritance diagram for oepdev::DMTPole:



## **Public Member Functions**

virtual ~DMTPole ()

Destructor.

• virtual bool has\_charges () const

Has distributed charges?

virtual bool has\_dipoles () const

Has distributed dipoles?

virtual bool has\_quadrupoles () const

Has distributed quadrupoles?

virtual bool has\_octupoles () const

Has distributed octupoles?

virtual bool has\_hexadecapoles () const

Has distributed hexadecapoles?

virtual psi::SharedMatrix centres () const

Get the positions of distribution centres.

• virtual psi::SharedMatrix origins () const

Get the positions of distribution origins.

virtual std::vector < psi::SharedMatrix > charges () const
 Get the distributed charges.

virtual std::vector< psi::SharedMatrix > dipoles () const
 Get the distributed dipoles.

virtual std::vector < psi::SharedMatrix > quadrupoles () const
 Get the distributed quadrupoles.

virtual std::vector< psi::SharedMatrix > octupoles () const

Get the distributed octupoles.

virtual std::vector < psi::SharedMatrix > hexadecapoles () const
 Get the distributed hexadecapoles.

virtual psi::SharedMatrix charges (int i) const

Get the distributed charges for the ith distribution.

virtual psi::SharedMatrix dipoles (int i) const

Get the distributed dipoles for the ith distribution.

virtual psi::SharedMatrix quadrupoles (int i) const

Get the distributed quadrupoles for the ith distribution.

virtual psi::SharedMatrix octupoles (int i) const

Get the distributed octupoles for the ith distribution.

virtual psi::SharedMatrix hexadecapoles (int i) const

Get the distributed hexadecapoles for the ith distribution.

virtual int n\_sites () const

Get the number of distributed sites.

virtual int n\_dmtp () const

Get the number of distributions.

void set\_charges (std::vector< psi::SharedMatrix > M)

Set the distributed charges.

void set\_dipoles (std::vector < psi::SharedMatrix > M)

Set the distributed dipoles.

void set\_quadrupoles (std::vector< psi::SharedMatrix > M)

Set the distributed quadrupoles.

void set\_octupoles (std::vector< psi::SharedMatrix > M)

Set the distributed octupoles.

void set\_hexadecapoles (std::vector < psi::SharedMatrix > M)

Set the distributed hexadecapoles.

void set\_charges (psi::SharedMatrix M, int i)

Set the distributed charges for the ith distribution.

void set\_dipoles (psi::SharedMatrix M, int i)

Set the distributed dipoles for the ith distribution.

void set\_quadrupoles (psi::SharedMatrix M, int i)

Set the distributed quadrupoles for the ith distribution.

void set\_octupoles (psi::SharedMatrix M, int i)

Set the distributed octupoles for the ith distribution.

void set\_hexadecapoles (psi::SharedMatrix M, int i)

Set the distributed hexadecapoles for the ith distribution.

- virtual void recenter (psi::SharedMatrix new\_origins)
- void translate (psi::SharedVector transl)

Translate the DMTP sets.

void rotate (psi::SharedMatrix rotmat)

Rotate the DMTP sets.

void superimpose (psi::SharedMatrix ref\_xyz, std::vector< int > suplist)

Superimpose the DMTP sets.

virtual void compute (psi::SharedMatrix D, bool transition, int i)=0

Compute DMTP's from the one-particle density matrix.

void compute (std::vector < psi::SharedMatrix > D, std::vector < bool > transition)

Compute DMTP's from the set of the one-particle density matrices.

void compute (void)

Compute DMTP's from the sum of the ground-state alpha and beta one-particle density matrices (transition=false, i=0)

 std::shared\_ptr< MultipoleConvergence > energy (std::shared\_ptr< DMTPole > other, MultipoleConvergence::ConvergenceLevel max\_clevel=MultipoleConvergence::R5)

Evaluate the generalized interaction energy.

 std::shared\_ptr< MultipoleConvergence > potential (std::shared\_ptr< DMTPole > other, MultipoleConvergence::ConvergenceLevel max\_clevel=MultipoleConvergence::R5)

Evaluate the generalized potential.

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

static std::shared\_ptr< DMTPole > build (const std::string &type, std::shared\_ptr< psi::Wavefunction > wfn, int n=1)

Build an empty DMTP object from the wavefunction.

#### **Protected Member Functions**

DMTPole (std::shared\_ptr< psi::Wavefunction > wfn, int n)

Construct an empty DMTP object from the wavefunction.

void compute\_integrals ()

Compute multipole integrals.

void compute\_order ()

Compute maximum order of the integrals.

virtual void recenter (psi::SharedMatrix new\_origins, int i)

Change origins of the distributed multipole moments of ith set.

virtual void allocate ()

Initialize and allocate memory.

#### **Protected Attributes**

std::string name\_

Name of the distribution method.

• psi::SharedMolecule mol\_

Molecule associated with this DMTP.

psi::SharedWavefunction wfn\_

Wavefunction associated with this DMTP.

psi::SharedBasisSet primary\_

Basis set (primary)

int nDMTPs\_

Number of DMTP's.

int nSites\_

Number of DMTP sites.

int order\_

Maximum order of the multipole.

std::vector < psi::SharedMatrix > mpInts\_

Multipole integrals.

bool hasCharges\_

Has distributed charges?

bool hasDipoles\_

Has distributed dipoles?

bool hasQuadrupoles\_

Has distributed quadrupoles?

bool hasOctupoles\_

Has distributed octupoles?

bool hasHexadecapoles\_

Has distributed hexadecapoles?

psi::SharedMatrix centres\_

DMTP centres.

• psi::SharedMatrix origins\_

DMTP origins.

std::vector < psi::SharedMatrix > charges\_

DMTP charges.

std::vector < psi::SharedMatrix > dipoles\_

DMTP dipoles.

std::vector < psi::SharedMatrix > quadrupoles\_

DMTP quadrupoles.

std::vector < psi::SharedMatrix > octupoles\_

DMTP octupoles.

std::vector< psi::SharedMatrix > hexadecapoles\_

DMTP hexadecapoles.

#### **Friends**

class MultipoleConvergence

## 17.15.1 Detailed Description

Handles the distributed multipole expansions up to hexadecapoles. Distributed centres as well as DMTP origins are allowed to be located in arbitrary points in space. The object describes a set of *N* DMTP's, that can be generated by providing one-particle density matrices in AO basis. Nuclear contributions can be switched on or off separately for each DMTP within a set. The following operations on the DMTP sets are available through the API:

- · translation
- rotation
- superimposition
- · recentering the origins
- · computing the generalized property from another DMTP set

#### 17.15.2 Constructor & Destructor Documentation

## 17.15.2.1 DMTPole()

#### **Parameters**

wfn	- wavefunction
n	- number of DMTP sets

Do not use this constructor. Use the DMTPole::build method.

## 17.15.3 Member Function Documentation

## 17.15.3.1 build()

#### **Parameters**

type	- DMTP method. Available: CAMM.
wfn	- wavefunction
n	- number of DMTP sets

#### **Returns**

**DMTP** distribution

### 17.15.3.2 energy()

### **Parameters**

other	- interacting DMTP distribution.
max_clevel	- maximum convergence level (see below).

#### **Returns**

The generalized interaction energy convergence (A.U. units)

The following convergence levels are available:

- MultipoleConvergence::R1: includes qq terms.
- MultipoleConvergence::R2: includes dq terms and above.
- MultipoleConvergence::R3: includes qQ, dd terms and above.
- MultipoleConvergence::R4: includes qO, dQ terms and above.
- MultipoleConvergence:: R5: includes qH, dO, QQ terms and above.

## 17.15.3.3 potential()

#### **Parameters**

other	- interacting DMTP distribution.
max_clevel	- maximum convergence level (see below).

#### **Returns**

The generalized potential convergence (A.U. units)

The following convergence levels are available:

- MultipoleConvergence::R1: includes qq terms.
- MultipoleConvergence::R2: includes dq terms and above.
- MultipoleConvergence::R3: includes qQ, dd terms and above.
- MultipoleConvergence::R4: includes qO, dQ terms and above.
- MultipoleConvergence::R5: includes qH, dO, QQ terms and above.

#### 17.15.3.4 recenter()

Change origins of the distributed multipole moments of all sets

## **Parameters**

```
new_origins - matrix with coordinates of the new origins \{\mathbf{r}_{\mathrm{new}}\}.
```

Note

The number of origins has to be equal to the number of distributed centres.

Recentering of the multipoles affects the distributed dipoles and higher moments. The moments are given as

$$\begin{split} q_{\text{new}} &= q_{\text{old}} \\ \mu_{\text{new}} &= \mu_{\text{old}} - q_{\text{old}} \Delta^{(1)} \\ \Theta_{\text{new}} &= \Theta_{\text{old}} + q_{\text{old}} \Delta^{(2)} - \sum_{\mathscr{P}_2} \mathscr{P}_2 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}} + \mu_{\text{old}} \right) \otimes \Delta^{(1)} \right] \\ \Omega_{\text{new}} &= \Omega_{\text{old}} - q_{\text{old}} \Delta^{(3)} + \sum_{\mathscr{P}_3} \mathscr{P}_3 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}} + \mu_{\text{old}} \right) \otimes \Delta^{(2)} \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(1)} \right] \\ \Xi_{\text{new}} &= \Xi_{\text{old}} + q_{\text{old}} \Delta^{(4)} - \sum_{\mathscr{P}_3} \mathscr{P}_3 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}} + \mu_{\text{old}} \right) \otimes \Delta^{(3)} \right] + \sum_{\mathscr{P}_3} \mathscr{P}_3 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}}^2 + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}} + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \right) \otimes \Delta^{(2)} \right] \right] - \sum_{\mathscr{P}_6} \mathscr{P}_6 \left[ \left( q_{\text{old}} \mathbf{r}_{\text{old}} + \mu_{\text{old}} \otimes \mathbf{r}_{\text{old}} + \Theta_{\text{old}} \otimes \mathbf{r}_{\text{old}} \right) \otimes \Delta^{(2)} \right] \right]$$

where

$$\Delta^{(1)} \equiv \mathbf{r}_{\text{new}} - \mathbf{r}_{\text{old}}$$
 $\Delta^{(2)} \equiv \mathbf{r}_{\text{new}}^2 - \mathbf{r}_{\text{old}}^2$ 
 $\Delta^{(3)} \equiv \mathbf{r}_{\text{new}}^3 - \mathbf{r}_{\text{old}}^3$ 
 $\Delta^{(4)} \equiv \mathbf{r}_{\text{new}}^4 - \mathbf{r}_{\text{old}}^4$ 

In the above equations, the distributed centre label was omitted (redundant) as each distributed site of multipoles is independent of the others. TODO - Finish for octupoles and hexadecapoles! -> define the permutation operators!

#### 17.15.4 Friends And Related Function Documentation

#### 17.15.4.1 MultipoleConvergence

friend class MultipoleConvergence [friend]

Convergence of multipole moment series.

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

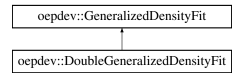
- oepdev/lib3d/dmtp.h
- oepdev/lib3d/dmtp\_base.cc

# 17.16 oepdev::DoubleGeneralizedDensityFit Class Reference

Generalized Density Fitting Scheme - Double Fit.

#include <oep\_gdf.h>

Inheritance diagram for oepdev::DoubleGeneralizedDensityFit:



#### **Public Member Functions**

- DoubleGeneralizedDensityFit (std::shared\_ptr< psi::BasisSet > bs\_auxiliary, std::shared\_ptr< psi::BasisSet > bs\_intermediate, std::shared\_ptr< psi::Matrix > v\_vector)
- std::shared\_ptr< psi::Matrix > compute (void)
   Perform the generalized density fit.

## **Additional Inherited Members**

## 17.16.1 Detailed Description

The density fitting map projects the OEP onto an arbitrary (not necessarily complete) auxiliary basis set space through application of the self energy minimization technique. The resulting three-electron repulsion integrals are computed by utilizing the resolution of identity in an intermediate, nearly-complete basis set space, hence performing an internal density fitting in nearly complete basis. Refer to density fitting specialized for OEP's for more details.

#### 17.16.2 Determination of the OEP matrix

Coefficients **G** are computed by using the following relation

$$G = A^{-1} \cdot R \cdot H$$

where the intermediate projection matrix is given by

$$\mathbf{H} = \mathbf{S}^{-1} \cdot \mathbf{V}$$

In the above equations,

$$A_{\xi\xi'} = (\xi||\xi')$$

$$R_{\xi\varepsilon} = (\xi||\varepsilon)$$

$$S_{\varepsilon\varepsilon'} = (\varepsilon|\varepsilon')$$

$$V^{\varepsilon i} = (\varepsilon|\hat{v}i)$$

The following labeling convention is used here:

• *i* denotes the arbitrary state vector

- $\xi$  denotes the auxiliary basis set element
- $\varepsilon$  denotes the intermediate (nearly complete) basis set element

In the above, | denotes the single integration over electron coordinate, i.e.,

$$(a|b) \equiv \int d\mathbf{r} \phi_a^*(\mathbf{r}) \phi_b(\mathbf{r})$$

whereas || acts as shown below:

$$(a||b) \equiv \iint d\mathbf{r}' d\mathbf{r}'' \frac{\phi_a^*(\mathbf{r}')\phi_b(\mathbf{r}'')}{|\mathbf{r}' - \mathbf{r}''|}$$

The spatial form of the potential operator  $\hat{v}$  can be expressed by

$$v(\mathbf{r}) \equiv \int d\mathbf{r}' \frac{\rho(\mathbf{r}')}{|\mathbf{r}' - \mathbf{r}|}$$

with  $\rho(\mathbf{r})$  being the effective one-electron density associated with  $\hat{v}$ .

#### 17.16.2.1 Theory behind the double GDF scheme

In order to perform the generalized density fitting in an incomplete auxiliary basis set, one must apply the following formula:

$$\mathbf{G} = \mathbf{A}^{-1} \cdot \mathbf{B}$$

where one encounters the need of evaluation of the following three-electron integrals

$$B_{\xi i} = (\xi || \hat{v}i) \equiv \iiint d\mathbf{r}' d\mathbf{r}'' d\mathbf{r}''' \phi_{\xi}^*(\mathbf{r}') \frac{1}{|\mathbf{r}' - \mathbf{r}''|} \rho(\mathbf{r}''') \frac{1}{|\mathbf{r}''' - \mathbf{r}''|} \phi_i(\mathbf{r}'')$$

Computation of all the necessery integrals of this kind is very costly and impractical for larger molecules. However, one can use the same trick that is a kernel of the OEP technique introduced in the OEPDev project, i.e., introduce the effective potential in order to get rid of one integration. This can be done by performing the generalized density fitting in the nearly complete intermediate basis

$$\hat{v}|i)\cong\sum_{m{arepsilon}}H_{m{arepsilon}i}|m{arepsilon})$$

Note that this is done just for the sake of factorizing the triple integral and computing the OEP matrix for the incomplete auxiliary basis. Therefore, the intermediate basis set is used just for a while during density fitting and is no longer necessary later on. By inserting the above identity to the triple integral one can transform it into a sum of the two-electron integrals that are much easier to evaluate. This leads to equations given in the beginning of this section.

#### 17.16.3 Member Function Documentation

#### 17.16.3.1 compute()

#### **Returns**

The OEP coefficients  $G_{\xi_i}$ 

Implements oepdev::GeneralizedDensityFit.

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

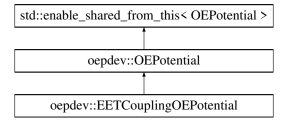
- oepdev/liboep/oep\_gdf.h
- oepdev/liboep/oep\_gdf.cc

## 17.17 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, SharedBasisSet intermediate, Options & options)
- **EETCouplingOEPotential** (SharedWavefunction wfn, Options & options)
- virtual void compute (const std::string &oepType) override

Compute matrix forms of all OEP's within a specified OEP type.

virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, std::shared\_ptr< psi::Vector > &v) override

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

virtual void print\_header () const override

## **Additional Inherited Members**

## 17.17.1 Detailed Description

Contains the following OEP types:

- Fujimoto.ET1
- Fujimoto.ET2
- Fujimoto.HT1
- Fujimoto.HT1
- Fujimoto.HT2
- Fujimoto.CT1
- Fujimoto.CT2

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

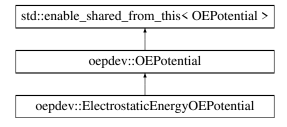
- oepdev/liboep/oep.h
- oepdev/liboep/oep\_coupling\_eet.cc

## 17.18 oepdev::ElectrostaticEnergyOEPotential Class Reference

Generalized One-Electron Potential for Electrostatic Energy.

```
#include <oep.h>
```

Inheritance diagram for oepdev::ElectrostaticEnergyOEPotential:



## **Public Member Functions**

- ElectrostaticEnergyOEPotential (SharedWavefunction wfn, Options & options)
   Only ESP-based potential is worth implementing.
- virtual void compute (const std::string &oepType) override
   Compute matrix forms of all OEP's within a specified OEP type.

virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, std::shared\_ptr< psi::Vector > &v) override

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

virtual void print\_header () const override

#### **Additional Inherited Members**

## 17.18.1 Detailed Description

Contains the following OEP types:

V

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

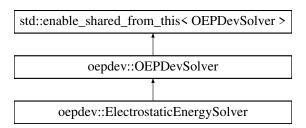
- oepdev/liboep/oep.h
- oepdev/liboep/oep\_energy\_coul.cc

## 17.19 oepdev::ElectrostaticEnergySolver Class Reference

Compute the Coulombic interaction energy between unperturbed wavefunctions.

```
#include <solver.h>
```

Inheritance diagram for oepdev::ElectrostaticEnergySolver:



## **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**

## 17.19.1 Detailed Description

The implemented methods are shown in below

Table 17.25: Methods available in the Solver

Keyword	Method Description	
	Day abwaady Mathada	
	Benchmark Methods	
AO_EXPANDED	*Default*. Exact Coulombic energy from atomic orbital expansions.	
MO_EXPANDED	Exact Coulombic energy from molecular orbital expansions	
	OEP-Based Methods	
ESP_SYMMETRIZED	*Default*. Coulombic energy from ESP charges interacting with nu-	
	clei and electronic density. Symmetrized with respect to monomers.	
CAMM	Coulombic energy from CAMM distributions.	

Below the detailed description of the above methods is given.

## **Benchmark Methods**

Exact Coulombic energy from atomic orbital expansions.

The Coulombic interaction energy is given by

$$E^{\text{Coul}} = E^{\text{Nuc-Nuc}} + E^{\text{Nuc-El}} + E^{\text{El-El}}$$

where the nuclear-nuclear repulsion energy is

$$E^{\text{Nuc-Nuc}} = \sum_{x \in A} \sum_{y \in R} \frac{Z_x Z_y}{|\mathbf{r}_x - \mathbf{r}_y|}$$

the nuclear-electronic attraction energy is

$$E^{\text{Nuc-El}} = \sum_{x \in A} \sum_{\lambda, \sigma \in B} Z_x V_{\lambda, \sigma}^{(x)} \left( D_{\lambda, \sigma}^{(\alpha)} + D_{\lambda, \sigma}^{(\beta)} \right) + \sum_{v \in B} \sum_{\mu, v \in A} Z_y V_{\mu, v}^{(v)} \left( D_{\mu, v}^{(\alpha)} + D_{\mu, v}^{(\beta)} \right)$$

and the electron-electron repulsion energy is

$$E^{\mathrm{El-El}} = \sum_{\mu\nu\in A} \sum_{\lambda\sigma\in B} \left\{ D_{\mu\nu}^{(\alpha)} + D_{\mu\nu}^{(\beta)} \right\} \left\{ D_{\lambda\sigma}^{(\alpha)} + D_{\lambda\sigma}^{(\beta)} \right\} (\mu\nu|\lambda\sigma)$$

In the above equations,

$$V_{\lambda\sigma}^{(x)} \equiv \int \frac{\varphi_{\lambda}^{*}(\mathbf{r})\varphi_{\sigma}(\mathbf{r})}{|\mathbf{r} - \mathbf{r}_{x}|} d\mathbf{r}$$

Exact Coulombic energy from molecular orbital expansion.

This approach is fully equivalent to the atomic orbital expansion shown above. For the closed shell case, the Coulombic interaction energy is given by

$$E^{\text{Coul}} = E^{\text{Nuc-Nuc}} + E^{\text{Nuc-El}} + E^{\text{El-El}}$$

where the nuclear-nuclear repulsion energy is

$$E^{\text{Nuc-Nuc}} = \sum_{x \in A} \sum_{y \in B} \frac{Z_x Z_y}{|\mathbf{r}_x - \mathbf{r}_y|}$$

the nuclear-electronic attraction energy is

$$E^{\text{Nuc-El}} = 2 \sum_{i \in A} \sum_{y \in B} V_{ii}^{(y)} + 2 \sum_{j \in B} \sum_{x \in A} V_{jj}^{(x)}$$

and the electron-electron repulsion energy is

$$E^{\text{El-El}} = 4\sum_{i \in A} \sum_{j \in B} (ii|jj)$$

## **OEP-Based Methods**

Coulombic energy from ESP charges interacting with nuclei and electronic density.

In this approach, nuclear and electronic density of either species is approximated by ESP charges. In order to achieve symmetric expression, the interaction is computed twice (ESP of A interacting with density matrix and nuclear charges of B and vice versa) and then divided by 2. Thus,

$$E^{\text{Coul}} \approx \frac{1}{2} \left[ \sum_{x \in A} \sum_{y \in B} \frac{Z_x q_y}{|\mathbf{r}_x - \mathbf{r}_y|} + \sum_{y \in B} \sum_{\mu \nu \in A} q_y V_{\mu \nu}^{(y)} \left( D_{\mu \nu}^{(\alpha)} + D_{\mu \nu}^{(\beta)} \right) + \sum_{y \in B} \sum_{x \in A} \frac{q_x Z_y}{|\mathbf{r}_x - \mathbf{r}_y|} + \sum_{x \in A} \sum_{\lambda \sigma \in B} q_x V_{\lambda \sigma}^{(x)} \left( D_{\lambda \sigma}^{(\alpha)} + D_{\lambda \sigma}^{(\beta)} \right) \right]$$

If the basis set is large and the number of ESP centres  $q_{x(y)}$  is sufficient, the sum of first two contributions equals the sum of the latter two contributions.

Notes:

• This solver also computes and prints the ESP-ESP point charge interaction energy,

$$E^{\text{Coul,ESP}} \approx \sum_{x \in A} \sum_{y \in B} \frac{q_x q_y}{|\mathbf{r}_x - \mathbf{r}_y|}$$

for reference purposes.

• In order to construct this solver, **always** use the OEPDevSolver::build static factory method.

#### 17.19.2 Member Function Documentation

## 17.19.2.1 compute\_benchmark()

Each solver object has one DEFAULT benchmark method

#### **Parameters**

```
method - benchmark method
```

Implements oepdev::OEPDevSolver.

## 17.19.2.2 compute\_oep\_based()

Each solver object has one DEFAULT OEP-based method.

#### **Parameters**

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

Implements oepdev::OEPDevSolver.

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

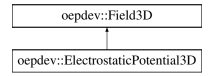
- oepdev/libsolver/solver.h
- oepdev/libsolver/solver\_energy\_coul.cc

# 17.20 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 &padding, psi::SharedWavefunction wfn, psi::Options &options)
- ElectrostaticPotential3D (const int &nx, const int &ny, const int &nz, const double &px, const double &py, const double &pz, psi::SharedWavefunction wfn, psi::Options &options)
- virtual std::shared\_ptr< psi::Vector > compute\_xyz (const double &x, const double &y, const double &z)

Compute a value of 3D field at point (x, y, z)

virtual void print () const

Print information of the object to Psi4 output.

#### **Additional Inherited Members**

## 17.20.1 Detailed Description

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

$$v_{
m nuc}(\mathbf{r}) = \sum_{x} rac{Z_x}{|\mathbf{r} - \mathbf{r}_x|}$$
 $v_{
m el}(\mathbf{r}) = \sum_{\mu\nu} \left\{ D_{\mu\nu}^{(lpha)} + D_{\mu\nu}^{(eta)} 
ight\} V_{
u\mu}(\mathbf{r})$ 

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

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

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

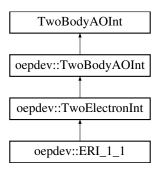
- oepdev/lib3d/space3d.h
- oepdev/lib3d/space3d.cc

# 17.21 oepdev::ERI\_1\_1 Class Reference

2-centre ERI of the form (a|O(2)|b) where O(2) = 1/r12.

#include <eri.h>

Inheritance diagram for oepdev::ERI\_1\_1:



## **Public Member Functions**

- ERI\_1\_1 (const IntegralFactory \*integral, int deriv=0, bool use\_shell\_pairs=false)
   Constructor. Use oepdev::IntegralFactory to generate this object.
- ~ERI\_1\_1 ()
   Destructor.

## **Protected Member Functions**

size\_t compute\_doublet (int, int)
 Compute ERI's between 2 shells.

## **Protected Attributes**

- double \* mdh\_buffer\_1\_
   Buffer for McMurchie-Davidson-Hermite coefficents for monomial expansion (shell 1)
- double \* mdh\_buffer\_2\_
   Buffer for McMurchie-Davidson-Hermite coefficents for monomial expansion (shell 2)

## 17.21.1 Detailed Description

ERI's are computed for a shell doublet (P|Q) and stored in the <code>target\_full\_buffer</code>, accessible through <code>buffer()</code> method:

For each 
$$(n_1,l_1,m_1)\in P$$
:  
For each  $(n_2,l_2,m_2)\in Q$ :  
 $\mathrm{ERI}=(A|B)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}]$ 

For detailed description of the McMurchie-Davidson scheme, refer to The Integral Package Library.

## 17.21.2 Implementation

A set of ERI's in a shell is decontracted as

$$(A|B)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}] = \sum_{ij} c_i(\alpha_1)c_j(\alpha_2)(i|j)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}]$$

where the primitive ERI is given by

$$(i|j)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}] = \sum_{N_1=0}^{n_1} \sum_{L_1=0}^{l_1} \sum_{M_1=0}^{m_1} \sum_{N_2=0}^{n_2} \sum_{L_2=0}^{l_2} \sum_{M_2=0}^{m_2} d_{N_1}^{n_1} d_{L_1}^{l_1} d_{M_1}^{m_1} d_{N_2}^{n_2} d_{L_2}^{l_2} d_{M_2}^{m_2} [N_1 L_1 M_1 | N_2 L_2 M_2]$$

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

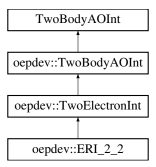
- · oepdev/libints/eri.h
- · oepdev/libints/eri.cc

## 17.22 oepdev::ERI\_2\_2 Class Reference

4-centre ERI of the form (ab|O(2)|cd) where O(2) = 1/r12.

#include <eri.h>

Inheritance diagram for oepdev::ERI\_2\_2:



### **Public Member Functions**

- ERI\_2\_2 (const IntegralFactory \*integral, int deriv=0, bool use\_shell\_pairs=false)

  Constructor. Use oepdev::IntegralFactory to generate this object.
- ∼ERI\_2\_2 ()

Destructor.

## **Protected Member Functions**

size\_t compute\_quartet (int, int, int, int)
 Compute ERI's between 4 shells.

#### **Protected Attributes**

double \* mdh\_buffer\_12\_

Buffer for McMurchie-Davidson-Hermite coefficents for binomial expansion (shells 1 and 2)

double \* mdh\_buffer\_34\_

Buffer for McMurchie-Davidson-Hermite coefficents for binomial expansion (shells 3 and 4)

## 17.22.1 Detailed Description

ERI's are computed for a shell quartet (PQ|RS) and stored in the target\_full\_buffer, accessible through buffer() method:

For each 
$$(n_1,l_1,m_1)\in P$$
:
For each  $(n_2,l_2,m_2)\in Q$ :
For each  $(n_3,l_3,m_3)\in R$ :
For each  $(n_4,l_4,m_4)\in S$ :
 $\mathrm{ERI}=(AB|CD)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}]$ 

For detailed description of the McMurchie-Davidson scheme, refer to The Integral Package Library.

## 17.22.2 Implementation

A set of ERI's in a shell is decontracted as

$$(AB|CD)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}] = \sum_{ijkl} c_i(\alpha_1)c_j(\alpha_2)c_k(\alpha_3)c_l(\alpha_4)(ij|kl)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}]$$

where the primitive ERI is given by

$$(ij|kl)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}] = E_{ij}(\alpha_1,\alpha_2)E_{kl}(\alpha_3,\alpha_4)$$

$$\times \sum_{N_1=0}^{n_1+n_2} \sum_{L_1=0}^{l_1+l_2} \sum_{M_1=0}^{m_1+m_2} \sum_{N_2=0}^{n_3+n_4} \sum_{L_2=0}^{l_3+l_4} \sum_{M_2=0}^{m_1n_2} d_{N_1}^{l_1l_2} d_{M_1}^{m_1m_2} d_{N_2}^{n_3n_4} d_{L_2}^{l_3l_4} d_{M_2}^{m_3m_4} [N_1L_1M_1|N_2L_2M_2]$$

In the above equation, the multiplicative constants are given as

$$E_{ij}(\alpha_1, \alpha_2) = \exp\left[-\frac{\alpha_1 \alpha_2}{\alpha_1 + \alpha_2} |\mathbf{A} - \mathbf{B}|^2\right]$$
$$E_{kl}(\alpha_3, \alpha_4) = \exp\left[-\frac{\alpha_3 \alpha_4}{\alpha_3 + \alpha_4} |\mathbf{C} - \mathbf{D}|^2\right]$$

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

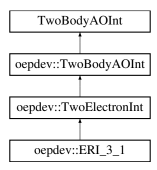
- · oepdev/libints/eri.h
- · oepdev/libints/eri.cc

## 17.23 oepdev::ERI\_3\_1 Class Reference

4-centre ERI of the form (abc|O(2)|d) where O(2) = 1/r12.

#include <eri.h>

Inheritance diagram for oepdev::ERI\_3\_1:



#### **Public Member Functions**

- ERI\_3\_1 (const IntegralFactory \*integral, int deriv=0, bool use\_shell\_pairs=false)
   Constructor. Use oepdev::IntegralFactory to generate this object.
- ∼ERI\_3\_1 ()

Destructor.

#### **Protected Member Functions**

• size\_t compute\_quartet (int, int, int, int)

Compute ERI's between 4 shells.

## **Protected Attributes**

double \* mdh\_buffer\_123\_

Buffer for McMurchie-Davidson-Hermite coefficents for trinomial expansion (shells 1, 2 and 3)

double \* mdh\_buffer\_4\_

Buffer for McMurchie-Davidson-Hermite coefficents for monomial expansion (shell 4)

## 17.23.1 Detailed Description

ERI's are computed for a shell quartet (PQR|S) and stored in the target\_full\_buffer, accessible through buffer() method:

For each 
$$(n_1,l_1,m_1)\in P$$
:  
For each  $(n_2,l_2,m_2)\in Q$ :  
For each  $(n_3,l_3,m_3)\in R$ :  
For each  $(n_4,l_4,m_4)\in S$ :  
 $\mathrm{ERI}=(ABC|D)[\{\alpha\},\mathbf{n},\mathbf{l},\mathbf{m}]$ 

For detailed description of the McMurchie-Davidson scheme, refer to The Integral Package Library.

## 17.23.2 Implementation

A set of ERI's in a shell is decontracted as

$$(ABC|D)[\{\alpha\}, \mathbf{n}, \mathbf{l}, \mathbf{m}] = \sum_{ijkl} c_i(\alpha_1)c_j(\alpha_2)c_k(\alpha_3)c_l(\alpha_4)(ijk|l)[\{\alpha\}, \mathbf{n}, \mathbf{l}, \mathbf{m}]$$

where the primitive ERI is given by

$$(ijk|l)[\{\boldsymbol{\alpha}\},\mathbf{n},\mathbf{l},\mathbf{m}] = E_{ijk}(\alpha_1,\alpha_2,\alpha_3) \times \sum_{N_1=0}^{n_1+n_2+n_3} \sum_{L_1=0}^{l_1+l_2+l_3} \sum_{M_1=0}^{m_1+m_2+m_3} \sum_{N_2=0}^{n_4} \sum_{L_2=0}^{m_4} \sum_{M_2=0}^{m_4} d_{N_1}^{n_1n_2n_3} d_{L_1}^{l_1l_2l_3} d_{M_1}^{m_1m_2m_3} d_{N_2}^{n_4} d_{L_2}^{l_4} d_{M_2}^{m_4} [N_1L_1M_1|N_2L_2M_2]$$

In the above equation, the multiplicative constants are given as

$$E_{ijk}(\alpha_1, \alpha_2, \alpha_3) = \exp\left[-\frac{\alpha_1 \alpha_2}{\alpha_1 + \alpha_2} |\mathbf{A} - \mathbf{B}|^2\right] \exp\left[-\frac{(\alpha_1 + \alpha_2) \alpha_3}{\alpha_1 + \alpha_2 + \alpha_3} |\mathbf{P} - \mathbf{C}|^2\right]$$

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

- · oepdev/libints/eri.h
- · oepdev/libints/eri.cc

# 17.24 oepdev::ESPSolver Class Reference

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

#### **Public Member Functions**

ESPSolver (SharedField3D field)

Construct from 3D vector field.

• ESPSolver (SharedField3D field, psi::SharedMatrix centres)

Construct from 3D vector field.

virtual ∼ESPSolver ()

Destructor.

virtual psi::SharedMatrix charges () const

Get the (fit) charges.

virtual psi::SharedMatrix centres () const

Get the charge distribution centres.

virtual void set\_charge\_sums (psi::SharedVector s)

Set the charge sums  $Q_p$ .

virtual void set\_charge\_sums (const double &s)

Set the charge sums  $Q_p$  (equal to all fields)

virtual void compute ()

Perform fitting of effective charges.

#### **Protected Attributes**

const int nCentres\_

Number of fit centres.

const int nFields\_

Number of fields to fit.

SharedField3D field\_

Scalar field.

psi::SharedMatrix charges\_

Charges to be fit.

psi::SharedMatrix centres\_

Centres, at which fit charges will reside.

psi::SharedVector charge\_sums\_

Vector of sums of partial charges.

## 17.24.1 Detailed Description

Solves the least-squares problem to fit the generalized charges  $q_{m;p}$ , that reproduce the reference generalized potential  $v_p^{\rm ref}({\bf r})$  supplied by the Field3D object:

$$\int d\mathbf{r}' \left[ v_p^{\rm ref}(\mathbf{r}') - \sum_m \frac{q_{m;p}}{|\mathbf{r}' - \mathbf{r}_m|} \right]^2 \to {\rm minimize}$$

The charges are subject to the following constraint:

$$\sum_m q_{m;p} = Q_p$$
 for all  $p$ 

Method description.

M generalized charges is found by solving the matrix equation

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

where the **A** matrix of dimension  $(M+1) \times (M+1)$  and  $\mathbf{b}_p$  vector or length M+1 are given as

$$A_{mn} = \sum_{i} \frac{1}{r_{im}r_{in}}$$

$$b_{m;p} = \sum_{i} \frac{v_{p}^{\text{ref}}(\mathbf{r}_{m})}{r_{im}}$$

In the above equation, summations run over all sample points, at which reference potential is known. The solution is stored in the  $M \times N$  matrix, where N is the dimensionality of the 3D vector field (i.e., the number of potentials supplied,  $p_{\rm max}$ ). As a default,  $Q_p=0$  for all potentials. This can be set by oepdev::ESPSolver::set\_charge\_sums method.

Note

#### Useful options:

- ESP\_PAD\_SPHERE Padding spherical radius for random points selection. Default: 10.0 [A.U.]
- ESP\_NPOINTS\_PER\_ATOM Number of random points per atom in a molecule. Detault: 1500
- ESP\_VDW\_RADIUS\_C The vdW radius for carbon atom. Default: 3.0 [A.U.]
- ESP\_VDW\_RADIUS\_H The vdW radius for hydrogen atom. Default: 4.0 [A.U.]
- ESP\_VDW\_RADIUS\_N The vdW radius for nitrogen atom. Default: 2.4 [A.U.]
- ESP\_VDW\_RADIUS\_O The vdW radius for oxygen atom. Default: 5.6 [A.U.]
- ESP\_VDW\_RADIUS\_F The vdW radius for fluorium atom. Default: 2.3 [A.U.]
- ESP\_VDW\_RADIUS\_CL The vdW radius for chlorium atom. Default: 2.9 [A.U.]

#### 17.24.2 Constructor & Destructor Documentation

```
17.24.2.1 ESPSolver() [1/2]

oepdev::ESPSolver::ESPSolver (

SharedField3D field)
```

Assume that the centres are on atoms associated with the 3D vector field.

#### **Parameters**

field
-------

## **17.24.2.2 ESPSolver()** [2/2]

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

#### **Parameters**

field	- oepdev 3D vector field object
centres	- matrix with coordinates of charge distribution centres

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

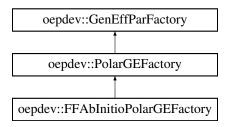
- · oepdev/lib3d/esp.h
- · oepdev/lib3d/esp.cc

# 17.25 oepdev::FFAbInitioPolarGEFactory Class Reference

Polarization GEFP Factory from First Principles: Finite-Difference Model. Arbitrary level of theory.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::FFAbInitioPolarGEFactory:



#### **Public Member Functions**

- FFAbInitioPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
- virtual std::shared\_ptr< GenEffPar > compute (void)
   Compute the density matrix susceptibility tensors.

#### **Additional Inherited Members**

## 17.25.1 Detailed Description

Implements creation of the density matrix susceptibility tensors. Does not guarantee the idempotency of the density matrix in LCAO-MO variation, but for weak electric fields the idempotency is to be expected up to first order. The density matrix susceptibility tensor is represented by:

$$\delta D_{\alpha\beta} = \mathbf{B}_{\alpha\beta}^{(1)} \cdot \mathbf{F} + \mathbf{B}_{\alpha\beta}^{(2)} : \mathbf{F} \otimes \mathbf{F}$$

where  $\mathbf{B}_{lphaeta}^{(1)}$  is the density matrix dipole polarizability defined as

$$\mathbf{B}_{\alpha\beta}^{(1)} = \frac{\partial D_{\alpha\beta}}{\partial \mathbf{F}} \Big|_{\mathbf{F}=\mathbf{0}}$$

whereas  $\mathbf{B}_{\alpha\beta}^{(2)}$  is the density matrix dipole-dipole hyperpolarizability,

$$\mathbf{B}_{\alpha\beta}^{(2)} = \frac{1}{2} \frac{\partial^2 D_{\alpha\beta}}{\partial \mathbf{F} \otimes \partial \mathbf{F}} \Big|_{\mathbf{F} = \mathbf{0}}$$

The first derivative is evaluated numerically from central finite-field 3-point formula,

$$f' = \frac{f(h) - f(-h)}{2h} + \mathfrak{O}(h^2)$$

where h is the differentiation step. Second derivatives are evaluated from the following formulae:

$$f_{uu} = \frac{f(h) + f(-h) - 2f(0)}{h^2} + \mathfrak{O}(h^2)$$

$$f_{uw} = \frac{f(h,h) + f(-h,-h) + 2f(0) - f(h,0) - f(-h,0) - f(0,h) - f(0,-h)}{2h^2} + \mathfrak{O}(h^2)$$

As long as the second-order susceptibility is considered, this susceptibility model works well for uniform weak, moderate and strong electric fields.

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

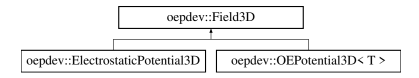
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_ffabinitio.cc

# 17.26 oepdev::Field3D Class Reference

General Vector Dield in 3D Space. Abstract base.

#include <space3d.h>

Inheritance diagram for oepdev::Field3D:



# **Public Member Functions**

 Field3D (const int &ndim, const int &np, const double &pad, psi::SharedWavefunction wfn, psi::Options &options)

Construct potential on random grid by providing wavefunction. Excludes space within vdW volume.

 Field3D (const int &ndim, const int &nx, const int &ny, const int &nz, const double &px, const double &py, const double &pz, std::shared\_ptr< psi::Wavefunction > wfn, psi::Options &options)

Construct potential on cube grid by providing wavefunction.

virtual ∼Field3D ()

Destructor.

virtual int npoints () const

Get the number of points at which the 3D 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  $\{[x, y, z, f_1(x, y, z), f_2(x, y, z), ..., f_n(x, y, z)]\}$  where n = ndim.

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.

int dimension () const

Get the number of fields.

virtual void compute ()

Compute the 3D field in each point from the point collection.

virtual std::shared\_ptr< psi::Vector > compute\_xyz (const double &x, const double &y, const double &z)=0

Compute a value of 3D 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< Field3D > build (const std::string &type, const int &np, const double &pad, psi::SharedWavefunction wfn, psi::Options &options, const int &ndim=1)

Build 3D field of random points. vdW volume is excluded.

static shared\_ptr< Field3D > build (const std::string &type, const int &nx, const int &ny, const int &nz, const double &px, const double &py, const double &pz, psi::SharedWavefunction wfn, psi::Options &options, const int &ndim=1)

Build 3D field of points on a g09-cube grid.

# **Protected Attributes**

std::shared\_ptr< PointsCollection3D > pointsCollection\_

Collection of points at which the 3D field is to be computed.

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

The data matrix in a form  $\{[x, y, z, f_1(x, y, z), f_2(x, y, z), ..., f_n(x, y, z)]\}$  where  $n = nDim_1$ .

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.

int nDim\_

Dimensionality of the 3D field (1: scalar field, 2>: vector field)

bool isComputed\_

Has data already computed?

# 17.26.1 Detailed Description

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

- Electrostatic potential computes electrostatic potential (requires wavefunction)
- Template of generic classes compute custom vector 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 following types of 3D vector fields are currently implemented:

• ELECTROSTATIC POTENTIAL

# 17.26.2 Constructor & Destructor Documentation

# 17.26.2.1 Field3D()

Construct potential on random grid by providing molecule. Excludes space within vdW volume Field3D(const int& ndim, const int& np, const double& pad, psi::SharedMolecule mol, psi::Options& options);

# 17.26.3 Member Function Documentation

# **17.26.3.1** build() [1/2]

# **Parameters**

ndim	- dimensionality of 3D field (1: scalar field, >2: vector field)	
type	- type of 3D field	
np	- number of points	
pad	- radius padding of a minimal sphere enclosing the molecule	
wfn	- Psi4 Wavefunction containing the molecule	
options	- Psi4 options	

# **17.26.3.2** build() [2/2]

```
std::shared_ptr< Field3D > oepdev::Field3D::build (
    const std::string & type,
    const int & nx,
    const int & ny,
    const int & nz,
    const double & px,
    const double & py,
    const double & pz,
    psi::SharedWavefunction wfn,
    psi::Options & options,
    const int & ndim = 1 ) [static]
```

#### **Parameters**

ndim	- dimensionality of 3D field (1: scalar field, >2: vector field)
type	- type of 3D field
nx	- number of points along x direction
ny	- number of points along y direction
nz	- number of points along z direction
рх	- padding distance along x direction
ру	- padding distance along y direction
pz	- padding distance along z direction
wfn	- Psi4 Wavefunction containing the molecule
options	- Psi4 options

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

- · oepdev/lib3d/space3d.h
- oepdev/lib3d/space3d.cc

# 17.27 oepdev::Fourier9 Struct Reference

Simple structure to hold the Fourier series expansion coefficients for N=4.

```
#include <unitary_optimizer.h>
```

# **Public Attributes**

- double a0
- · double a1
- · double a2
- · double a3
- · double a4

- double b1
- double b2
- double b3
- double b4

# 17.27.1 Detailed Description

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

· oepdev/libutil/unitary\_optimizer.h

# 17.28 oepdev::GenEffFrag Class Reference

Generalized Effective Fragment. Container Class.

```
#include <gefp.h>
```

# **Public Member Functions**

· GenEffFrag ()

Initialize with default name of GEFP (Default)

• GenEffFrag (std::string name)

Initialize with custom name of GEFP.

∼GenEffFrag ()

Destruct.

void rotate (std::shared\_ptr< psi::Matrix > R)

Rotate.

void translate (std::shared\_ptr< psi::Vector > T)

Translate.

- void superimpose (std::shared\_ptr< psi::Matrix > targetXYZ, std::vector< int > supList)
   Superimpose.
- void set\_gefp\_polarization (const std::shared\_ptr< GenEffPar > &par)

Set the Density Matrix Susceptibility Tensor Object.

void set\_dmat\_dipole\_polarizability (const std::vector< std::vector< std::shared\_ptr<
psi::Matrix >>> &susc)

Set the Density Matrix Dipole Polarizability.

void set\_dmat\_dipole\_dipole\_hyperpolarizability (const std::vector< std::vector< std::vector< std::shared\_ptr< psi::Matrix >>> &susc)

Set the Density Matrix Dipole-Dipole Hyperpolarizability.

void set\_dmat\_quadrupole\_polarizability (const std::vector< std::vector< std::shared\_ptr<</li>
 psi::Matrix >>> &susc)

Set the Density Matrix Quadrupole Polarizability.

std::shared\_ptr< psi::Matrix > susceptibility (int fieldRank, int fieldGradientRank, int i, int x) const

Grab the Density Matrix Susceptibility.

std::vector< std::shared\_ptr< psi::Matrix >> susceptibility (int fieldRank, int fieldGradientRank, int i) const

Grab the Density Matrix Susceptibility.

std::vector < std::vector < std::shared\_ptr < psi::Matrix > > susceptibility (int fieldRank, int fieldGradientRank) const

Grab the Density Matrix Susceptibility.

# **Public Attributes**

std::map< std::string, std::shared\_ptr< GenEffPar >> parameters
 Dictionary of All GEF Parameters.

# **Protected Attributes**

std::string name\_

Name of GEFP.

std::shared\_ptr< GenEffPar > densityMatrixSusceptibilityGEF\_

Density Matrix Susceptibility Tensor.

std::shared\_ptr< GenEffPar > electrostaticEnergyGEF\_

Electrostatic Energy Effective One-Electron Potential.

std::shared\_ptr< GenEffPar > repulsionEnergyGEF\_

Exchange-Repulsion Effective One-Electron Potential.

std::shared\_ptr< GenEffPar > chargeTransferEnergyGEF\_

Charge-Transfer Effective One-Electron Potential.

std::shared\_ptr< GenEffPar > EETCouplingConstantGEF\_

EET Coupling Effective One-Electron Potential.

# 17.28.1 Detailed Description

Describes the GEFP fragment that is in principle designed to work at correlated levels of theory.

# 17.28.2 Member Function Documentation

# **17.28.2.1** susceptibility() [1/3]

```
std::shared_ptr<psi::Matrix> oepdev::GenEffFrag::susceptibility (
    int fieldRank,
    int fieldGradientRank,
    int i,
    int x ) const [inline]
```

#### **Parameters**

fieldRank	- power dependency with respect to the electric field
fieldGradientRank	- power dependency with respect to the electric field gradient
i	- id of the distributed site
X	- id of the composite Cartesian component

# 17.28.2.2 susceptibility() [2/3]

```
std::vector<std::shared_ptr<psi::Matrix> > oepdev::GenEffFrag::susceptibility
(
          int fieldRank,
          int fieldGradientRank,
          int i) const [inline]
```

# **Parameters**

fieldRank	- power dependency with respect to the electric field
fieldGradientRank	- power dependency with respect to the electric field gradient
i	- id of the distributed site

# **17.28.2.3** susceptibility() [3/3]

```
std::vector<std::vector<std::shared_ptr<psi::Matrix> > oepdev::GenEffFrag::susceptib.
(
    int fieldRank,
    int fieldGradientRank) const [inline]
```

# **Parameters**

fieldRank	- power dependency with respect to the electric field
fieldGradientRank	- power dependency with respect to the electric field gradient

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

- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp.cc

# 17.29 oepdev::GenEffPar Class Reference

Generalized Effective Fragment Parameters. Container Class.

```
#include <gefp.h>
```

#### **Public Member Functions**

GenEffPar (std::string name)

Create with name of this parameter type.

∼GenEffPar ()

Destruct.

void set\_susceptibility (int fieldRank, int fieldGradientRank, const std::vector< std::vector< std::shared\_ptr< psi::Matrix >>> &susc)

Set the Density Matrix Susceptibility.

void set\_dipole\_polarizability (const std::vector< std::vector< std::shared\_ptr< psi::Matrix</li>
 >>> &susc)

Set The Density Matrix Dipole Polarizability.

void set\_dipole\_hyperpolarizability (const std::vector< std::vector< std::shared\_ptr<</li>
 psi::Matrix >>> &susc)

Set The Density Matrix Dipole-Dipole Hyperpolarizability.

void set\_quadrupole\_polarizability (const std::vector< std::vector< std::shared\_ptr< psi::Matrix >>> &susc)

Set The Density Matrix Quadrupole Polarizability.

void set\_centres (const std::vector < std::shared\_ptr < psi::Vector >> &centres)

Set the distributed centres' positions.

void allocate (int fieldRank, int fieldGradientRank, int nsites, int nbf)

Allocate the Density Matrix Susceptibility.

void allocate\_dipole\_polarizability (int nsites, int nbf)

Allocate The Density Matrix Dipole Polarizability.

void allocate\_dipole\_dipole\_hyperpolarizability (int nsites, int nbf)

Allocate The Density Matrix Dipole-Dipole Hyperpolarizability.

void allocate\_quadrupole\_polarizability (int nsites, int nbf)

Allocate The Density Matrix Quadrupole Polarizability.

- bool hasDensityMatrixDipolePolarizability () const
- bool hasDensityMatrixDipoleDipoleHyperpolarizability () const
- bool hasDensityMatrixQuadrupolePolarizability () const
- std::shared\_ptr< psi::Matrix > susceptibility (int fieldRank, int fieldGradientRank, int i, int x) const

Grab the Density Matrix Susceptibility.

std::vector< std::shared\_ptr< psi::Matrix >> susceptibility (int fieldRank, int fieldGradientRank, int i) const

Grab the Density Matrix Susceptibility.

std::vector < std::vector < std::shared\_ptr < psi::Matrix > > susceptibility (int fieldRank, int fieldGradientRank) const

Grab the Density Matrix Susceptibility.

- std::vector < std::shared\_ptr < psi::Matrix >> > dipole\_polarizability () const
   Grab the density matrix dipole polarizability tensor.
- std::vector< std::shared\_ptr< psi::Matrix >> dipole\_polarizability (int i) const
   Grab the density matrix dipole polarizability tensor's x-th component.
- std::shared\_ptr< psi::Matrix > dipole\_polarizability (int i, int x) const
   Grab the density matrix dipole polarizability tensor's x-th component of the i-th distributed site.
- std::vector< std::shared\_ptr< psi::Matrix >> > dipole\_dipole\_hyperpolarizability
   () const

Grab the density matrix dipole-dipole hyperpolarizability tensor.

- std::vector< std::shared\_ptr< psi::Matrix >> dipole\_dipole\_hyperpolarizability (int i) const

  Grab the density matrix dipole-dipole hyperpolarizability tensor's x-th component.
- std::shared\_ptr< psi::Matrix > dipole\_dipole\_hyperpolarizability (int i, int x) const
   Grab the density matrix dipole-dipole hyperpolarizability tensor's x-th component of the i-th distributed site.
- std::vector< std::shared\_ptr< psi::Matrix >> > quadrupole\_polarizability () const

Grab the density matrix quadrupole polarizability tensor.

- std::vector< std::shared\_ptr< psi::Matrix >> quadrupole\_polarizability (int i) const
   Grab the density matrix quadrupole polarizability tensor's x-th component.
- std::shared\_ptr< psi::Matrix > quadrupole\_polarizability (int i, int x) const
   Grab the density matrix quadrupole polarizability tensor's x-th component of the i-th distributed
- std::vector < std::shared\_ptr < psi::Vector > > centres () const
   Grab the centres' positions.
- std::shared\_ptr< psi::Vector > centre (int i) const
   Grab the position of the i-th distributed site.
- std::shared\_ptr< psi::Matrix > compute\_density\_matrix (std::shared\_ptr< psi::Vector > field)

Compute the density matrix due to the uniform electric field perturbation.

- std::shared\_ptr< psi::Matrix > compute\_density\_matrix (double fx, double fy, double fz)

  Compute the density matrix due to the uniform electric field perturbation.
- std::shared\_ptr< psi::Matrix > compute\_density\_matrix (std::vector< std::shared\_ptr< psi::Vector >> fields)

Compute the density matrix due to the non-uniform electric field perturbation.

std::shared\_ptr< psi::Matrix > compute\_density\_matrix (std::vector< std::shared\_ptr< psi::Vector >> fields, std::vector< std::shared\_ptr< psi::Matrix >> grads)

Compute the density matrix due to the non-uniform electric field perturbation.

# **Protected Attributes**

std::string name\_

The Name of Parameter Type.

 std::vector< std::vector< std::shared\_ptr< psi::Matrix >> > densityMatrixDipolePolarizability\_

The Density Matrix Dipole Polarizability.

 std::vector< std::shared\_ptr< psi::Matrix >> > densityMatrixDipoleDipole-Hyperpolarizability\_

The Density Matrix Dipole-Dipole Hyperpolarizability.

 std::vector< std::shared\_ptr< psi::Matrix >>> densityMatrixQuadrupole-Polarizability\_

The Density Matrix Quadrupole Polarizability.

std::vector < std::shared\_ptr < psi::Vector > > distributedCentres\_

The Positions of the Distributed Centres.

- bool hasDensityMatrixDipolePolarizability\_
- bool hasDensityMatrixDipoleDipoleHyperpolarizability\_
- bool hasDensityMatrixQuadrupolePolarizability\_

# 17.29.1 Detailed Description

# 17.29.2 Member Function Documentation

# 17.29.2.1 allocate()

# **Parameters**

fieldRank	- power dependency with respect to the electric field ${f F}$
	hh
fieldGradientRank	- power dependency with respect to the electric field gradient $ abla\otimes \mathbf{F}$
nsites	- number of distributed sites
nbf	- number of basis functions in the basis set

The following susceptibilities are supported (fieldRank, fieldGradientRank):

- (1, 0) dipole polarizability, interacts with **F**
- (2, 0) dipole-dipole hyperpolarizability, interacts with  ${f F}\otimes {f F}$
- (0, 1) quadrupole polarizability, interacts with  $\nabla \otimes \mathbf{F}$

# **17.29.2.2** compute\_density\_matrix() [1/4]

#### **Parameters**

*field* - the uniform electric field vector (A.U.)

# 17.29.2.3 compute\_density\_matrix() [2/4]

#### **Parameters**

fx	- x-th Cartesian component of the uniform electric field vector (A.U.)
fy	- y-th Cartesian component of the uniform electric field vector (A.U.)
fz	- z-th Cartesian component of the uniform electric field vector (A.U.)

# **17.29.2.4** compute\_density\_matrix() [3/4]

# **Parameters**

fields	- the list of non-uniform electric field vector (A.U.) evaluated at the distributed
	DMatPol sites

# 17.29.2.5 compute\_density\_matrix() [4/4]

```
std::shared_ptr< psi::Matrix > oepdev::GenEffPar::compute_density_matrix (
    std::vector< std::shared_ptr< psi::Vector >> fields,
    std::vector< std::shared_ptr< psi::Matrix >> grads )
```

#### **Parameters**

fields	- the list of electric field vectors (A.U.) evaluated at the distributed DMatPol sites
grads	- the list of electric field gradient matrices (A.U.) evaluated at the distributed DMatPol
	sites

# 17.29.2.6 set\_susceptibility()

#### **Parameters**

fieldRank	- power dependency with respect to the electric field ${f F}$
fieldGradientRank	- power dependency with respect to the electric field gradient $ abla\otimes \mathbf{F}$
susc	- the susceptibility tensor

The following susceptibilities are supported (fieldRank, fieldGradientRank):

- (1, 0) dipole polarizability, interacts with **F**
- (2, 0) dipole-dipole hyperpolarizability, interacts with  $F \otimes F$
- (0, 1) quadrupole polarizability, interacts with  $\nabla \otimes F$

# **17.29.2.7** susceptibility() [1/3]

```
std::shared_ptr<psi::Matrix> oepdev::GenEffPar::susceptibility (
    int fieldRank,
    int fieldGradientRank,
    int i,
    int x ) const [inline]
```

#### **Parameters**

fieldRank	- power dependency with respect to the electric field
fieldGradientRank	- power dependency with respect to the electric field gradient
i	- id of the distributed site
X	- id of the composite Cartesian component

The following susceptibilities are supported (fieldRank, fieldGradientRank):

- (1, 0) dipole polarizability, interacts with **F**
- (2, 0) dipole-dipole hyperpolarizability, interacts with  ${f F} \otimes {f F}$
- (0, 1) quadrupole polarizability, interacts with  $\nabla \otimes \mathbf{F}$

The distributed sites are assumed to be atomic sites or molecular orbital centroids (depending on the polarization factory used). For the electric field, the composite Cartesian index is just an ordinary Cartesian index. For the electric field gradient and electric field squared, the composite Cartesian index is given as

$$I(x, y) = 3x + y$$

where the values of 0, 1 and 2 correspond to x, y and z Cartesian components, respectively. Therefore, in the latter case, there is 9 distinct composite Cartesian components.

```
17.29.2.8 susceptibility() [2/3]
```

#### **Parameters**

fieldRank	- power dependency with respect to the electric field
fieldGradientRank	- power dependency with respect to the electric field gradient
i	- id of the distributed site

The following susceptibilities are supported (fieldRank, fieldGradientRank):

- (1, 0) dipole polarizability, interacts with **F**
- (2, 0) dipole-dipole hyperpolarizability, interacts with  $\mathbf{F} \otimes \mathbf{F}$
- (0, 1) quadrupole polarizability, interacts with  $\nabla \otimes F$

The distributed sites are assumed to be atomic sites or molecular orbital centroids (depending on the polarization factory used).

# 17.29.2.9 susceptibility() [3/3]

```
std::vector<std::shared_ptr<psi::Matrix> > oepdev::GenEffPar::susceptibile
(
    int fieldRank,
    int fieldGradientRank) const [inline]
```

#### **Parameters**

fieldRank	- power dependency with respect to the electric field
fieldGradientRank	- power dependency with respect to the electric field gradient

The following susceptibilities are supported (fieldRank, fieldGradientRank):

- (1, 0) dipole polarizability, interacts with F
- (2, 0) dipole-dipole hyperpolarizability, interacts with  $F \otimes F$
- (0, 1) quadrupole polarizability, interacts with  $\nabla \otimes \mathbf{F}$

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

- oepdev/libgefp/gefp.h
- · oepdev/libgefp/gefp.cc

# 17.30 oepdev::GenEffParFactory Class Reference

Generalized Effective Fragment Factory. Abstract Base.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::GenEffParFactory:



# **Public Member Functions**

- GenEffParFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
   Construct from wavefunction and Psi4 options.
- virtual  $\sim$ GenEffParFactory ()

Destruct.

- virtual std::shared\_ptr< GenEffPar > compute (void)=0
   Compute the fragment parameters.
- virtual std::shared\_ptr< psi::Wavefunction > wfn (void) const

Grab wavefunction.

virtual psi::Options & options (void) const

Grab options.

std::shared\_ptr< CPHF > cphf\_solver () const

Grab the CPHF object.

# **Static Public Member Functions**

static std::shared\_ptr< GenEffParFactory > build (const std::string &type, std::shared\_ptr< psi::Wavefunction > wfn, psi::Options &opt)

Build Density Matrix Susceptibility Generalized Factory.

#### **Protected Member Functions**

virtual double random\_double ()

Draw random number.

virtual std::shared\_ptr< psi::Vector > draw\_random\_point ()

Draw random point in 3D space, excluding the vdW region.

virtual bool is\_in\_vdWsphere (double x, double y, double z) const

Is the point inside a vdW region?

# **Protected Attributes**

std::shared\_ptr< psi::Wavefunction > wfn\_

Wavefunction.

• psi::Options & options\_

Psi4 Options.

std::default\_random\_engine randomNumberGenerator\_

Random number generators.

- std::uniform\_real\_distribution< double > randomDistribution\_
- std::shared\_ptr< psi::Matrix > excludeSpheres\_

Matrix with vdW sphere information.

std::map< std::string, double > vdwRadius\_

Map with vdW radii.

double cx\_

Centre-of-mass coordinates.

- double cy\_
- double cz\_
- double radius\_

Radius of padding sphere around the molecule.

const int nbf\_

Number of basis functions.

std::shared\_ptr< CPHF > cphfSolver\_

```
The CPHF object.
```

std::shared\_ptr< oepdev::GenEffParFactory > abInitioPolarizationSusceptibilitiesFactory\_

Ab initio polarization susceptibility factory.

# 17.30.1 Detailed Description

Describes the GEFP fragment that is in principle designed to work at correlated levels of theory.

# 17.30.2 Member Function Documentation

# 17.30.2.1 build()

#### **Parameters**

type	- Type of factory
wfn	- Psi4 wavefunction
opt	- Psi4 options

# Available factory types:

• POLARIZATION - creates the polarization generalized effective fragment parameters' factory Factory subtype is specified in Psi4 options (input file).

#### Note

# Useful options:

- POLARIZATION factory type:
  - DMATPOL\_TRAINING\_MODE training mode. Default: EFIELD
  - DMATPOL\_NSAMPLES number of random samples (field or test charges sets).
     Default: 30
  - DMATPOL\_FIELD\_SCALE electric field scale factor (relevant if training mode is EFIELD). Default: 0.01 [au]

- DMATPOL\_NTEST\_CHARGE number of test charges per sample (relevant if training mode is CHARGES). Default: 1
- DMATPOL\_TEST\_CHARGE test charge value (relevant if training mode is CHARGES). Default: 0.001 [au]
- DMATPOL\_FIELD\_RANK electric field rank. Default: 1
- DMATPOL\_GRADIENT\_RANK electric field gradient rank. Default: 0
- DMATPOL\_TEST\_FIELD\_X test electric field in X direction. Default: 0.000
   [au]
- DMATPOL\_TEST\_FIELD\_Y test electric field in Y direction. Default: 0.000
   [au]
- DMATPOL\_TEST\_FIELD\_Z test electric field in Z direction. Default: 0.008
   [au]
- DMATPOL\_OUT\_STATS output file name for statistical evaluation results. Default: dmatpol.stats.dat
- DMATPOL\_DO\_AB\_INITIO compute ab initio susceptibilities and evaluate statistics for it. Default: false
- DMATPOL\_OUT\_STATS\_AB\_INITIO output file name for statistical evaluation results of ab initio model. Default: dmatpol.stats.abinitio.dat

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

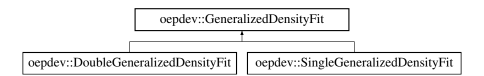
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp.cc

# 17.31 oepdev::GeneralizedDensityFit Class Reference

Generalized Density Fitting Scheme. Abstract Base.

#include <oep\_gdf.h>

Inheritance diagram for oepdev::GeneralizedDensityFit:



# **Public Member Functions**

GeneralizedDensityFit ()

Constructor. Initializes the pointers.

virtual ∼GeneralizedDensityFit ()

Destructor.

virtual std::shared\_ptr< psi::Matrix > compute (void)=0

Perform the generalized density fit.

std::shared\_ptr< psi::Matrix > G (void) const

Extract the  $G_{\xi_i}$  coefficients.

#### Static Public Member Functions

static std::shared\_ptr< GeneralizedDensityFit > build (std::shared\_ptr< psi::BasisSet > bs\_auxiliary, std::shared\_ptr< psi::Matrix > v\_vector)

Factory for Single GDF Computer.

static std::shared\_ptr< GeneralizedDensityFit > build (std::shared\_ptr< psi::BasisSet > bs\_auxiliary, std::shared\_ptr< psi::BasisSet > bs\_intermediate, std::shared\_ptr< psi::Matrix > v\_vector)

Factory for Double GDF Computer.

#### **Protected Member Functions**

void invert\_matrix (std::shared\_ptr< psi::Matrix > &M)

Invert a square matrix and check if the inverse is acceptable.

# **Protected Attributes**

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

The OEP coefficients  $G_{\xi_i}$ .

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

The intermediate DF coefficients for  $\hat{v}|i\rangle$ .

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

The V matrix  $(\xi | \hat{v}i)$ .

int n\_a\_

Number of auxiliary basis set functions.

int n\_i\_

Number of intermediate basis set functions.

• int n\_o\_

Number of OEP's.

std::shared\_ptr< psi::BasisSet > bs\_a\_

Basis set: auxiliary.

std::shared\_ptr< psi::BasisSet > bs\_i\_

Basis set: intermediate.

std::shared\_ptr< oepdev::IntegralFactory > ints\_aa\_

Integral factory: aux - aux.

std::shared\_ptr< oepdev::IntegralFactory > ints\_ai\_

Integral factory: aux - int.

std::shared\_ptr< oepdev::IntegralFactory > ints\_ii\_
 Integral factory: int - int.

# 17.31.1 Detailed Description

Performs the following map:

$$\hat{v}|i)\cong\sum_{m{\eta}}G_{m{\eta}i}|m{\eta})$$

where  $\hat{v}$  is the effective one-electron potential (OEP) operator,  $|i\rangle$  is an arbitrary state vector and  $|\eta\rangle$  is an auxiliary basis vector. The coefficients  $G_{\eta i}$  are stored and define the OEP acting on the state i. The mapping onto the auxiliary space can be done in two ways:

- Single Density Fit. This method requires the auxiliary basis set to be nearly complete.
- Double Density Fit. This method can be used to arbitrary auxiliary basis sets.

# 17.31.2 Member Function Documentation

```
17.31.2.1 build() [1/2]
```

```
std::shared_ptr< GeneralizedDensityFit > GeneralizedDensityFit::build (
    std::shared_ptr< psi::BasisSet > bs_auxiliary,
    std::shared_ptr< psi::Matrix > v_vector ) [static]
```

#### **Parameters**

bs_auxiliary	- auxiliary basis set
v_vector	- the matrix with $V_{\xi i}$ elements

#### Returns

Generalized Density Fit Computer.

# **17.31.2.2** build() [2/2]

```
std::shared_ptr< GeneralizedDensityFit > GeneralizedDensityFit::build (
    std::shared_ptr< psi::BasisSet > bs_auxiliary,
    std::shared_ptr< psi::BasisSet > bs_intermediate,
    std::shared_ptr< psi::Matrix > v_vector ) [static]
```

# **Parameters**

148

bs_auxiliary	- auxiliary basis set
bs₋intermediate	- intermediate basis set
v_vector	- the matrix with $V_{arepsilon i}$ elements

# Returns

Generalized Density Fit Computer.

# 17.31.2.3 compute()

#### Returns

The OEP coefficients  $G_{\xi_i}$ 

Implemented in oepdev::DoubleGeneralizedDensityFit, and oepdev::SingleGeneralizedDensityFit.

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

- oepdev/liboep/oep\_gdf.h
- oepdev/liboep/oep\_gdf.cc

# 17.32 oepdev::GeneralizedPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::GeneralizedPolarGEFactory:



# **Classes**

struct StatisticalSet

A structure to handle statistical data.

# **Public Member Functions**

GeneralizedPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)

Construct from Psi4 wavefunction and options.

virtual ∼GeneralizedPolarGEFactory ()

Destruct.

virtual std::shared\_ptr< GenEffPar > compute (void)

Pefrorm Least-Squares Fit.

bool has\_dipole\_polarizability () const

Dipole Polarizability (interacting with **F**)

bool has\_dipole\_dipole\_hyperpolarizability () const

Dipole-Dipole Hyperpolarizability (interacting with  $\mathbf{F}^2$ )

• bool has\_quadrupole\_polarizability () const

Quadrupole Polarizability (interacting with  $\nabla \otimes \mathbf{F}$ )

• bool has\_ab\_initio\_dipole\_polarizability () const

Ab Initio Dipole Polarizability (interacting with F)

• double Zinit () const

Grab initial summaric Z value.

• double Z () const

Grab final summaric Z value.

# **Protected Member Functions**

void allocate (void)

Allocate memory.

void invert\_hessian (void)

Invert Hessian (do also the identity test)

void compute\_electric\_field\_sums (void)

Compute electric field sum set.

void compute\_electric\_field\_gradient\_sums (void)

Compute electric field gradient sum set.

void compute\_statistics (void)

Run the statistical evaluation of results.

void set\_distributed\_centres (void)

Set the distributed centres.

void compute\_parameters (void)

Compute the parameters.

void fit (void)

Perform least-squares fit.

void compute\_ab\_initio (void)

Compute ab initio parameters.

void save (int i, int j)

Save susceptibility tensors associated with the i-th and j-th basis set function.

virtual void compute\_samples (void)=0

Compute samples of density matrices and select electric field distributions.

virtual void compute\_gradient (int i, int j)=0

Compute Gradient vector associated with the i-th and j-th basis set function.

virtual void compute\_hessian (void)=0

Compute Hessian matrix (independent on the parameters)

# **Protected Attributes**

int nBlocks\_

Number of parameter blocks.

int nSites\_

Number of distributed sites.

int nSitesAbInitio\_

Number of distributed sites of Ab Initio model (FF - single site (com); distributed: LMO sites)

int nParameters\_

Dimensionality of entire parameter space.

std::vector< int > nParametersBlock\_

Dimensionality of parameter space per block.

const int nSamples\_

Number of statistical samples.

const double symmetryNumber\_ [6]

Symmetry number for matrix susceptibilities.

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

Gradient.

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

Hessian.

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

Parameters.

std::shared\_ptr< oepdev::GenEffPar > PolarizationSusceptibilities\_

Density Matrix Susceptibility Tensors Object.

std::shared\_ptr< oepdev::GenEffPar > abInitioPolarizationSusceptibilities\_

Density Matrix Susceptibility Tensors Object for Ab Initio Model.

bool hasDipolePolarizability\_

Has Dipole Polarizability?

bool hasDipoleDipoleHyperpolarizability\_

Has Dipole-Dipole Hyperpolarizability?

bool hasQuadrupolePolarizability\_

Has Quadrupole Polarizability?

bool hasAbInitioDipolePolarizability\_

Has Ab Initio Dipole Polarizability?

StatisticalSet referenceStatisticalSet\_

Reference statistical data.

StatisticalSet referenceDpolStatisticalSet\_

Multipole reference statistical data.

StatisticalSet modelStatisticalSet\_

Model statistical data.

StatisticalSet abInitioModelStatisticalSet\_

Ab Initio Model statistical data.

std::vector< std::shared\_ptr< psi::Matrix >> VMatrixSet\_

Potential matrix set.

std::vector < std::vector < std::shared\_ptr < Vector > > electricFieldSet\_

Electric field set.

std::vector< std::shared\_ptr< Matrix >> > electricFieldGradientSet\_

Electric field gradient set.

std::vector< std::vector< double >> electricFieldSumSet\_

Electric field sum set.

 std::vector< std::vector< std::shared\_ptr< psi::Vector > > electricFieldGradientSum-Set

Electric field gradient sum set.

std::vector < std::shared\_ptr < Vector > > abInitioModelElectricFieldSet\_

Electric field set for Ab Initio Model (LMO-distributed)

const double mField\_

Level shifters for Hessian blocks.

double Zinit\_

Initial summaric Z value.

double Z\_

Final summaric Z value.

std::shared\_ptr< psi::JK > jk\_

Computer of generalized JK objects.

# **Additional Inherited Members**

# 17.32.1 Detailed Description

Implements a general class of methods for the density matrix susceptibility tensors represented by:

$$\delta D_{\alpha\beta} = \sum_{i} \left\{ \mathbf{B}_{i;\alpha\beta}^{(10)} \cdot \mathbf{F}(\mathbf{r}_{i}) + \mathbf{B}_{i;\alpha\beta}^{(20)} : \mathbf{F}(\mathbf{r}_{i}) \otimes \mathbf{F}(\mathbf{r}_{i}) + \mathbf{B}_{i;\alpha\beta}^{(01)} : \nabla_{i} \otimes \mathbf{F}(\mathbf{r}_{i}) + \ldots \right\}$$

where:

•  $\mathbf{B}_{i:\alpha\beta}^{(10)}$  is the density matrix dipole polarizability

- $\mathbf{B}^{(20)}_{i:lphaeta}$  is the density matrix dipole-dipole hyperpolarizability
- ${\bf B}^{(01)}_{i:\alpha\beta}$  is the density matrix quadrupole polarizability

all defined for the generalized distributed site at  $\mathbf{r}_i$ .

Available models:

- 1. Training against uniform electric fields
  - oepdev::LinearUniformEFieldPolarGEFactory linear with respect to electric field
  - oepdev::QuadraticUniformEFieldPolarGEFactory quadratic with respect to electric field
- 2. Training against non-uniform electric fields
  - oepdev::LinearNonUniformEFieldPolarGEFactory linear with respect to electric field, distributed site model
  - oepdev::QuadraticNonUniformEFieldPolarGEFactory quadratic with respect to electric field, distributed site model
  - oepdev::LinearGradientNonUniformEFieldPolarGEFactory linear with respect to electric field and linear with respect to electric field gradient, distributed site model. This model does not function now.
  - oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory linear with respect to electric field and linear with respect to electric field gradient, distributed site model. This model does not function now.

For the non-linear field training, a set of point charges in each training sample is assumed. Distributed models use atomic centers as expansion points.

Determination of the generalized susceptibilities

Let  $\left\{\mathbf{F}^{(1)}(\mathbf{r}), \mathbf{F}^{(2)}(\mathbf{r}), \dots, \mathbf{F}^{(N)}(\mathbf{r}), \dots\right\}$  be a set of  $N_{\text{max}}$  distinct and randomly sampled spatial distributions of electric field. It is assumed that the exact difference one-particle density matrices (with respect to the unperturbed state) defined as

$$\delta \overline{\mathbf{D}}^{(N)} \equiv \overline{\mathbf{D}}^{(N)} - \overline{\mathbf{D}}^{(0)}$$

are known for each sample (overline symbolizes the exact estimate). Now, for each pair of the AO indices the following parameterization is constructed:

$$\delta D^{(N)} = \sum_{i}^{M} \left\{ \sum_{u}^{x,y,z} s_{iu}^{[1]} F_{iu}^{(N)} + \sum_{u}^{x,y,z} \sum_{w < u} r_{uw} s_{iuw}^{[2]} F_{iu}^{(N)} F_{iw}^{(N)} + \dots \right\}$$

(the Greek subscripts were omitted here for notational simplicity). In the above equation,  $B_u^{(i;1)} = s_{iu}^{[1]}$  and  $B_{uw}^{(i;2)} = r_{uw}s_{iuw}^{[2]}$ , where  $r_{uw}$  is the symmetry factor equal to 1 for diagonal elements and 2 for off-diagonal elements of  $B_{uw}^{(i;2)}$ . The multiple parameter blocks ( $\mathbf{s}^{[1]}$ ,  $\mathbf{s}^{[2]}$  and so on) appear in the first power, allowing for linear least-squares regression. The square bracket superscripts denote the block of the parameter space.

To determine the optimum set,  $\mathbf{s} = \begin{pmatrix} \mathbf{s}^{[1]} & \mathbf{s}^{[2]} & \cdots \end{pmatrix}^T$ , a loss function Z that is subject to the least-squares minimization, is defined as

$$Z(\mathbf{s}) = \sum_{N}^{N_{\text{max}}} \left( \delta D^{(N)} - \delta \overline{D}^{(N)} \right)^2 .$$

The Hessian of Z computed with respect to the parameters is parameter-independent (constant) and generally non-singular as long as the electric fields on all distributed sites are different. Therefore, the exact solution for the optimal parameters is given by the Newton equation

$$\mathbf{s} = -\mathbf{H}^{-1} \cdot \mathbf{g} \, .$$

where  $\mathbf{g}$  and  $\mathbf{H}$  are the gradient vector and the Hessian matrix, respectively. Note that in this case the dimensions of parameter space for the block 1 and 2 are equal to 3M and 6M, respectively. The explicit forms of the gradient and Hessian up to second-order are given in the next section.

**Explicit Formulae for Gradient and Hessian Blocks in Linear Regression DMS Model** 

The gradient vector  $\mathbf{g}$  and the Hessian matrix  $\mathbf{H}$  are built from blocks associated with a particular type of parameters, i.e.,

$$\mathbf{g} = \begin{pmatrix} \mathbf{g}^{[1]} \\ \mathbf{g}^{[2]} \end{pmatrix}, \quad \mathbf{H} = \begin{pmatrix} \mathbf{H}^{[11]} & \mathbf{H}^{[12]} \\ \mathbf{H}^{[21]} & \mathbf{H}^{[22]} \end{pmatrix}$$

where the block indices 1 and 2 correspond to the first- and second-order susceptibilities, respectively. Note that the second derivatives of  $\delta D^{(N)}$  with respect to the adjustable parameters vanish due to the linear functional form of the parameterization formula given in the previous section. Thus, the gradient element of the r-th block and Hessian element of the (rs)-th block read

$$\begin{split} g^{[r]} &\equiv \frac{\partial Z}{\partial s^{[r]}} = -2\sum_{N} \overline{\delta D}^{(N)} \frac{\partial \left[\delta D^{(N)}\right]}{\partial s^{[r]}} \;, \\ H^{[rs]} &\equiv \frac{\partial^{2} Z}{\partial s^{[r]} \partial s^{[s]}} = 2\sum_{N} \frac{\partial \left[\delta D^{(N)}\right]}{\partial s^{[r]}} \frac{\partial \left[\delta D^{(N)}\right]}{\partial s^{[s]}} \;. \end{split}$$

The explicit formulae for the gradient are

$$\begin{split} g_{ku}^{[1]} &= -2\sum_{N} \overline{\delta D}^{(N)} F_{ku}^{(N)} , \\ g_{kuw}^{[2]} &= -2r_{uw} \sum_{N} \overline{\delta D}^{(N)} F_{ku}^{(N)} F_{kw}^{(N)} . \end{split}$$

The Hessian subsequently follows to be %

$$\begin{split} H_{ku,lw}^{[11]} &= 2\sum_{N} F_{ku}^{(N)} F_{lw}^{(N)} \;, \\ H_{ku,lu'w'}^{[12]} &= 2r_{u'w'} \sum_{N} F_{ku}^{(N)} F_{lu'}^{(N)} F_{lw'}^{(N)} \;, \\ H_{kuw,lu'w'}^{[22]} &= 2r_{uw} r_{u'w'} \sum_{N} F_{ku}^{(N)} F_{kw}^{(N)} F_{lu'}^{(N)} F_{lw'}^{(N)} \;. \end{split}$$

Note that due to the symmetry of the Hessian matrix, the block 21 is a transpose of the block 12. The composite indices ku and kuw are constructed from the distributed site index k and the appropriate symmetry-adapted (w < u) Cartesian component of a particular DMS tensor: u for the first-order, and uw for the second-order susceptibility tensor, respectively. The method described above can be easily extended to third and higher orders.

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

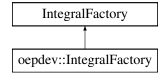
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_base.cc

# 17.33 oepdev::IntegralFactory Class Reference

Extended IntegralFactory for computing integrals.

#include <integral.h>

Inheritance diagram for oepdev::IntegralFactory:



# **Public Member Functions**

- IntegralFactory (std::shared\_ptr< psi::BasisSet > bs1, std::shared\_ptr< psi::BasisSet > bs2, std::shared\_ptr< psi::BasisSet > bs3, std::shared\_ptr< psi::BasisSet > bs4)
   Initialize integral factory given a BasisSet for each center. Becomes (bs1 bs2 | bs3 bs4).
- IntegralFactory (std::shared\_ptr< psi::BasisSet > bs1, std::shared\_ptr< psi::BasisSet > bs2)

Initialize integral factory given a BasisSet for two centres. Becomes (bs1 bs2 | bs1 bs2).

- IntegralFactory (std::shared\_ptr< psi::BasisSet > bs1)
   Initialize integral factory given a BasisSet for two centres. Becomes (bs1 bs1 | bs1 bs1).
- virtual ~IntegralFactory ()
   Destructor.
- virtual oepdev::TwoBodyAOInt \* eri\_1\_1 (int deriv=0, bool use\_shell\_pairs=false)

Returns an ERI\_1\_1 integral object.

- virtual oepdev::TwoBodyAOInt \* eri\_2\_1 (int deriv=0, bool use\_shell\_pairs=false)
   Returns an ERI\_2\_1 integral object.
- virtual oepdev::TwoBodyAOInt \* eri\_2\_2 (int deriv=0, bool use\_shell\_pairs=false)
   Returns an ERI\_2\_2 integral object.
- virtual oepdev::TwoBodyAOInt \* eri\_3\_1 (int deriv=0, bool use\_shell\_pairs=false)
   Returns an ERI\_3\_1 integral object.

# 17.33.1 Detailed Description

In addition to integrals available in Psi4, oepdev::IntegralFactory enables to compute also:

- · OEI's:
  - none at that moment
- ERI's:
  - integrals of type (a|b) oepdev::ERI\_1\_1
  - integrals of type (ab|c) oepdev::ERI\_2\_1
  - integrals of type (abc|d) oepdev::ERI\_3\_1
  - integrals of type (ab|cd) oepdev::ERI\_2\_2 (also in Psi4 as psi::ERI)

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

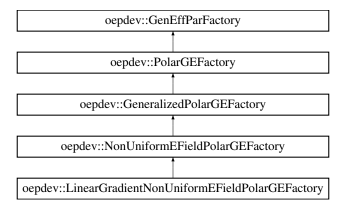
- · oepdev/libpsi/integral.h
- · oepdev/libpsi/integral.cc

# 17.34 oepdev::LinearGradientNonUniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::LinearGradientNonUniformEFieldPolarGEFactory:



# **Public Member Functions**

LinearGradientNonUniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)

void compute\_gradient (int i, int j)

Compute Gradient vector associated with the i-th and j-th basis set function.

void compute\_hessian (void)

Compute Hessian matrix (independent on the parameters)

# **Additional Inherited Members**

# 17.34.1 Detailed Description

Implements the generalized density matrix susceptibility model of the form

$$\delta D_{lphaeta} pprox \sum_{i} \left\{ \mathbf{B}_{i;lphaeta}^{(10)} \cdot \mathbf{F}(\mathbf{r}_{i}) + \mathbf{B}_{i;lphaeta}^{(01)} : \nabla_{i} \otimes \mathbf{F}(\mathbf{r}_{i}) 
ight\}$$

where:

- $\mathbf{B}_{i;lphaeta}^{(10)}$  is the density matrix dipole polarizability
- $\mathbf{B}_{i:\alpha\beta}^{(01)}$  is the density matrix quadrupole polarizability all defined for the distributed site at  $\mathbf{r}_i$ .

Note

This model is not available now and probably will be deprecated in the future.

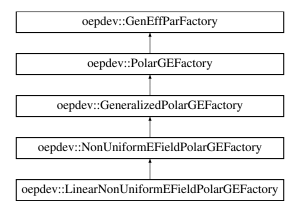
The documentation for this class was generated from the following files:

- · oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_nonuniform\_field\_1\_grad\_1.cc

# 17.35 oepdev::LinearNonUniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

Inheritance diagram for oepdev::LinearNonUniformEFieldPolarGEFactory:



# **Public Member Functions**

- LinearNonUniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options &opt)
- void compute\_gradient (int i, int j)

Compute Gradient vector associated with the i-th and j-th basis set function.

void compute\_hessian (void)

Compute Hessian matrix (independent on the parameters)

# **Additional Inherited Members**

# 17.35.1 Detailed Description

Implements the generalized density matrix susceptibility model of the form

$$\delta D_{lphaeta}pprox\sum_{i}\mathbf{B}_{i;lphaeta}^{(10)}\cdot\mathbf{F}(\mathbf{r}_{i})$$

where:

•  $\mathbf{B}_{i:\alpha\beta}^{(10)}$  is the density matrix dipole polarizability defined for the distributed site at  $\mathbf{r}_i$ .

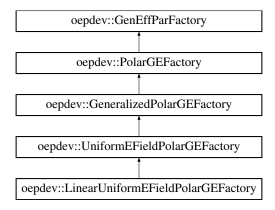
The documentation for this class was generated from the following files:

- · oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_nonuniform\_field\_1.cc

# 17.36 oepdev::LinearUniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

Inheritance diagram for oepdev::LinearUniformEFieldPolarGEFactory:



# **Public Member Functions**

- LinearUniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
- void compute\_gradient (int i, int j)

Compute Gradient vector associated with the i-th and j-th basis set function.

void compute\_hessian (void)

Compute Hessian matrix (independent on the parameters)

# **Additional Inherited Members**

# 17.36.1 Detailed Description

Implements the generalized density matrix susceptibility model of the form

$$\delta D_{lphaeta} pprox \mathbf{B}_{lphaeta}^{(10)} \cdot \mathbf{F}$$

where:

•  ${f B}^{(10)}_{lphaeta}$  is the density matrix dipole polarizability

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

- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_uniform\_field\_1.cc

# 17.37 oepdev::MultipoleConvergence Class Reference

Multipole Convergence.

#include <dmtp.h>

# **Public Types**

```
enum ConvergenceLevel {
R1, R2, R3, R4,
R5 }
enum Property { Energy, Potential }
```

# **Public Member Functions**

MultipoleConvergence (std::shared\_ptr< DMTPole > dmtp1, std::shared\_ptr< DMTPole > dmtp2, ConvergenceLevel max\_clevel=R5)

Construct from two shared DMTPole objects.

virtual ~MultipoleConvergence ()

Destructor.

- void compute (Property property=Energy)
- std::shared\_ptr< psi::Vector > level (ConvergenceLevel clevel=R5)

# **Protected Member Functions**

void compute\_energy ()
 Compute the generalized energy.

void compute\_potential ()

Void compute the generalized potential.

# **Protected Attributes**

ConvergenceLevel max\_clevel\_

Maximum allowed convergence level.

std::shared\_ptr< DMTPole > dmtp\_1\_

First DMTP set.

std::shared\_ptr< DMTPole > dmtp\_2\_

Second DMTP set.

std::map< std::string, std::shared\_ptr< psi::Vector >> convergenceList\_

Dictionary of available convergence level results.

# 17.37.1 Detailed Description

Handles the convergence of the distributed multipole expansions up to hexadecapole. Takes shared pointers to existing DMTPole objects and computes the generalized property:

energy

• potential from the DMTP sets. The results are stored in vector of length equal to the number of DMTP's in a set decribed by DMTPole objects given.

# Note

The number of DMTP's in each object has to be the same.

# 17.37.2 Member Enumeration Documentation

# 17.37.2.1 ConvergenceLevel

enum oepdev::MultipoleConvergence::ConvergenceLevel

Convergence level of the multipole expansion:

#### **Parameters**

R1	- qq term
R2	- qd and sum of the above
R3	- qQ, dd and sum of the above
R4	- qO, dQ and sum of the above
R5	- qH, dO, QQ and sum of the above

# 17.37.2.2 Property

enum oepdev::MultipoleConvergence::Property

Property to be evaluated from interacting DMTP's:

# **Parameters**

Energy	- generalized energy
Potential	- generalized potential

# 17.37.3 Constructor & Destructor Documentation

# 17.37.3.1 MultipoleConvergence()

```
oepdev::MultipoleConvergence::MultipoleConvergence (
    std::shared_ptr< DMTPole > dmtp1,
    std::shared_ptr< DMTPole > dmtp2,
    MultipoleConvergence::ConvergenceLevel max_clevel = R5 )
```

#### **Parameters**

dmtp1	- first DMTPole object
dmtp2	- second DMTPole object
max_clevel	- maximul allowed convergence level

# 17.37.4 Member Function Documentation

# 17.37.4.1 compute()

# Compute the generalized property

#### **Parameters**

```
property - generalized Property
```

# 17.37.4.2 level()

Grab the generalized property at specified level of convergence

# **Parameters**

```
clevel - ConvergenceLevel
```

#### **Returns**

vector of results (each element corresponds to each DMTP pair in a set)

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

- oepdev/lib3d/dmtp.h
- oepdev/lib3d/dmtp\_base.cc

# 17.38 oepdev::NonUniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::NonUniformEFieldPolarGEFactory:



# **Public Member Functions**

- NonUniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
- void compute\_samples (void)

Compute samples of density matrices and select electric field distributions.

virtual void compute\_gradient (int i, int j)=0

Compute Gradient vector associated with the i-th and j-th basis set function.

virtual void compute\_hessian (void)=0

Compute Hessian matrix (independent on the parameters)

# **Additional Inherited Members**

# 17.38.1 Detailed Description

Implements a class of density matrix susceptibility models for parameterization in the non-uniform electric field generated by point charges.

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

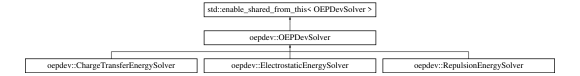
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_nonuniform\_base.cc

# 17.39 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, SharedWave-functionUnion 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.

# 17.39.1 Detailed Description

Uses only a wavefunction union object to initialize. Available solvers:

- ELECTROSTATIC ENERGY
- REPULSION ENERGY

• CHARGE TRANSFER ENERGY

Options controlling the generalized density fitting (GDF):

- OEPDEV\_DF\_TYPE type of the GDF. Default: DOUBLE.
- DF\_BASIS\_OEP auxiliary basis set. Default: sto-3g.
- DF\_BASIS\_INT intermediate basis set. Relevant only if double GDF is used. Default: aug-cc-pVDZ-jkfit. Note that intermediate basis set should be nearly complete.

# 17.39.2 Constructor & Destructor Documentation

# 17.39.2.1 OEPDevSolver()

#### **Parameters**

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

# 17.39.3 Member Function Documentation

# 17.39.3.1 build()

# **Parameters**

target	- target property
wfn₋union	- wavefunction union of isolated molecular wavefunctions

# Implemented target properties:

- ELECTROSTATIC\_ENERGY Coulombic interaction energy between unperturbed wavefunctions
- REPULSION\_ENERGY Pauli repulsion interaction energy between unperturbed wavefunctions.

#### See also

ElectrostaticEnergySolver

#### 17.39.3.2 compute\_benchmark()

Each solver object has one DEFAULT benchmark method

#### **Parameters**

```
method - benchmark method
```

Implemented in oepdev::ChargeTransferEnergySolver, oepdev::RepulsionEnergySolver, and oepdev::ElectrostaticEnergySolver.

#### 17.39.3.3 compute\_oep\_based()

Each solver object has one DEFAULT OEP-based method.

#### **Parameters**

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

Implemented in oepdev::ChargeTransferEnergySolver, oepdev::RepulsionEnergySolver, and oepdev::ElectrostaticEnergySolver.

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

- oepdev/libsolver/solver.h
- oepdev/libsolver/solver\_base.cc
- oepdev/libsolver/solver\_energy\_pauli.cc

# 17.40 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)

Fully ESP-based OEP object.

OEPotential (SharedWavefunction wfn, SharedBasisSet auxiliary, SharedBasisSet intermediate, Options & Options)

General OEP object.

virtual ∼OEPotential ()

Destructor.

virtual void compute (void)

Compute matrix forms of all OEP's within all OEP types.

virtual void compute (const std::string &oepType)=0

Compute matrix forms of all OEP's within a specified OEP type.

virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, std::shared\_ptr< psi::Vector > &v)=0

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

 std::shared\_ptr< OEPotential3D< OEPotential > > make\_oeps3d (const std::string &oep-Type)

Create 3D vector field with OEP.

virtual void write\_cube (const std::string &oepType, const std::string &fileName)

Write potential to a cube file.

• 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.

OEPType oep (const std::string &oepType) const

Retrieve the potentials.

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

Retrieve the potentials in a matrix form.

SharedWavefunction wfn () const

Retrieve wavefunction object.

- void set\_name (const std::string &name)
- virtual void print\_header () const =0

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

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

Build fully ESP-based OEP object.

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

Build general OEP object.

#### **Protected Attributes**

Options options\_

Psi4 options.

SharedWavefunction wfn\_

Wavefunction.

SharedBasisSet primary\_

Promary Basis set.

SharedBasisSet auxiliary\_

Auxiliary Basis set.

• SharedBasisSet intermediate\_

Intermediate Basis set.

std::string name\_

Name of this OEP;.

std::map< std::string, OEPType > oepTypes\_

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

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< oepdev::PotentialInt > potInt\_

One-electron potential shared pointer.

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

Occupied orbitals.

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

Virtual orbitals.

# 17.40.1 Detailed Description

Manages OEP's in matrix and 3D forms. Available OEP categories:

- ELECTROSTATIC ENERGY
- REPULSION ENERGY
- CHARGE TRANSFER ENERGY
- EET COUPLING CONSTANT

## 17.40.2 Constructor & Destructor Documentation

```
17.40.2.1 OEPotential() [1/2]

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

#### **Parameters**

wfn	- wavefunction
options	- Psi4 options

#### 17.40.2.2 **OEPotential()** [2/2]

#### **Parameters**

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

#### 17.40.3 Member Function Documentation

#### **Parameters**

type	- OEP category
wfn	- wavefunction
options	- Psi4 options

# **17.40.3.2** build() [2/2]

#### **Parameters**

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

## 17.40.3.3 make\_oeps3d()

#### **Parameters**

	оерТуре	- type of OEP. ESP-based OEP is assumed.
--	---------	--

#### **Returns**

Vector field 3D with the OEP values.

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

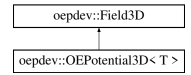
- · oepdev/liboep/oep.h
- oepdev/liboep/oep\_base.cc

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

Class template for OEP 3D fields.

#include <space3d.h>

Inheritance diagram for oepdev::OEPotential3D< T >:



#### **Public Member Functions**

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

Construct random spherical collection of 3D field of type T.

 OEPotential3D (const int &ndim, const int &nx, const int &ny, const int &nz, const double &px, const double &py, const double &pz, std::shared\_ptr< T > oep, const std::string &oepType, psi::Options &options)

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

virtual ∼OEPotential3D ()

Destructor.

virtual std::shared\_ptr< psi::Vector > compute\_xyz (const double &x, const double &y, const double &z)

Compute a value of 3D 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 3D field type stored in instance of  ${\mathbb T}$ 

#### **Additional Inherited Members**

## 17.41.1 Detailed Description

```
template < class T > class oepdev::0EPotential3D < T >
```

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

with the descriptor of a certain 3D field type, x, y, z the points in 3D space in which the scalar or vector 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 oepdev 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)
```

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

oepdev/lib3d/space3d.h

# 17.42 oepdev::OEPType Struct Reference

Container to handle the type of One-Electron Potentials.

```
#include <oep.h>
```

#### **Public Attributes**

std::string name

Name of this type of OEP.

bool is\_density\_fitted

Is this OEP DF-based?

• int n

Number of OEP's within a type.

SharedMatrix matrix

All OEP's of this type gathered in a matrix form.

SharedDMTPole dmtp

Distributed Multipole Object.

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

· oepdev/liboep/oep.h

# 17.43 oepdev::PerturbCharges Struct Reference

Structure to hold perturbing charges.

```
#include <scf_perturb.h>
```

#### **Public Attributes**

• std::vector< double > charges

Vector of charge values.

std::vector < std::shared\_ptr < psi::Vector > > positions

Vector of charge position vectors.

## 17.43.1 Detailed Description

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

• oepdev/libutil/scf\_perturb.h

# 17.44 oepdev::Points3Dlterator::Point Struct Reference

#### **Public Attributes**

- double x
- double y
- double z
- int index

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

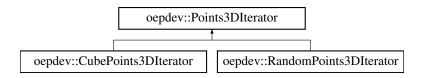
oepdev/lib3d/space3d.h

# 17.45 oepdev::Points3Dlterator Class Reference

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

#include <space3d.h>

Inheritance diagram for oepdev::Points3DIterator:



#### Classes

struct Point

#### **Public Member Functions**

Points3DIterator (const int &np)

Plain constructor. Initializes the abstract features.

virtual ~Points3Dlterator ()

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, 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< Points3Dlterator > 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\_

# 17.45.1 Detailed Description

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.

#### 17.45.2 Constructor & Destructor Documentation

#### 17.45.2.1 Points3Dlterator()

#### **Parameters**

*np* - number of points this iterator is constructed for

#### 17.45.3 Member Function Documentation

# 

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

## **17.45.3.2** build() [2/3]

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

#### **Parameters**

CX	- coordinate x of sphere's centre
су	- coordinate y of sphere's centre
CZ	- coordinate z of sphere's centre

#### **17.45.3.3** build() [3/3]

```
shared_ptr< Points3DIterator > oepdev::Points3DIterator::build (
    const int & np,
    const double & pad,
    psi::SharedMolecule mol ) [static]
```

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:

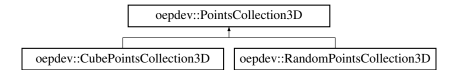
- oepdev/lib3d/space3d.h
- oepdev/lib3d/space3d.cc

# 17.46 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 &cy=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, 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

#### 17.46.1 Detailed Description

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

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

#### 17.46.2 Constructor & Destructor Documentation

## 17.46.2.1 PointsCollection3D()

#### **Parameters**

```
np - number of points to be created
```

## 17.46.3 Member Function Documentation

```
17.46.3.1 build() [1/3]
```

```
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)
```

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
cy	- coordinate y of sphere's centre
CZ	- coordinate z of sphere's centre

#### **17.46.3.2** build() [2/3]

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

#### **17.46.3.3** build() [3/3]

```
std::shared_ptr< PointsCollection3D > oepdev::PointsCollection3D::build (
    const int & nx,
    const int & ny,
    const int & nz,
    const double & px,
    const double & py,
    const double & pz,
    psi::SharedBasisSet bs,
    psi::Options & options ) [static]
```

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
рх	- 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/lib3d/space3d.h
- oepdev/lib3d/space3d.cc

# 17.47 oepdev::PolarGEFactory Class Reference

Polarization GEFP Factory. Abstract Base.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::PolarGEFactory:



#### **Public Member Functions**

180

- PolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options &opt)
   Construct from Psi4 options.
- virtual ~PolarGEFactory ()

Destruct.

virtual std::shared\_ptr< GenEffPar > compute (void)=0
 Compute the density matrix susceptibility tensors.

#### **Protected Member Functions**

- std::shared\_ptr< psi::Vector > draw\_field ()
   Randomly draw electric field value.
- double draw\_charge ()

Randomly draw charge value.

std::shared\_ptr< oepdev::RHFPerturbed > perturbed\_state (const std::shared\_ptr< psi::Vector > &field)

Solve SCF equations to find perturbed state due to uniform electric field.

std::shared\_ptr< oepdev::RHFPerturbed > perturbed\_state (const std::shared\_ptr< psi::Vector > &pos, const double &charge)

Solve SCF equations to find perturbed state due to point charge.

std::shared\_ptr< oepdev::RHFPerturbed > perturbed\_state (const std::shared\_ptr< psi::Matrix > &charges)

Solve SCF equations to find perturbed state due to set of point charges.

std::shared\_ptr< psi::Vector > field\_due\_to\_charges (const std::shared\_ptr< psi::Matrix > &charges, const double &x, const double &y, const double &z)

Evaluate electric field at point (x,y,z) due to point charges.

- std::shared\_ptr< psi::Vector > field\_due\_to\_charges (const std::shared\_ptr< psi::Matrix > &charges, const std::shared\_ptr< psi::Vector > &pos)
- std::shared\_ptr< psi::Matrix > field\_gradient\_due\_to\_charges (const std::shared\_ptr< psi::Matrix > &charges, const double &x, const double &y, const double &z)

Evaluate electric field gradient at point (x,y,z) due to point charges.

std::shared\_ptr< psi::Matrix > field\_gradient\_due\_to\_charges (const std::shared\_ptr< psi::Matrix > &charges, const std::shared\_ptr< psi::Vector > &pos)

#### **Additional Inherited Members**

## 17.47.1 Detailed Description

Basic interface for the polarization density matrix susceptibility parameters.

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

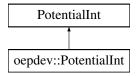
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_base.cc

# 17.48 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 &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.

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.

#### 17.48.1 Constructor & Destructor Documentation

#### **17.48.1.1 PotentialInt()** [1/3]

```
oepdev::PotentialInt::PotentialInt (
    std::vector< psi::SphericalTransform > & st,
    std::shared_ptr< psi::BasisSet > bs1,
    std::shared_ptr< psi::BasisSet > bs2,
    int deriv = 0 )
```

#### **Parameters**

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

#### **17.48.1.2 PotentialInt()** [2/3]

```
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 )
```

#### **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

#### **17.48.1.3 PotentialInt()** [3/3]

```
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)
```

#### **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

## 17.48.2 Member Function Documentation

#### 17.48.2.1 set\_charge\_field()

```
void oepdev::PotentialInt::set_charge_field ( const double & x, const double & y, const double & z, const double & q = 1.0)
```

#### **Parameters**

X	- 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:

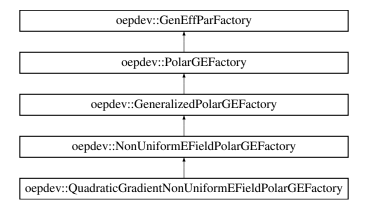
- oepdev/libpsi/potential.h
- · oepdev/libpsi/potential.cc

# 17.49 oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory:



#### **Public Member Functions**

- QuadraticGradientNonUniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
- void compute\_gradient (int i, int j)

Compute Gradient vector associated with the i-th and j-th basis set function.

void compute\_hessian (void)

Compute Hessian matrix (independent on the parameters)

#### **Additional Inherited Members**

## 17.49.1 Detailed Description

Implements the generalized density matrix susceptibility model of the form

$$\delta D_{\alpha\beta} \approx \sum_{i} \left\{ \mathbf{B}_{i;\alpha\beta}^{(10)} \cdot \mathbf{F}(\mathbf{r}_{i}) + \mathbf{B}_{i;\alpha\beta}^{(20)} : \mathbf{F} \otimes \mathbf{F}(\mathbf{r}_{i}) + \mathbf{B}_{i;\alpha\beta}^{(01)} : \nabla_{i} \otimes \mathbf{F}(\mathbf{r}_{i}) \right\}$$

where:

- $\mathbf{B}_{i:\alpha\beta}^{(10)}$  is the density matrix dipole polarizability
- $\mathbf{B}_{i:\alpha\beta}^{(20)}$  is the density matrix dipole-dipole hyperpolarizability
- ${f B}_{i;lphaeta}^{(01)}$  is the density matrix quadrupole polarizability all defined for the distributed site at  ${f r}_i$ .

**Note** 

This model is not available now and probably will be deprecated in the future.

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

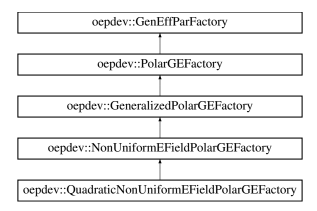
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_nonuniform\_field\_2\_grad\_1.cc

# 17.50 oepdev::QuadraticNonUniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

#include <gefp.h>

Inheritance diagram for oepdev::QuadraticNonUniformEFieldPolarGEFactory:



#### **Public Member Functions**

- QuadraticNonUniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
- void compute\_gradient (int i, int j)

Compute Gradient vector associated with the i-th and j-th basis set function.

void compute\_hessian (void)

Compute Hessian matrix (independent on the parameters)

#### **Additional Inherited Members**

#### 17.50.1 Detailed Description

Implements the generalized density matrix susceptibility model of the form

$$\delta D_{\alpha\beta} \approx \sum_{i} \left\{ \mathbf{B}_{i;\alpha\beta}^{(10)} \cdot \mathbf{F}(\mathbf{r}_{i}) + \mathbf{B}_{i;\alpha\beta}^{(20)} : \mathbf{F}(\mathbf{r}_{i}) \otimes \mathbf{F}(\mathbf{r}_{i}) \right\}$$

where:

- $\mathbf{B}_{i;lphaeta}^{(10)}$  is the density matrix dipole polarizability
- ${\bf B}_{i;\alpha\beta}^{(20)}$  is the density matrix dipole-dipole hyperpolarizability all defined for the distributed site at  ${\bf r}_i$ .

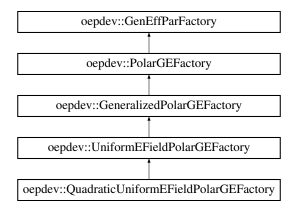
The documentation for this class was generated from the following files:

- · oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_nonuniform\_field\_2.cc

# 17.51 oepdev::QuadraticUniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

Inheritance diagram for oepdev::QuadraticUniformEFieldPolarGEFactory:



#### **Public Member Functions**

- QuadraticUniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options &opt)
- void compute\_gradient (int i, int j)

Compute Gradient vector associated with the i-th and j-th basis set function.

void compute\_hessian (void)

Compute Hessian matrix (independent on the parameters)

## **Additional Inherited Members**

## 17.51.1 Detailed Description

Implements the generalized density matrix susceptibility model of the form

$$\delta D_{\alpha\beta} pprox \mathbf{B}_{\alpha\beta}^{(10)} \cdot \mathbf{F} + \mathbf{B}_{\alpha\beta}^{(20)} : \mathbf{F} \otimes \mathbf{F}$$

where:

- $\mathbf{B}_{lphaeta}^{(10)}$  is the density matrix dipole polarizability
- ${f B}^{(20)}_{lphaeta}$  is the density matrix dipole-dipole hyperpolarizability

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

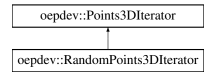
- oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_uniform\_field\_2.cc

# 17.52 oepdev::RandomPoints3DIterator 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 &cx, 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< double > randomDistribution\_

#### **Additional Inherited Members**

# 17.52.1 Detailed Description

**Note:** Always create instances by using static factory method from Points3DIterator. Do not use constructors of this class.

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

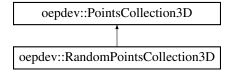
- oepdev/lib3d/space3d.h
- oepdev/lib3d/space3d.cc

# 17.53 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**

# 17.53.1 Detailed Description

**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:

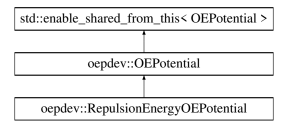
- · oepdev/lib3d/space3d.h
- oepdev/lib3d/space3d.cc

# 17.54 oepdev::RepulsionEnergyOEPotential Class Reference

Generalized One-Electron Potential for Pauli Repulsion Energy.

#include <oep.h>

Inheritance diagram for oepdev::RepulsionEnergyOEPotential:



#### **Public Member Functions**

- RepulsionEnergyOEPotential (SharedWavefunction wfn, SharedBasisSet auxiliary, SharedBasisSet intermediate, Options & options)
- RepulsionEnergyOEPotential (SharedWavefunction wfn, Options & options)
- virtual void compute (const std::string &oepType) override

Compute matrix forms of all OEP's within a specified OEP type.

virtual void compute\_3D (const std::string &oepType, const double &x, const double &y, const double &z, std::shared\_ptr< psi::Vector > &v) override

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

virtual void print\_header () const override

#### **Additional Inherited Members**

#### 17.54.1 Detailed Description

Contains the following OEP types:

- Murrell-etal.S1
- Otto-Ladik.S2

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

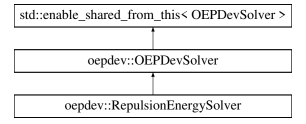
- oepdev/liboep/oep.h
- oepdev/liboep/oep\_energy\_pauli.cc

# 17.55 oepdev::RepulsionEnergySolver Class Reference

Compute the Pauli-Repulsion interaction energy between unperturbed wavefunctions.

#include <solver.h>

Inheritance diagram for oepdev::RepulsionEnergySolver:



## **Public Member Functions**

- **RepulsionEnergySolver** (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**

## 17.55.1 Detailed Description

The implemented methods are shown below

Table 17.74: Methods available in the Solver

Keyword	Method Description	
Benchmark Methods		

Keyword	Method Description	
HAYES_STONE	*Default*. Pauli Repulsion energy at HF level from Hayes and Stone	
	(1984).	
DENSITY_BASED	Pauli Repulsion energy at HF level from Mandado and Hermida-	
	Ramon (2012).	
MURRELL_ETAL	Approximate Pauli Repulsion energy at HF level from Murrell et al	
	(1967).	
OTTO_LADIK	Approximate Pauli Repulsion energy at HF level from Otto and Ladik	
	(1975).	
EFP2	Approximate Pauli Repulsion energy at HF level from EFP2 model.	
OEP-Based Methods		
MURRELL_ETAL_MIX	*Default*. OEP-Murrell et al's: S1 term via DF-OEP, S2 term via	
	ESP-OEP.	
MURRELL_ETAL_ESP	OEP-Murrell et al's: S1 and S2 via ESP-OEP	

#### Note:

- This solver also computes and prints the exchange energy at HF level (formulae are given below) for reference purposes.
- In order to construct this solver, **always** use the OEPDevSolver::build static factory method.

Below the detailed description of the implemented equations is given for each of the above provided methods. In the formulae across, it is assumed that the orbitals are real. The Coulomb notation for electron repulsion integrals (ERI's) is adopted; i.e,

$$(ac|bd) = \iint d\mathbf{r}_1 d\mathbf{r}_2 \phi_a(\mathbf{r}_1) \phi_c(\mathbf{r}_1) \frac{1}{r_{12}} \phi_b(\mathbf{r}_2) \phi_d(\mathbf{r}_2)$$

Greek subscripts denote basis set orbitals whereas Italic subscripts denote the occupied molecular orbitals.

#### **Benchmark Methods**

Pauli Repulsion energy at HF level by Hayes and Stone (1984).

For a closed-shell system, equation of Hayes and Stone (1984) becomes

$$E^{\text{Rep}} = 2\sum_{kl} \left( V_{kl}^A + V_{kl}^B + T_{kl} \right) \left[ [\mathbf{S}^{-1}]_{lk} - \delta_{lk} \right] + \sum_{klmn} (kl|mn) \left\{ 2[\mathbf{S}^{-1}]_{kl} [\mathbf{S}^{-1}]_{mn} - [\mathbf{S}^{-1}]_{kn} [\mathbf{S}^{-1}]_{lm} - 2\delta_{kl} \delta_{mn} + \delta_{kn} \delta_{mn} \right\}$$

where  ${\bf S}$  is the overlap matrix between the doubly-occupied orbitals. The exact, pure exchange energy is for a closed shell case given as

$$E^{\text{Ex,pure}} = -2\sum_{a \in A} \sum_{b \in B} (ab|ba)$$

Similarity transformation of molecular orbitals does not affect the resulting energies. The overall exchange-repulsion interaction energy is then (always net repulsive)

$$E^{\text{Ex-Rep}} = E^{\text{Ex,pure}} + E^{\text{Rep}}$$

Repulsion energy of Mandado and Hermida-Ramon (2011)

At the Hartree-Fock level, the exchange-repulsion energy from the density-based scheme of Mandado and Hermida-Ramon (2011) is fully equivalent to the method by Hayes and Stone (1984). However, density-based method enables to compute exchange-repulsion energy at any level of theory. It is derived based on the Pauli deformation density matrix,

$$\Delta \mathbf{D}^{\text{Pauli}} \equiv \mathbf{D}^{oo} - \mathbf{D}$$

where  $\mathbf{D}^{oo}$  and  $\mathbf{D}$  are the density matrix formed from mutually orthogonal sets of molecular orbitals within the entire aggregate (formed by symmetric orthogonalization of MO's) and the density matrix of the unperturbed system (that can be understood as a Hadamard sum  $\mathbf{D} \equiv \mathbf{D}^A \oplus \mathbf{D}^B$ ).

At HF level, the Pauli deformation density matrix is given by

$$\Delta \mathbf{D}^{Pauli} = \mathbf{C} \left[ \mathbf{S}^{-1} - \mathbf{1} \right] \mathbf{C}^{\dagger}$$

whereas the density matrix constructed from mutually orthogonal orbitals is

$$\mathbf{D}^{oo} = \mathbf{C}\mathbf{S}^{-1}\mathbf{C}^{\dagger}$$

In the above equations, S is the overlap matrix between doubly occupied molecular orbitals of the entire aggregate.

Here, the expressions for the exchange-repulsion energy at any level of theory are shown for the case of open-shell system. The net repulsive energy is given as

$$E^{\text{Ex-Rep}} = E^{\text{Rep},1} + E^{\text{Rep},2} + E^{\text{Ex}}$$

where the one- and two-electron part of the repulsion energy is

$$E^{\text{Rep},1} = E^{\text{Rep},\text{Kin}} + E^{\text{Rep},\text{Nuc}}$$
  
 $E^{\text{Rep},2} = E^{\text{Rep},\text{el}-\Delta} + E^{\text{Rep},\Delta-\Delta}$ 

The kinetic and nuclear contributions are

$$\begin{split} E^{\text{Rep,Kin}} &= 2 \sum_{\alpha\beta \in A,B} \Delta D_{\alpha\beta}^{\text{Pauli}} T_{\alpha\beta} \\ E^{\text{Rep,Nuc}} &= 2 \sum_{\alpha\beta \in A,B} \Delta D_{\alpha\beta}^{\text{Pauli}} \sum_{z \in A,B} V_{\alpha\beta}^{(z)} \end{split}$$

whereas the electron-deformation and deformation-deformation interaction contributions are

$$\begin{split} E^{\text{Rep,el}-\Delta} &= 4 \sum_{\alpha\beta\gamma\delta \in A,B} \Delta D^{\text{Pauli}}_{\alpha\beta} D_{\gamma\delta}(\alpha\beta|\gamma\delta) \\ E^{\text{Rep},\Delta-\Delta} &= 2 \sum_{\alpha\beta\gamma\delta \in A,B} \Delta D^{\text{Pauli}}_{\alpha\beta} \Delta D^{\text{Pauli}}_{\gamma\delta}(\alpha\beta|\gamma\delta) \end{split}$$

The associated exchange energy is given by

$$E^{\mathrm{Ex}} = -\sum_{lphaeta\gamma\delta\in A,B} \left[ D^{oo}_{lpha\delta}D^{oo}_{eta\gamma} - D^{A}_{lpha\delta}D^{A}_{eta\gamma} - D^{B}_{lpha\delta}D^{B}_{eta\gamma} 
ight] (lphaeta|\gamma\delta)$$

It is important to emphasise that, although, at HF level, the particular 'repulsive' and 'exchange' energies computed by using either Hayes and Stone or Mandado and Hermida-Ramon methods are not equal to each other, they sum up to exactly the same exchange-repulsion energy,  $E^{\rm Ex-Rep}$ . Therefore, these methods at HF level are fully equivalent but the nature of partitioning of repulsive and exchange parts is different. It is also noted that the orbital localization does *not* affect the resulting energies, as opposed to the few approximate methods described below (Otto-Ladik and EFP2 methods).

Approximate Pauli Repulsion energy at HF level from Murrell et al.

By expanding the overlap matrix in a Taylor series one can show that the Pauli repulsion energy is approximately given as

$$E^{\text{Rep}} = E^{\text{Rep}}(\mathscr{O}(S)) + E^{\text{Rep}}(\mathscr{O}(S^2))$$

where the first-order term is

$$E^{\operatorname{Rep}}(\mathscr{O}(S)) = -2\sum_{a \in A}\sum_{b \in B}S_{ab}\left\{V_{ab}^A + \sum_{c \in A}\left[2(ab|cc) - (ac|bc)\right] + V_{ab}^B + \sum_{d \in B}\left[2(ab|dd) - (ad|bd)\right]\right\}$$

whereas the second-order term is

$$E^{\text{Rep}}(\mathscr{O}(S^2)) = 2\sum_{a \in A} \sum_{b \in B} S_{ab} \left\{ \sum_{c \in A} S_{bc} \left[ V_{ac}^B + 2\sum_{d \in B} (ac|dd) \right] + \sum_{d \in B} S_{ad} \left[ V_{bd}^A + 2\sum_{c \in A} (bd|cc) \right] - \sum_{c \in A} \sum_{d \in B} S_{cd}(ac|bd) \right\}$$

Thus derived repulsion energy is invariant with respect to transformation of molecular orbitals, similarly as Hayes-Stone's method and density-based method. By using OEP technique, the above theory can be exactly re-cast *without* any further approximations.

Approximate Pauli Repulsion energy at HF level from Otto and Ladik (1975).

The Pauli repulsion energy is approximately given as

$$E^{\mathsf{Rep}} = E^{\mathsf{Rep}}(\mathscr{O}(S)) + E^{\mathsf{Rep}}(\mathscr{O}(S^2))$$

where the first-order term is

$$E^{\text{Rep}}(\mathscr{O}(S)) = -2\sum_{a \in A}\sum_{b \in B}S_{ab}\left\{V_{ab}^A + 2\sum_{c \in A}(ab|cc) - (ab|aa) + V_{ab}^B + 2\sum_{d \in B}(ab|dd) - (ab|bb)\right\}$$

whereas the second-order term is

$$E^{\text{Rep}}(\mathscr{O}(S^2)) = 2\sum_{a \in A} \sum_{b \in B} S_{ab}^2 \left\{ V_{aa}^B + V_{bb}^A + 2\sum_{c \in A} (cc|bb) + 2\sum_{d \in B} (aa|dd) - (aa|bb) \right\}$$

Thus derived repulsion energy is *not* invariant with respect to transformation of molecular orbitals, in contrast to Hayes-Stone's method and density-based method. It was shown that good results are obtained when using localized molecular orbitals, whereas using canonical molecular orbitals brings poor results. By using OEP technique, the above theory can be exactly re-cast *without* any further approximations.

Approximate Pauli Repulsion energy at HF level from Jensen and Gordon (1996).

The Pauli repulsion energy used within the EFP2 approach is approximately given as

$$E^{\text{Rep}} = E^{\text{Rep}}(\mathscr{O}(S)) + E^{\text{Rep}}(\mathscr{O}(S^2))$$

where the first-order term is

$$E^{\text{Rep}}(\mathscr{O}(S)) = -2\sum_{a \in A} \sum_{b \in B} S_{ab} \left\{ \sum_{c \in A} F_{ac}^A S_{cb} + \sum_{d \in B} F_{bd}^B S_{da} - 2T_{ab} \right\}$$

whereas the second-order term is

$$E^{\text{Rep}}(\mathscr{O}(S^2)) = 2\sum_{a \in A} \sum_{b \in B} S_{ab}^2 \left\{ \sum_{x \in A} \frac{-Z_x}{R_{xb}} + \sum_{y \in B} \frac{-Z_y}{R_{ya}} + \sum_{c \in A} \frac{2}{R_{bc}} + \sum_{d \in B} \frac{2}{R_{ad}} - \frac{1}{R_{ab}} \right\}$$

Thus derived repulsion energy is *not* invariant with respect to transformation of molecular orbitals, in contrast to Hayes-Stone's method and density-based method. It was shown that good results are obtained when using localized molecular orbitals, whereas using canonical molecular orbitals brings poor results.

In EFP2, exchange energy is approximated by spherical Gaussian approximation (SGO). The result of this is the following formula for the exchange energy:

$$E^{\rm Ex} \approx -4 \sum_{a \in A} \sum_{b \in R} \sqrt{\frac{-2 \ln |S_{ab}|}{\pi}} \frac{S_{ab}^2}{R_{ab}}$$

In all the above formulas,  $R_{ij}$  are distances between position vectors of \*i\*th and \*j\*th point. The LMO centroids are defined by

$$\mathbf{r}_a = (a|\mathbf{r}|a)$$

where a denotes the occupied molecular orbital.

#### **OEP-Based Methods**

The Murrell et al's theory of Pauli repulsion for S-1 term and the Otto-Ladik's theory for S-2 term is here re-cast by introducing OEP's. The S-1 term is expressed via DF-OEP, whereas the S-2 term via ESP-OEP.

S-1 term (Murrell et al.)

The OEP reduction without any approximations leads to the following formula

$$E^{\mathrm{Rep}}(\mathscr{O}(S^1)) = -2\sum_{a \in A}\sum_{b \in B}S_{ab}\left\{\sum_{\xi \in A}S_{b\xi}G^A_{\xi a} + \sum_{\eta \in B}S_{a\eta}G^B_{\eta b}\right\}$$

where the OEP matrices are given as

$$G_{\xi a}^{A} = \sum_{\xi' \in A} \left[ \mathbf{S}^{-1} \right]_{\xi \xi'} \sum_{\alpha \in A} \left\{ C_{\alpha a} V_{\alpha \xi'}^{A} + \sum_{\mu \nu \in A} \left[ 2C_{\alpha a} D_{\mu \nu} - C_{\nu a} D_{\alpha \mu} \right] (\alpha \xi' | \mu \nu) \right\}$$

and analogously for molecule B. Here, the nuclear attraction integrals are denoted by  $V_{lpha\xi'}^A$ .

S-2 term (Otto-Ladik)

After the OEP reduction, this contribution under Otto-Ladik approximation has the following form:

$$E^{\text{Rep}}(\mathcal{O}(S^2)) = 2\sum_{a \in A} \sum_{b \in B} S_{ab}^2 \left\{ \sum_{x \in A} q_{xa} V_{bb}^{(x)} + \sum_{y \in B} q_{yb} V_{aa}^{(y)} \right\}$$

where the ESP charges associated with each occupied molecular orbital reproduce the *effective* potential of molecule in question, i.e.,

$$\sum_{x \in A} \frac{q_{xa}}{|\mathbf{r} - \mathbf{r}_x|} \cong v_a^A(\mathbf{r})$$

where the potential is given by

$$v_a^A(\mathbf{r}) = \sum_{x \in A} \frac{-Z_x}{|\mathbf{r} - \mathbf{r}_x|} + 2\sum_{c \in A} \int \frac{\phi_c(\mathbf{r}')\phi_c(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}' - \frac{1}{2} \int \frac{\phi_a(\mathbf{r}')\phi_a(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}'$$

#### 17.55.2 Member Function Documentation

#### 17.55.2.1 compute\_benchmark()

Each solver object has one DEFAULT benchmark method

#### **Parameters**

```
method - benchmark method
```

Implements oepdev::OEPDevSolver.

#### 17.55.2.2 compute\_oep\_based()

Each solver object has one DEFAULT OEP-based method.

#### **Parameters**

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

Implements oepdev::OEPDevSolver.

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

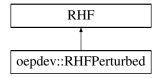
- oepdev/libsolver/solver.h
- oepdev/libsolver/solver\_energy\_pauli.cc

# 17.56 oepdev::RHFPerturbed Class Reference

RHF theory under electrostatic perturbation.

```
#include <scf_perturb.h>
```

Inheritance diagram for oepdev::RHFPerturbed:



#### **Public Member Functions**

RHFPerturbed (std::shared\_ptr< psi::Wavefunction > ref\_wfn, std::shared\_ptr< psi::SuperFunctional > functional)

Build from wavefunction and superfunctional.

RHFPerturbed (std::shared\_ptr< psi::Wavefunction > ref\_wfn, std::shared\_ptr< psi::SuperFunctional > functional, psi::Options &options, std::shared\_ptr< psi::PSIO > psio)

Build from wavefunction and superfunctional + options and psio.

virtual ∼RHFPerturbed ()

Clear memory.

virtual double compute\_energy ()

Compute total energy.

virtual void set\_perturbation (std::shared\_ptr< psi::Vector > field)

Perturb the system with external electric field.

virtual void <u>set\_perturbation</u> (const double &fx, const double &fy, const double &fz)

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

 virtual void set\_perturbation (std::shared\_ptr< psi::Vector > position, const double &charge)

Perturb the system with a point charge.

 virtual void set\_perturbation (const double &rx, const double &ry, const double &rz, const double &charge)

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

- std::shared\_ptr< psi::Matrix > Vpert () const
   Get a copy of the perturbation potential one-electron matrix.
- double nuclear\_interaction\_energy () const
   Get the interaction energy of the nuclei with the perturbing potential.

#### **Protected Member Functions**

virtual void perturb\_Hcore ()
 Add the electrostatic perturbation to the Hcore matrix.

#### **Protected Attributes**

- std::shared\_ptr< psi::Vector > perturbField\_ Perturbing electric field.
- std::shared\_ptr< PerturbCharges > perturbCharges\_ Perturbing charges.
- std::shared\_ptr< psi::Matrix > Vpert\_
   Perturbation potential one-electron matrix.
- double nuclearInteractionEnergy\_
   Electrostatic interaction energy due to nuclei.

## 17.56.1 Detailed Description

Compute RHF wavefunction under the following conditions:

- · external uniform electric field
- set of point charges The mixed conditions can also be used.

## **Theory**

The electrostatic perturbation is here understood as a distribution of external (generally non-uniform) electric field. It is assumed that this perturbation is one-electron in nature. Therefore, the one-electron Hamiltonian is changed according to the following

$$\mathbf{H}^{\text{core}} \to \mathbf{H}^{\text{core}} + \sum_{n} q_{n} \mathbf{V}^{(n)} - \mathbb{M} \cdot \mathbf{F}$$

where  $q_n$  is the external classical point charge,  $\mathbf{V}^{(n)}$  is the associated matrix of potential integrals,  $\mathbb{M}$  is the vector of dipole integrals and  $\mathbf{F}$  is an external uniform electric field. The total energy is then computed by performing an SCF procedure on the above one-electron Hamiltionian. The contribution due to nuclei is included, i.e.,

$$E_{
m Nuc} 
ightarrow E_{
m Nuc-Nuc} + \sum_{In} rac{q_n Z_I}{r_{In}} - \mu_{
m Nuc} \cdot {f F}$$

where  $\mu_{\mathrm{Nuc}}$  is the nuclear dipole moment and  $Z_I$  is the atomic number of the Ith nucleus. It is added in the nuclear repulsion energy  $E_{\mathrm{Nuc-Nuc}}$  (note that the resulting energy can be negative as well depending on the electric field direction and configuration of point charges.

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

- oepdev/libutil/scf\_perturb.h
- oepdev/libutil/scf\_perturb.cc

# 17.57 oepdev::ShellCombinationsIterator Class Reference

Iterator for Shell Combinations. Abstract Base.

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

Inheritance diagram for oepdev::ShellCombinationsIterator:

```
oepdev::ShellCombinationsIterator

oepdev::AllAOShellCombinationsIterator_4
```

#### **Public Member Functions**

• ShellCombinationsIterator (int nshell)

Constructor.

virtual ∼ShellCombinationsIterator ()

Destructor.

virtual void first (void)=0

First iteration.

virtual void next (void)=0

Next iteration.

- virtual std::shared\_ptr< psi::BasisSet > bs\_1 (void) const
  - Grab the basis set of axis 1.
- virtual std::shared\_ptr< psi::BasisSet > bs\_2 (void) const

Grab the basis set of axis 2.

- virtual std::shared\_ptr< psi::BasisSet > bs\_3 (void) const
   Grab the basis set of axis 3.
- virtual std::shared\_ptr< psi::BasisSet > bs\_4 (void) const
   Grab the basis set of axis 4.
- virtual int P (void) const

Grab the current shell P index.

virtual int Q (void) const

Grab the current shell Q index.

virtual int R (void) const

Grab the current shell R index.

virtual int S (void) const

Grab the current shell S index.

virtual bool is\_done (void)

Return status of an iterator.

· virtual const int nshell (void) const

Return number of shells this iterator is for.

- virtual std::shared\_ptr< AOIntegralsIterator > ao\_iterator (std::string mode="ALL") const
- virtual void compute\_shell (std::shared\_ptr< oepdev::TwoBodyAOInt > tei) const =0
- virtual void compute\_shell (std::shared\_ptr< psi ::TwoBodyAOInt > tei) const =0

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

static std::shared\_ptr< ShellCombinationsIterator > build (const IntegralFactory &ints, std::string mode="ALL", int nshell=4)

Build shell iterator from oepdev::IntegralFactory.

static std::shared\_ptr< ShellCombinationsIterator > build (std::shared\_ptr< IntegralFactory > ints, std::string mode="ALL", int nshell=4)

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

static std::shared\_ptr< ShellCombinationsIterator > build (const psi::IntegralFactory &ints, std::string mode="ALL", int nshell=4)

Build shell iterator from psi::IntegralFactory.

static std::shared\_ptr< ShellCombinationsIterator > build (std::shared\_ptr< psi::IntegralFactory > ints, std::string mode="ALL", int nshell=4)

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

## **Protected Attributes**

SharedBasisSet bs\_1\_

Basis set of axis 1.

SharedBasisSet bs\_2\_

Basis set of axis 2.

SharedBasisSet bs\_3\_

Basis set of axis 3.

SharedBasisSet bs\_4\_

Basis set of axis 4.

const int nshell\_

Number of shells this iterator is for.

bool done

Status of an iterator.

# 17.57.1 Detailed Description

Date

2018/03/01 17:22:00

## 17.57.2 Constructor & Destructor Documentation

#### 17.57.2.1 ShellCombinationsIterator()

#### **Parameters**

*nshell* - number of shells this iterator is for

#### 17.57.3 Member Function Documentation

```
17.57.3.1 ao_iterator()
```

Make an AO integral iterator based on current shell

#### **Parameters**

```
mode - either "ALL" or "UNIQUE" (iterate over all or unique integrals)
```

#### **Returns**

iterator over AO integrals

#### **Parameters**

ints	- integral factory
mode	- mode of iteration (either ALL or UNIQUE)
nshell	- number of shells to iterate through

#### **Returns**

shell iterator

# **Examples:**

example\_integrals\_iter.cc.

#### **Parameters**

ints	- integral factory
mode	- mode of iteration (either ALL or UNIQUE)
nshell	- number of shells to iterate through

#### **Returns**

shell iterator

#### 17.57.3.4 compute\_shell()

Compute integrals in a current shell. Works both for oepdev::TwoBodyAOInt and psi::TwoBodyAOInt

#### **Parameters**

```
tei | - two body integral object
```

Implemented in oepdev::AllAOShellCombinationsIterator\_2, and oepdev::AllAOShellCombinationsIterator\_4. The documentation for this class was generated from the following files:

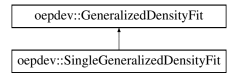
- oepdev/libutil/integrals\_iter.h
- oepdev/libutil/integrals\_iter.cc

# 17.58 oepdev::SingleGeneralizedDensityFit Class Reference

Generalized Density Fitting Scheme - Single Fit.

```
#include <oep_gdf.h>
```

Inheritance diagram for oepdev::SingleGeneralizedDensityFit:



#### **Public Member Functions**

- SingleGeneralizedDensityFit (std::shared\_ptr< psi::BasisSet > bs\_auxiliary, std::shared\_ptr< psi::Matrix > v\_vector)
- std::shared\_ptr< psi::Matrix > compute (void)
   Perform the generalized density fit.

#### **Additional Inherited Members**

# 17.58.1 Detailed Description

The density fitting map projects the OEP onto the auxiliary, nearly complete basis set space through application of the resolution of identity. Refer to density fitting in complete space for more details.

#### 17.58.2 Determination of the OEP matrix

Coefficients G are computed by using the following relation

$$\mathbf{G}^{(i)} = \mathbf{v}^{(i)} \cdot \mathbf{S}^{-1}$$

where

$$S_{\xi\eta} = (\xi|\eta)$$
  
 $v_{\xi}^{(i)} = (\xi|\hat{v}i)$ 

In the above, | denotes the single integration over electron coordinate, i.e.,

$$(a|b) \equiv \int d\mathbf{r} \phi_a^*(\mathbf{r}) \phi_b(\mathbf{r})$$

whereas the spatial form of the potential operator  $\hat{v}$  can be expressed by

$$v(\mathbf{r}) \equiv \int d\mathbf{r}' \frac{\rho(\mathbf{r}')}{|\mathbf{r}' - \mathbf{r}|}$$

with  $\rho(\mathbf{r})$  being the effective one-electron density associated with  $\hat{v}$ .

#### 17.58.3 Member Function Documentation

#### 17.58.3.1 compute()

**Returns** 

The OEP coefficients  $G_{\xi_i}$ 

Implements oepdev::GeneralizedDensityFit.

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

- oepdev/liboep/oep\_qdf.h
- oepdev/liboep/oep\_gdf.cc

# 17.59 oepdev::GeneralizedPolarGEFactory::StatisticalSet Struct Reference

A structure to handle statistical data.

```
#include <gefp.h>
```

# **Public Attributes**

std::vector < double > InducedInteractionEnergySet
 Interaction energy set.

- std::vector < std::shared\_ptr < psi::Matrix > > DensityMatrixSet
   Density matrix set.
- std::vector < std::shared\_ptr < psi::Vector > > InducedDipoleSet
   Induced dipole moment set.
- std::vector < std::shared\_ptr < psi::Vector > > InducedQuadrupoleSet
   Induced quadrupole moment set.
- std::vector < std::shared\_ptr < psi::Matrix > > JKMatrixSet
   Sum of J and K matrix set.

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

oepdev/libgefp/gefp.h

# 17.60 oepdev::test::Test Class Reference

#### Manages test routines.

```
#include <test.h>
```

# **Public Member Functions**

- Test (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options &options)
   Construct the tester.
- ∼Test ()

Destructor.

double run (void)

Pefrorm the test.

### **Protected Member Functions**

double test\_basic (void)

Test the basic functionalities of OEPDev.

double test\_cphf (void)

Test the CPHF method.

double test\_dmatPol (void)

Test the density matrix susceptibility (X = 1)

double test\_dmatPolX (void)

Test the density matrix susceptibility.

double test\_eri\_1\_1 (void)

Test the oepdev::ERI\_1\_1 class against psi::ERI.

double test\_eri\_2\_2 (void)

Test the oepdev::ERI\_2\_2 class against psi::ERI.

double test\_eri\_3\_1 (void)

Test the oepdev::ERI\_3\_1 class against psi::ERI.

double test\_unitaryOptimizer (void)

Test the oepdev::UnitaryOptimizer class.

double test\_unitaryOptimizer\_4\_2 (void)

Test the oepdev::UnitaryOptimizer\_4\_2 class.

double test\_scf\_perturb (void)

Test the oepdev::RHFPerturbed class.

double test\_camm (void)

Test the oepdev::CAMM class.

double test\_dmtp\_energy (void)

Test the oepdev::DMTP class for energy calculations.

double test\_custom (void)

Test the custom code.

#### **Protected Attributes**

std::shared\_ptr< psi::Wavefunction > wfn\_

Wavefunction object.

psi::Options & options\_

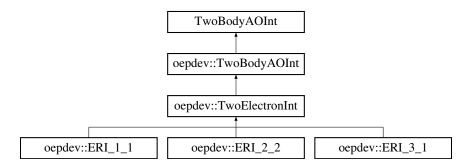
Psi4 Options.

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

- oepdev/libtest/test.h
- · oepdev/libtest/basic.cc
- oepdev/libtest/camm.cc
- · oepdev/libtest/cphf.cc
- oepdev/libtest/dmatpol.cc
- oepdev/libtest/dmatpolX.cc
- oepdev/libtest/dmtp\_energy.cc
- oepdev/libtest/eri\_1\_1.cc
- oepdev/libtest/eri\_2\_2.cc
- oepdev/libtest/eri\_3\_1.cc
- oepdev/libtest/scf\_perturb.cc
- oepdev/libtest/test.cc
- oepdev/libtest/test\_custom.cc
- oepdev/libtest/unitary\_optimizer.cc
- oepdev/libtest/unitary\_optimizer\_4\_2.cc

# 17.61 oepdev::TwoBodyAOInt Class Reference

Inheritance diagram for oepdev::TwoBodyAOInt:



#### **Public Member Functions**

- virtual void compute (std::shared\_ptr< psi::Matrix > &result, int ibs1=0, int ibs2=2)

  Compute two-body two-centre integral and put it into matrix.
- virtual void compute (psi::Matrix &result, int ibs1=0, int ibs2=2)
- virtual size\_t compute\_shell (int, int, int, int)=0
- virtual size\_t compute\_shell (int, int, int)=0
- virtual size\_t compute\_shell (int, int)=0
- virtual size\_t compute\_shell\_deriv1 (int, int, int, int)=0
- virtual size\_t compute\_shell\_deriv2 (int, int, int, int)=0
- virtual size\_t compute\_shell\_deriv1 (int, int, int)=0
- virtual size\_t compute\_shell\_deriv2 (int, int, int)=0
- virtual size\_t compute\_shell\_deriv1 (int, int)=0
- virtual size\_t compute\_shell\_deriv2 (int, int)=0

#### **Protected Member Functions**

- TwoBodyAOInt (const IntegralFactory \*intsfactory, int deriv=0)
- TwoBodyAOInt (const TwoBodyAOInt &rhs)

#### 17.61.1 Member Function Documentation

#### **Parameters**

result	- matrix where to store (i $  $ j) two-body integrals
ibs1	- first basis set axis
ibs2	- second basis set axis

# 17.61.1.2 compute() [2/2]

```
void oepdev::TwoBodyAOInt::compute (
    psi::Matrix & result,
    int ibs1 = 0,
    int ibs2 = 2 ) [virtual]
```

This is an overloaded member function, provided for convenience. It differs from the above function only in what argument(s) it accepts.

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

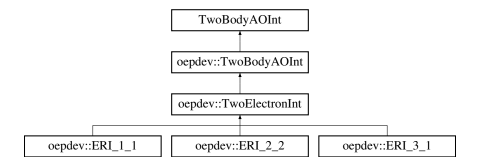
- oepdev/libpsi/integral.h
- · oepdev/libpsi/integral.cc

# 17.62 oepdev::TwoElectronInt Class Reference

General Two Electron Integral.

```
#include <eri.h>
```

Inheritance diagram for oepdev::TwoElectronInt:



#### **Public Member Functions**

- TwoElectronInt (const IntegralFactory \*integral, int deriv, bool use\_shell\_pairs)
- virtual size\_t compute\_shell (int, int)
   Compute ERI's between 2 shells. Result is stored in buffer.
- virtual size\_t compute\_shell (int, int, int)

Compute ERI's between 3 shells. Result is stored in buffer.

virtual size\_t compute\_shell (int, int, int, int)

Compute ERI's between 4 shells. Result is stored in buffer.

- virtual size\_t compute\_shell (const psi::AOShellCombinationsIterator &)
- virtual size\_t compute\_shell\_deriv1 (int, int)

Compute first derivatives of ERI's between 2 shells.

virtual size\_t compute\_shell\_deriv2 (int, int)

Compute second derivatives of ERI's between 2 shells.

virtual size\_t compute\_shell\_deriv1 (int, int, int)

Compute first derivatives of ERI's between 3 shells.

virtual size\_t compute\_shell\_deriv2 (int, int, int)

Compute second derivatives of ERI's between 3 shells.

virtual size\_t compute\_shell\_deriv1 (int, int, int, int)

Compute first derivatives of ERI's between 4 shells.

virtual size\_t compute\_shell\_deriv2 (int, int, int, int)

Compute second derivatives of ERI's between 4 shells.

#### **Protected Member Functions**

• int get\_cart\_am (int am, int n, int x)

Get the angular momentum per Cartesian component.

double get\_R (int N, int L, int M)

Get the (N,L,M)th McMurchie-Davidson coefficient.

virtual size\_t compute\_doublet (int, int)

Computes the ERI's between three shells.

virtual size\_t compute\_triplet (int, int, int)

Computes the ERI's between three shells.

virtual size\_t compute\_quartet (int, int, int, int)

Computes the ERI's between four shells.

#### **Protected Attributes**

const int max\_am\_

Maximum angular momentum.

const int n\_max\_am\_

Maximum number of angular momentum functions.

psi::Fjt \* fjt\_

Computes the fundamental: Boys function value at T for degree v.

bool use\_shell\_pairs\_

Should we use shell pair information?

const double cartMap\_ [60]

Map of Cartesian components per each am.

const double df\_ [8]

Double factorial array.

double \* mdh\_buffer\_R\_

Buffer for the McMurchie-Davidson-Hermite R coefficents.

# 17.62.1 Detailed Description

Implements the McMurchie-Davidson recursive scheme for all integral types. The integral can be defined for any number of Gaussian centres, thus it is not limited to 2-by-2 four-centre ERI. Currently implemented subtypes are:

```
    oepdev::ERI_1_1 - 2-centre electron-repulsion integral (i|j)
```

- oepdev::ERI\_2\_2 4-centre electron-repulsion integral (ij|kl)
- oepdev::ERI\_3\_1 4-centre electron-repulsion integral (ijk|I)

#### See also

The Integral Package Library

### 17.62.2 Member Function Documentation

```
17.62.2.1 compute_shell()
```

Compute ERIs between 4 shells. Result is stored in buffer. Only for use with ERI\_2\_2 and the same basis sets, otherwise shell pairs won't be compatible.

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

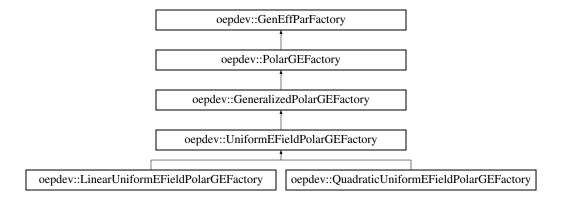
- · oepdev/libints/eri.h
- oepdev/libints/eri.cc

# 17.63 oepdev::UniformEFieldPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Parameterization.

```
#include <gefp.h>
```

Inheritance diagram for oepdev::UniformEFieldPolarGEFactory:



#### **Public Member Functions**

- UniformEFieldPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)
- void compute\_samples (void)

Compute samples of density matrices and select electric field distributions.

virtual void compute\_gradient (int i, int j)=0

Compute Gradient vector associated with the i-th and j-th basis set function.

virtual void compute\_hessian (void)=0

Compute Hessian matrix (independent on the parameters)

# **Additional Inherited Members**

### 17.63.1 Detailed Description

Implements a class of density matrix susceptibility models for parameterization in the uniform electric field.

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

- · oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_uniform\_base.cc

# 17.64 oepdev::UnitaryOptimizer Class Reference

Find the optimim unitary matrix of quadratic matrix equation.

#include <unitary\_optimizer.h>

#### **Public Member Functions**

• UnitaryOptimizer (double \*R, double \*P, int n, double conv=1.0e-6, int maxiter=100, bool verbose=true)

Create from R and P matrices and optimization options.

UnitaryOptimizer (std::shared\_ptr< psi::Matrix > R, std::shared\_ptr< psi::Vector > P, double conv=1.0e-6, int maxiter=100, bool verbose=true)

Create from R and P matrices and optimization options.

∼UnitaryOptimizer ()

Clear memory.

bool maximize ()

Run the minimization.

• bool minimize ()

Run the maximization.

std::shared\_ptr< psi::Matrix > X ()

Get the unitary matrix (solution)

double \* get\_X () const

Get the unitary matrix (pointer to solution)

double Z ()

Get the actual value of Z function.

• bool success () const

Get the status of the optimization.

#### **Protected Member Functions**

UnitaryOptimizer (int n, double conv, int maxiter, bool verbose)

Initialize the basic memory.

void common\_init\_ ()

Prepare the optimizer.

void run\_ (const std::string &opt)

Run the optimization (intermediate interface)

void optimize\_ (const std::string &opt)

Run the optimization (inner interface)

void refresh\_ ()

Restore the initial state of the optimizer.

void update\_conv\_ ()

Update the convergence.

void update\_iter\_ ()

Update the iterates.

void update\_Z\_ ()

Update Z value.

void update\_RP\_ ()

Uptade R and P matrices.

void update\_X\_ ()

Update the solution matrix X.

double eval\_Z\_ (double \*X, double \*R, double \*P)

Evaluate the objective Z function.

- double eval\_Z\_ ()
- double eval\_dZ\_ (double g, double \*R, double \*P, int i, int j)

Evaluate the change in Z.

double eval\_Z\_trial\_ (int i, int j, double gamma)

Evaluate the trial Z value.

void form\_X0\_ ()

Create identity matrix.

void form\_X\_ (int i, int j, double gamma)

Form unitary matrix X (store in buffer Xnew\_)

void form\_next\_X\_ (const std::string &opt)

Form the next unitary matrix X.

ABCD get\_ABCD\_ (int i, int j)

Retrieve ABCD parameters for root search.

void find\_roots\_boyd\_ (const ABCD &abcd)

Solve for all roots of equation A\*sin(g) + B\*cos(g) + C\*sin(2\*g) + D\*cos(2\*g) = 0 -> implements Boyd's method.

double find\_root\_halley\_ (double x0, const ABCD &abcd)

Solve for root of equation A\*sin(g) + B\*cos(g) + C\*sin(2\*g) + D\*cos(2\*g) = 0 -> implements Halley's method.

double find\_gamma\_ (const ABCD & abcd, int i, int j, const std::string & opt)

Compute gamma from roots of base equations.

bool It\_ (double a, double b)

less-than function

bool gt\_ (double a, double b)

greater-than function

double func\_0\_ (double g, const ABCD &abcd)

Function f(gamma) = d(dZ)/dgamma.

double func\_1\_ (double g, const ABCD &abcd)

Gradient of f(gamma)

double func\_2\_ (double g, const ABCD &abcd)

Hessian of f(gamma) - used only for Halley method (not implemented since Boyd method is more suitable here)

std::shared\_ptr< psi::Matrix > psi\_X\_ ()

Form the Psi4 matrix with the transformation matrix.

### **Protected Attributes**

const int n\_

Dimension of the problem.

const double conv\_

Convergence.

const int maxiter\_

Maximum number of iterations.

const bool verbose\_

Verbose mode.

double \* R\_

R matrix.

double \* P\_

P vector.

double \* R0\_

Reference R matrix.

double \* P0\_

Reference P vector.

double \* X\_

X Matrix (accumulated solution)

double \* W\_

Work place.

double \* Xold\_

Temporary X matrix.

double \* Xnew\_

Temporary X matrix.

• int niter\_

Current number of iterations.

double S<sub>-</sub> [4]

Current solutions.

double Zinit\_

Initial Z value.

double Zold\_

Old Z value.

double Znew\_

New Z value.

double conv\_current\_

Current convergence.

bool success\_

Status of optimization.

# 17.64.1 Detailed Description

The objective function of the orthogonal matrix **X** 

$$Z(\mathbf{X}) \equiv \sum_{ijkl} X_{ij} X_{kl} R_{jl} - \sum_{ij} X_{ij} P_j$$

is optimized by using the Jacobi iteration algorithm. In the above equation,  $\mathbf{R}$  is a square, general real matrix of size  $N \times N$  whereas  $\mathbf{P}$  is a real vector of length N.

Algorithm.

Optimization of X is factorized into a sequence of 2-dimensional rotations with one real parameter  $\gamma$ :

$$\mathbf{X}^{\text{New}} = \mathbf{U}(\gamma) \cdot \mathbf{X}^{\text{Old}}$$

where

$$\mathbf{U}(\gamma) \equiv egin{pmatrix} \ddots & & & & & & \\ & \cos(\gamma) & \cdots & \sin(\gamma) & & & \\ & \vdots & \ddots & \vdots & & \\ & -\sin(\gamma) & \cdots & \cos(\gamma) & & & \\ & & & \ddots & & \\ & & & & & \ddots \end{pmatrix}$$

is the Jacobi transformation matrix constructed for the  $\mathit{I}$ th and  $\mathit{J}$ th element from the entire  $\mathit{N}$ -dimensional set. For the sake of algirithmic simplicity, every iteration after  $U(\gamma)$  has been formed,  $\mathbf{X}^{\mathrm{Old}}$  is for a while assumed to be an identity matrix and the  $\mathbf{R}$  matrix and  $\mathbf{P}$  vector are transformed according to the following formulae

$$\mathbf{R} \to \mathbf{U}\mathbf{R}\mathbf{U}^T$$
$$\mathbf{P} \to \mathbf{U}\mathbf{P}$$

The full transformation matrix is accumulated in the memory buffer until convergence.

In each iteration, the optimum angle  $\gamma$  is found as follows: First, the roots of the finite Fourier series

$$A\sin(\gamma) + B\cos(\gamma) + C\sin(2\gamma) + D\cos(2\gamma) = 0$$

are found. In the above equations, the expansion coefficients are given as

$$A = P_I + P_J - \sum_{k \neq I,J} (R_{Ik} + R_{Jk} + R_{kI} + R_{kJ})$$

$$B = P_I - P_J - \sum_{k \neq I,J} (R_{Ik} - R_{Jk} + R_{kI} - R_{kJ})$$

$$C = -2(R_{IJ} + R_{JI})$$

$$D = -2(R_{II} - R_{JJ})$$

and I,J are the chosen indices in the Jacobi iteration subspace. The roots are evaluated by applying the Boyd's method[1], in which they are given as

$$\gamma_n = \Re\left[-i\ln(\lambda_n)\right]$$

where  $\lambda_n$  is an eivenvalue of the following 4 by 4 complex matrix:

$$\begin{pmatrix} 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \\ -\frac{D+iC}{D-iC} & -\frac{B+iC}{D-iC} & 0 & -\frac{B-iC}{D-iC} \end{pmatrix}$$

Once the four roots of the Fourier series equation are found, one solution out of four is chosen which satisfies the global optimum condition, i.e., the largest increase/decrease in the objective function given by

$$\delta Z = A(1 - \cos(\gamma)) + B\sin(\gamma) + C\sin^2(\gamma) + \frac{D}{2}\sin(2\gamma)$$

The discrimination between the minimae/maximae is performed based on the evaluation of the Hessian of Z with respect to  $\gamma$ ,

$$\frac{\partial^2 Z}{\partial \gamma^2} = A\cos(\gamma) - B\sin(\gamma) + 2C\cos(2\gamma) - 2D\sin(2\gamma)$$

All the N(N-1)/2 unique pairs of molecular orbitals are checked and the optimal set of  $\gamma, I, J$  is chosen to construct  $\mathbf{X}^{\text{New}}$ .

#### References:

[1] Boyd, J.P.; J. Eng. Math. (2006) 56, pp. 203-219

#### 17.64.2 Constructor & Destructor Documentation

### **17.64.2.1 UnitaryOptimizer()** [1/3]

#### **Parameters**

R	- R matrix
Р	- P vector
n	- dimensionality of the problem ( $N$ )
conv	- convergence in the ${\it Z}$ function
maxiter	- maximum number of iterations
verbose	- whether print information of iteration process or not Sets up the optimizer.

#### **17.64.2.2 UnitaryOptimizer()** [2/3]

```
oepdev::UnitaryOptimizer::UnitaryOptimizer (
    std::shared_ptr< psi::Matrix > R,
    std::shared_ptr< psi::Vector > P,
    double conv = 1.0e-6,
    int maxiter = 100,
    bool verbose = true )
```

#### **Parameters**

R	- R matrix
Р	- P vector
conv	- convergence in the ${\it Z}$ function
maxiter	- maximum number of iterations
verbose	- whether print information of iteration process or not Sets up the optimizer.

### **17.64.2.3 UnitaryOptimizer()** [3/3]

```
oepdev::UnitaryOptimizer::UnitaryOptimizer (
    int n,
    double conv,
    int maxiter,
    bool verbose ) [protected]
```

#### **Parameters**

n	- dimensionality of the problem ( $N$ )
conv	- convergence in the $Z$ function
maxiter	- maximum number of iterations
verbose	- whether print information of iteration process or not Sets up the optimizer.

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

- oepdev/libutil/unitary\_optimizer.h
- oepdev/libutil/unitary\_optimizer.cc

# 17.65 oepdev::UnitaryOptimizer\_4\_2 Class Reference

Find the optimim unitary matrix for quartic-quadratic matrix equation with trace.

```
#include <unitary_optimizer.h>
```

#### **Public Member Functions**

UnitaryOptimizer\_4\_2 (double \*R, double \*P, int n, double conv=1.0e-6, int maxiter=100, bool verbose=true)

Create from R and P matrices and optimization options.

~UnitaryOptimizer\_4\_2 ()

Clear memory.

• bool maximize ()

Run the minimization.

• bool minimize ()

Run the maximization.

std::shared\_ptr< psi::Matrix > X ()

Get the unitary matrix (solution)

double \* get\_X () const

Get the unitary matrix (pointer to solution)

double Z ()

Get the actual value of Z function.

• bool success () const

Get the status of the optimization.

### **Protected Member Functions**

• UnitaryOptimizer\_4\_2 (int n, double conv, int maxiter, bool verbose)

Initialize the basic memory.

void common\_init\_ ()

Prepare the optimizer.

void run\_ (const std::string &opt)

Run the optimization (intermediate interface)

void optimize\_ (const std::string &opt)

Run the optimization (inner interface)

void refresh\_ ()

Restore the initial state of the optimizer.

void update\_conv\_ ()

Update the convergence.

void update\_iter\_ ()

Update the iterates.

void update\_Z\_ ()

Update Z value.

void update\_RP\_ ()

Uptade R and P matrices.

void update\_X\_ ()

Update the solution matrix X.

double eval\_Z\_ (double \*X, double \*R, double \*P)

Evaluate the objective Z function.

- double eval\_Z\_ ()
- double eval\_dZ\_ (double g, double \*R, double \*P, int I, int J)

Evaluate the change in Z.

double eval\_Z\_trial\_ (int I, int J, double gamma)

Evaluate the trial Z value.

void form\_X0\_ ()

Create identity matrix.

void form\_X\_ (int I, int J, double gamma)

Form unitary matrix X (store in buffer Xnew\_)

void form\_next\_X\_ (const std::string &opt)

Form the next unitary matrix X.

• Fourier9 get\_fourier\_ (int I, int J)

Retrieve ABCD parameters for root search.

void find\_roots\_boyd\_ (const Fourier9 &abcd)

Solve for all roots of equation A\*sin(g) + B\*cos(g) + C\*sin(2\*g) + D\*cos(2\*g) = 0 -> implements Boyd's method.

double find\_root\_halley\_ (double x0, const Fourier9 &abcd)

Solve for root of equation A\*sin(g) + B\*cos(g) + C\*sin(2\*g) + D\*cos(2\*g) = 0 -> implements Halley's method.

double find\_gamma\_ (const Fourier9 &abcd, int i, int j, const std::string &opt)

Compute gamma from roots of base equations.

bool It\_ (double a, double b)

less-than function

bool gt\_ (double a, double b)

greater-than function

std::shared\_ptr< psi::Matrix > psi\_X\_ ()

Form the Psi4 matrix with the transformation matrix.

### **Protected Attributes**

const int n\_

Dimension of the problem.

const double conv\_

Convergence.

const int maxiter\_

Maximum number of iterations.

const bool verbose\_

Verbose mode.

double \* R\_

R tensor.

double \* P\_

P tensor.

double \* R0\_

Reference R tensor.

double \* P0\_

Reference P tensor.

double \* X\_

X Matrix (accumulated solution)

double \* W\_

Work place.

double \* Xold\_

Temporary X matrix.

double \* Xnew\_

Temporary X matrix.

int niter\_

Current number of iterations.

double S<sub>-</sub> [8]

Current solutions.

double Zinit\_

Initial Z value.

double Zold\_

Old Z value.

double Znew\_

New Z value.

double conv\_current\_

Current convergence.

bool success\_

Status of optimization.

# 17.65.1 Detailed Description

The objective function of the orthogonal matrix **X** 

$$Z(\mathbf{X}) \equiv \sum_{ijklmn} X_{ki} X_{lj} X_{mi} X_{nj} R_{ijklmn} + \sum_{ijk} X_{ji} X_{ki} P_{ijk}$$

is optimized by using the Jacobi iteration algorithm. In the above equation,  $\mathbf{R}$  is a general real sixth-rank tensor of size  $N^6$  whereas  $\mathbf{P}$  is a general real third-rank tensor of size  $N^3$ .

Algorithm.

Optimization of X is factorized into a sequence of 2-dimensional rotations with one real parameter  $\gamma$ :

$$\mathbf{X}^{\text{New}} = \mathbf{X}^{\text{Old}} \cdot \mathbf{U}(\gamma)$$

where

$$\mathbf{U}(\gamma) \equiv egin{pmatrix} \ddots & & & & & & \\ & \cos(\gamma) & \cdots & \sin(\gamma) & & & \\ & \vdots & \ddots & \vdots & & \\ & -\sin(\gamma) & \cdots & \cos(\gamma) & & & \\ & & & \ddots & & \\ & & & & & \ddots \end{pmatrix}$$

is the Jacobi transformation matrix constructed for the  $\mathit{I}$ th and  $\mathit{J}$ th element from the entire  $\mathit{N}$ -dimensional set. For the sake of algirithmic simplicity, every iteration after  $U(\gamma)$  has been formed,  $\mathbf{X}^{\mathrm{Old}}$  is for a while assumed to be an identity matrix and the  $\mathbf{R}$  as well as  $\mathbf{P}$  tensors are transformed according to the following formulae

$$egin{align*} R_{ijklmn} &
ightarrow \sum_{k'l'm'n'} R_{ijk'l'm'n'} X_{k'k} X_{l'l} X_{m'm} X_{n'n} \ P_{ijk} &
ightarrow \sum_{j'k'} P_{ij'k'} X_{j'j} X_{k'k} \ \end{aligned}$$

The full transformation matrix is accumulated in the memory buffer until convergence.

In each iteration, the optimum angle  $\gamma$  is found as follows: First, the roots of the finite Fourier series

$$a_0 + \sum_{p=1}^{4} \left\{ a_p \cos(px) + b_p \sin(px) \right\} = 0$$

are found. In the above equations, the expansion coefficients are calculated analytically as a function of I, J - the chosen indices in the Jacobi iteration subspace. The roots are evaluated by applying the Boyd's method[1], in which they are given as

$$\gamma_n = \Re\left[-i\ln(\lambda_n)\right]$$

where  $\lambda_n$  is an eivenvalue of the following 8 by 8 complex matrix:

$$\begin{pmatrix} 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ -\frac{a_4+ib_4}{a_4-ib_4} & -\frac{a_3+ib_3}{a_4-ib_4} & -\frac{a_2+ib_2}{a_4-ib_4} & -\frac{a_1+ib_1}{a_4-ib_4} & -\frac{2a_0}{a_4-ib_4} & -\frac{a_1-ib_1}{a_4-ib_4} & -\frac{a_2-ib_2}{a_4-ib_4} & -\frac{a_3-ib_3}{a_4-ib_4} \end{pmatrix}$$

Once the eight roots of the Fourier series equation are found, one solution out of eight is chosen which satisfies the global optimum condition, i.e., the largest increase/decrease in the objective function given by

$$\delta Z = Z(\mathbf{U}(\gamma)) - Z(\mathbf{1})$$

The Hessian is not computed. All the N(N-1)/2 unique pairs of molecular orbitals are checked and the optimal set of  $\gamma$ , I, J is chosen to construct  $\mathbf{X}^{\text{New}}$ .

#### References:

```
[1] Boyd, J.P.; J. Eng. Math. (2006) 56, pp. 203-219
```

### 17.65.2 Constructor & Destructor Documentation

# 17.65.2.1 UnitaryOptimizer\_4\_2() [1/2]

### **Parameters**

R	- <b>R</b> tensor (flattened row-wise)
Р	- P tensor (flattened row-wise)
n	- dimensionality of the problem ( $N$ )
conv	- convergence in the $Z$ function
maxiter	- maximum number of iterations
verbose	- whether print information of iteration process or not Sets up the optimizer.

### 17.65.2.2 UnitaryOptimizer\_4\_2() [2/2]

#### **Parameters**

n	dimensionality of the problem ( $N$ )	
conv	- convergence in the $Z$ function	
maxiter	- maximum number of iterations	
verbose	- whether print information of iteration process or not Sets up the optimizer.	

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

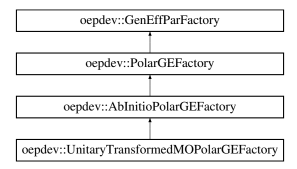
- oepdev/libutil/unitary\_optimizer.h
- · oepdev/libutil/unitary\_optimizer.cc

# 17.66 oepdev::UnitaryTransformedMOPolarGEFactory Class Reference

Polarization GEFP Factory with Least-Squares Scaling of MO Space.

#include <gefp.h>

Inheritance diagram for oepdev::UnitaryTransformedMOPolarGEFactory:



#### **Public Member Functions**

UnitaryTransformedMOPolarGEFactory (std::shared\_ptr< psi::Wavefunction > wfn, psi::Options & opt)

Construct from CPHF object and Psi4 options.

- virtual ~UnitaryTransformedMOPolarGEFactory ()
  - Destruct.
- std::shared\_ptr< GenEffPar > compute (void)

Pefrorm Least-Squares Fit.

#### **Additional Inherited Members**

### 17.66.1 Detailed Description

Implements creation of the density matrix susceptibility tensors for which  $X \neq 1$ . Guarantees the idempotency of the density matrix up to first-order in LCAO-MO variation.

#### Note

This method does not give better results than the X=1 method and is extremely time and memory consuming. Therefore, it is placed here only for future reference about solving unitary optimization problem in case it occurs.

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

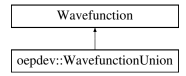
- · oepdev/libgefp/gefp.h
- oepdev/libgefp/gefp\_polar\_abinitio.cc

# 17.67 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.

WavefunctionUnion (SharedMolecule dimer, SharedBasisSet primary, SharedBasisSet auxiliary\_df, SharedBasisSet primary\_1, SharedBasisSet primary\_2, SharedBasisSet auxiliary\_1, SharedBasisSet auxiliary\_df\_1, SharedBasisSet auxiliary\_df\_2, SharedBasisSet intermediate\_1, SharedBasisSet intermediate\_2, SharedWavefunction wfn\_1, SharedWavefunction wfn\_2, 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)

void clear\_dpd ()

Close the DPD instance.

• int l\_nmo (int n) const

Get number of molecular orbitals of the \*n\*th fragment.

int l\_nso (int n) const

Get number of symmetry orbitals of the \*n\*th fragment.

• int l\_ndocc (int n) const

Get number of doubly occupied orbitals of the \*n\*th fragment.

int l\_nvir (int n) const

Get number of virtual orbitals of the \*n\*th fragment.

int l\_nalpha (int n) const

Get the number of the alpha electrons of the \*n\*th fragment.

int l\_nbeta (int n) const

Get the number of the beta electrons of the \*n\*th fragment.

int l\_nbf (int n) const

Get number of basis functions of the \*n\*th fragment.

• int l\_noffs\_ao (int n) const

Get the basis set offset of the \*n\*th fragment.

double l\_energy (int n) const

Get the reference energy of the \*n\*th fragment.

SharedMolecule l\_molecule (int n) const

Get the molecule object of the \*n\*th fragment.

SharedBasisSet I\_primary (int n) const

Get the primary basis set object of the \*n\*th fragment.

SharedBasisSet I\_auxiliary (int n) const

Get the auxiliary basis set object of the \*n\*th fragment.

SharedBasisSet Lintermediate (int n) const

Get the intermediate basis set object of the \*n\*th fragment.

SharedWavefunction Lwfn (int n) const

Get the wavefunction object of the \*n\*th fragment.

SharedMOSpace I\_mospace (int n, const std::string &label) const

Get the MO space named label (either OCC or VIR) of the \*n\*th fragment.

SharedLocalizer Llocalizer (int n) const

Get the orbital localizer object of the \*n\*th fragment.

SharedIntegralTransform integrals (void) const

Get the integral transform object of the entire union.

bool has\_localized\_orbitals (void) const

If union got its molecular orbital localized or not.

SharedBasisSet primary (void) const

Get the primary basis set for the entire union.

SharedMOSpace mospace (const std::string &label) const

Get the MO space named label (either OCC or VIR)

- SharedMatrix Ca\_subset (const std::string &basis="SO", const std::string &subset="ALL")
- SharedMatrix Cb\_subset (const std::string &basis="SO", const std::string &subset="ALL")
- SharedMatrix C\_subset\_helper (SharedMatrix C, const Dimension &noccpi, SharedVector epsilon, const std::string &basis, const std::string &subset)

Helpers for Ca\_ and Cb\_ matrix transformers.

 SharedVector epsilon\_subset\_helper (SharedVector epsilon, const Dimension &noccpi, const std::string &basis, const std::string &subset)

Helper for epsilon transformer.

void print\_header (void)

Print information about this wavefunction union.

void print\_mo\_integrals (void)

Print the MO ingegrals.

#### **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::map< const std::string, SharedMOSpace > mospacesUnion\_

Dictionary of MO spaces for the entire union (OCC and VIR)

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 < SharedBasisSet > Lintermediate\_

List of intermediate basis functions per molecule.

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

List of original isolated wavefunctions (electrons unrelaxed)

std::vector< std::string > l\_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 > l\_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 > l\_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 > Llocalizer\_

List of orbital localizers.

std::vector< std::map< const std::string, SharedMOSpace >> I\_mospace\_

List of dictionaries of MO spaces.

std::shared\_ptr< psi::OEProp > oeprop\_

One-Electron Property.

# 17.67.1 Detailed Description

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
- 1. Performs Hadamard sums on H\_, Fa\_, Da\_, Ca\_ and S\_ based on uncoupled wavefunctions.
- 2. 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!
- 3. Ca/Cb and epsilon subsets were reimplemented from psi::Wavefunction to remove sorting of orbitals. However, the corresponding member functions are not virtual in psi::Wavefunction. This could bring problems when upcasting.

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.

# 17.67.2 Constructor & Destructor Documentation

# **17.67.2.1** WavefunctionUnion() [1/2]

Provide wavefunction with molecule containing at least 2 fragments.

#### **Parameters**

ref_wfn	- reference wavefunction
options	- Psi4 options

#### **17.67.2.2** WavefunctionUnion() [2/2]

Provide molecule dimer and all the required monomer basis sets and wavefunctions.

#### **Parameters**

dimer	- molecule object
primary	- basis set object
auxiliary_df	- basis set object (for DF SCF)
primary_1	- basis set object for 1st monomer
primary_2	- basis set object for 2nd monomer
auxiliary_1	- basis set object for 1st monomer
auxiliary_2	- basis set object for 2nd monomer

#### **Parameters**

auxiliary_df_1	- basis set object for 1st monomer
auxiliary_df_2	- basis set object for 2nd monomer
intermediate <sub>-</sub> 1	- basis set object for 1st monomer
intermediate_2	- basis set object for 2nd monomer
wfn_1	- unperturbed wavefunction object
wfn_2	- unperturbed wavefunction object
options	- Psi4 options

# 17.67.3 Member Function Documentation

## 17.67.3.1 Ca\_subset()

Return a subset of the Ca matrix in a desired basis

#### **Parameters**

basis	the symmetry basis to use AO, SO	
subset	the subset of orbitals to return ALL, ACTIVE, FROZEN, OCC, VIR, FROZEN_OCC,	
	ACTIVE_OCC, ACTIVE_VIR, FROZEN_VIR	

#### **Returns**

the matrix in Pitzer order in the desired basis

# 17.67.3.2 Cb\_subset()

Return a subset of the Cb matrix in a desired basis

#### **Parameters**

basis	the symmetry basis to use AO, SO
subset	the subset of orbitals to return ALL, ACTIVE, FROZEN, OCC, VIR, FROZEN_OCC,
	ACTIVE_OCC, ACTIVE_VIR, FROZEN_VIR

# Returns

the matrix in Pitzer order in the desired basis

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

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

# **Chapter 18**

# **File Documentation**

# 18.1 include/oepdev\_files.h File Reference

#### **Macros**

#define OEPDEV\_USE\_PSI4\_DIIS\_MANAGER 0

Use DIIS from Psi4 (1) or OEPDev (0)?

#define OEPDEV\_MAX\_AM 8

L\_max.

• #define OEPDEV\_N\_MAX\_AM 17

 $2L_max+1$ 

#define OEPDEV\_CRIT\_ERI 1e-9

ERI criterion for E12, E34, E123 and lambda\*EXY coefficients.

#define OEPDEV\_SIZE\_BUFFER\_R 250563

Size of R buffer (OEPDEV\_N\_MAX\_AM\*OEPDEV\_N\_MAX\_AM\*OEPDEV\_N\_MAX\_AM\*OEPDEV\_N\_MAX\_AM\*3)

#define OEPDEV\_SIZE\_BUFFER\_D2 3264

Size of D2 buffer (3\*(OEPDEV\_MAX\_AM+1)\*(OEPDEV\_MAX\_AM+1)\*OEPDEV\_N\_MAX\_AM)

# 18.2 include/oepdev\_options.h File Reference

# **Namespaces**

• psi

Psi4 package namespace.

# **Functions**

PSI\_API int psi::read\_options (std::string name, Options & options)
 Options for the OEPDev plugin.

232 File Documentation

# 18.3 main.cc File Reference

```
#include <string>
#include "include/oepdev_files.h"
#include "psi4/liboptions/liboptions.h"
#include "psi4/libmints/wavefunction.h"
#include "include/oepdev_options.h"
#include "oepdev/liboep/oep.h"
#include "oepdev/libgefp/gefp.h"
#include "oepdev/libsolver/solver.h"
#include "oepdev/libtest/test.h"
#include <pybind11/pybind11.h>
```

# **Namespaces**

• psi

Psi4 package namespace.

# **Typedefs**

- using SharedWavefunction = std::shared\_ptr< psi::Wavefunction >
- using SharedUnion = std::shared\_ptr< oepdev::WavefunctionUnion >
- using SharedOEPotential = std::shared\_ptr< oepdev::OEPotential >
- using SharedGEFPFactory = std::shared\_ptr< oepdev::GenEffParFactory >
- using SharedGEFPParameters = std::shared\_ptr< oepdev::GenEffPar >

#### **Functions**

- void psi::export\_dmtp (py::module &)
- void psi::export\_cphf (py::module &)
- void psi::export\_solver (py::module &)
- PSI\_API SharedWavefunction psi::oepdev (SharedWavefunction ref\_wfn, Options &options)

Main routine of the OEPDev plugin.

psi::PYBIND11\_MODULE (oepdev, m)

# 18.4 oepdev/lib3d/dmtp.h File Reference

```
#include "psi4/libmints/wavefunction.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/molecule.h"
```

```
#include "psi4/libmints/matrix.h"
#include "psi4/libmints/vector.h"
```

#### **Classes**

· class oepdev::MultipoleConvergence

Multipole Convergence.

class oepdev::DMTPole

Distributed Multipole Analysis Container and Computer. Abstract Base.

· class oepdev::CAMM

Cumulative Atomic Multipole Moments.

# Namespaces

• psi

Psi4 package namespace.

• oepdev

OEPDev module namespace.

# **Typedefs**

using psi::SharedBasisSet = std::shared\_ptr< BasisSet >

# 18.5 oepdev/lib3d/esp.h File Reference

```
#include "space3d.h"
```

#### **Classes**

· class oepdev::ESPSolver

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

### **Namespaces**

oepdev

OEPDev module namespace.

234 File Documentation

# **Typedefs**

using oepdev::SharedField3D = std::shared\_ptr< oepdev::Field3D >

# 18.6 oepdev/libgefp/gefp.h File Reference

```
#include <vector>
#include <string>
#include <random>
#include <cmath>
#include "psi4/libpsi4util/PsiOutStream.h"
#include "psi4/libpsi4util/process.h"
#include "psi4/libmints/wavefunction.h"
#include "psi4/libmints/watrix.h"
#include "psi4/libmints/vector.h"
#include "psi4/libmints/vector3.h"
#include "../libmints/vector3.h"
#include "../libutil/util.h"
#include "../libutil/util.h"
#include "../libutil/cphf.h"
#include "../libutil/scf_perturb.h"
```

### **Classes**

· class oepdev::GenEffPar

Generalized Effective Fragment Parameters. Container Class.

class oepdev::GenEffFrag

Generalized Effective Fragment. Container Class.

class oepdev::GenEffParFactory

Generalized Effective Fragment Factory. Abstract Base.

class oepdev::PolarGEFactory

Polarization GEFP Factory. Abstract Base.

class oepdev::AbInitioPolarGEFactory

Polarization GEFP Factory from First Principles. Hartree-Fock Approximation.

class oepdev::FFAbInitioPolarGEFactory

Polarization GEFP Factory from First Principles: Finite-Difference Model. Arbitrary level of theory.

class oepdev::GeneralizedPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

struct oepdev::GeneralizedPolarGEFactory::StatisticalSet

A structure to handle statistical data.

class oepdev::UniformEFieldPolarGEFactory

Polarization GEFP Factory with Least-Squares Parameterization.

- class oepdev::NonUniformEFieldPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Parameterization.
- class oepdev::LinearUniformEFieldPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Parameterization.
- class oepdev::QuadraticUniformEFieldPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Parameterization.
- class oepdev::LinearNonUniformEFieldPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Parameterization.
- · class oepdev::QuadraticNonUniformEFieldPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Parameterization.
- class oepdev::LinearGradientNonUniformEFieldPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Parameterization.
- class oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Parameterization.
- class oepdev::UnitaryTransformedMOPolarGEFactory
  - Polarization GEFP Factory with Least-Squares Scaling of MO Space.

# **Namespaces**

oepdev

OEPDev module namespace.

# 18.7 oepdev/libints/eri.h File Reference

```
#include "psi4/libpsi4util/exception.h"
#include "psi4/libmints/integral.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/fjt.h"
#include "../libpsi/integral.h"
#include "recurr.h"
```

#### Classes

- class oepdev::TwoElectronInt
  - General Two Electron Integral.
- class oepdev::ERI\_1\_1
  - 2-centre ERI of the form (a|O(2)|b) where O(2) = 1/r12.
- class oepdev::ERI\_2\_2
  - 4-centre ERI of the form (ab|O(2)|cd) where O(2) = 1/r12.
- class oepdev::ERI\_3\_1
  - 4-centre ERI of the form (abc|O(2)|d) where O(2) = 1/r12.

236 File Documentation

# **Namespaces**

oepdev

OEPDev module namespace.

# 18.8 oepdev/libints/recurr.h File Reference

# **Namespaces**

oepdev

OEPDev module namespace.

#### **Macros**

#define D1\_INDEX(x, i, n) ((81\*(x))+(9\*(i))+(n))

Get the index of McMurchie-Davidson-Hermite D1 coefficient stored in the  $mdh\_buffer\_$ , that is attributed to the x Cartesian coordinate from angular momentum i of function 1, and the Hermite index n.

• #define D2\_INDEX(x, i, j, n) ((1377\*(x))+(153\*(i))+(17\*(j))+(n))

Get the index of McMurchie-Davidson-Hermite D2 coefficient stored in the mdh\_buffer\_, that is attributed to the x Cartesian coordinate from angular momenta i, j of function 1 and 2, and the Hermite index n.

#define D3\_INDEX(x, i, j, k, n) ((18225\*(x))+(2025\*(i))+(225\*(j))+(25\*(k))+(n))

Get the index of McMurchie-Davidson-Hermite D3 coefficient stored in the mdh\_buffer\_, that is attributed to the x Cartesian coordinate from angular momenta i, j and k of function 1, 2 and 3, and the Hermite index n.

#define R\_INDEX(n, I, m, j) ((14739\*(n))+(867\*(l))+(51\*(m))+(j))

Get the index of McMurchie-Davidson R coefficient stored in the mdh\_buffer\_R\_ from angular momenta n, I and m and the Boys index j.

### **Functions**

- double oepdev::d\_N\_n1\_n2 (int N, int n1, int n2, double PA, double PB, double aP)

  Compute McMurchie-Davidson-Hermite (MDH) coefficient for binomial expansion.
- void oepdev::make\_mdh\_D1\_coeff (int n1, double aPd, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for monomial expansion.

void oepdev::make\_mdh\_D2\_coeff (int n1, int n2, double aPd, double \*PA, double \*PB, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for binomial expansion.

void oepdev::make\_mdh\_D3\_coeff (int n1, int n2, int n3, double aPd, double \*PA, double \*PB, double \*PC, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for trinomial expansion.

void oepdev::make\_mdh\_D2\_coeff\_explicit\_recursion (int n1, int n2, double aP, double \*PA, double \*PB, double \*buffer)

Compute the McMurchie-Davidson-Hermite coefficients for binomial expansion by explicit recursion. This function makes the same changes to buffers as oepdev::make\_mdh\_D2\_coeff, but implements it through explicit recursion by calls to oepdev::d\_N\_n1\_n2. Therefore, it is slightly slower. Here for debugging purposes.

void oepdev::make\_mdh\_R\_coeff (int N, int L, int M, double alpha, double a, double b, double c, double \*F, double \*buffer)

Compute the McMurchie-Davidson R coefficients.

# 18.9 oepdev/liboep/oep.h File Reference

```
#include <cstdio>
#include <string>
#include <vector>
#include <map>
#include "psi4/liboptions/liboptions.h"
#include "psi4/libpsi4util/PsiOutStream.h"
#include "psi4/libmints/wavefunction.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/vector.h"
#include "psi4/libmints/vector.h"
#include "../libpsi/integral.h"
#include "../libpsi/potential.h"
#include "../lib3d/space3d.h"
#include "../lib3d/dmtp.h"
```

#### Classes

struct oepdev::OEPType

Container to handle the type of One-Electron Potentials.

· class oepdev::OEPotential

Generalized One-Electron Potential: Abstract base.

• class oepdev::ElectrostaticEnergyOEPotential

Generalized One-Electron Potential for Electrostatic Energy.

class oepdev::RepulsionEnergyOEPotential

Generalized One-Electron Potential for Pauli Repulsion Energy.

class oepdev::ChargeTransferEnergyOEPotential

Generalized One-Electron Potential for Charge-Transfer Interaction Energy.

class oepdev::EETCouplingOEPotential

Generalized One-Electron Potential for EET coupling calculations.

## **Namespaces**

· oepdev

OEPDev module namespace.

## **Typedefs**

- using oepdev::SharedWavefunction = std::shared\_ptr< Wavefunction >
- using oepdev::SharedBasisSet = std::shared\_ptr< BasisSet >
- using oepdev::SharedMatrix = std::shared\_ptr< Matrix >
- using oepdev::SharedVector = std::shared\_ptr< Vector >
- using oepdev::SharedDMTPole = std::shared\_ptr< DMTPole >

# 18.10 oepdev/liboep/oep\_gdf.h File Reference

```
#include <cstdio>
#include <string>
#include "psi4/liboptions/liboptions.h"
#include "psi4/libmints/wavefunction.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/matrix.h"
#include "../libpsi/integral.h"
```

#### Classes

class oepdev::GeneralizedDensityFit

Generalized Density Fitting Scheme. Abstract Base.

class oepdev::SingleGeneralizedDensityFit

Generalized Density Fitting Scheme - Single Fit.

• class oepdev::DoubleGeneralizedDensityFit

Generalized Density Fitting Scheme - Double Fit.

## **Namespaces**

oepdev

# 18.11 oepdev/libpsi/integral.h File Reference

```
#include "psi4/libmints/integral.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/matrix.h"
```

## **Classes**

- · class oepdev::TwoBodyAOInt
- · class oepdev::IntegralFactory

Extended IntegralFactory for computing integrals.

## **Namespaces**

oepdev

OEPDev module namespace.

# 18.12 oepdev/libpsi/potential.h File Reference

```
#include <vector>
#include "psi4/psi4-dec.h"
#include "psi4/liboptions/liboptions.h"
#include "psi4/libpsio/psio.h"
#include "psi4/libmints/typedefs.h"
#include "psi4/libmints/onebody.h"
#include "psi4/libmints/potential.h"
#include "psi4/libmints/sointegral_onebody.h"
#include "psi4/libmints/osrecur.h"
```

#### Classes

· class oepdev::PotentialInt

Computes potential integrals.

## **Namespaces**

oepdev

# 18.13 oepdev/libsolver/solver.h File Reference

```
#include <cstdio>
#include <string>
#include <map>
#include "psi4/psi4-dec.h"
#include "psi4/liboptions/liboptions.h"
#include "psi4/libpsio/psio.h"
#include "psi4/libmints/potential.h"
#include "psi4/libmints/integral.h"
#include "psi4/libpsi4util/PsiOutStream.h"
#include "../libutil/wavefunction_union.h"
#include "../libutil/integrals_iter.h"
#include "../libpsi/integral.h"
#include "../liboep/oep.h"
```

### Classes

· class oepdev::OEPDevSolver

Solver of properties of molecular aggregates. Abstract base.

class oepdev::ElectrostaticEnergySolver

Compute the Coulombic interaction energy between unperturbed wavefunctions.

class oepdev::RepulsionEnergySolver

Compute the Pauli-Repulsion interaction energy between unperturbed wavefunctions.

class oepdev::ChargeTransferEnergySolver

Compute the Charge-Transfer interaction energy between unperturbed wavefunctions.

## **Namespaces**

oepdev

OEPDev module namespace.

## **Typedefs**

- using oepdev::SharedWavefunctionUnion = std::shared\_ptr< WavefunctionUnion >
- using oepdev::SharedOEPotential = std::shared\_ptr< OEPotential >

# 18.14 oepdev/libtest/test.h File Reference

```
#include <vector>
#include "psi4/psi4-dec.h"
```

```
#include "psi4/liboptions/liboptions.h"
#include "psi4/libpsio/psio.h"
#include "psi4/libpsi4util/PsiOutStream.h"
#include "psi4/libmints/integral.h"
#include "psi4/libmints/matrix.h"
#include "psi4/libmints/wavefunction.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libqt/qt.h"
#include "../libpsi/integral.h"
#include "../libpsi/integrals_iter.h"
```

#### **Classes**

class oepdev::test::Test
 Manages test routines.

## **Namespaces**

· oepdev

OEPDev module namespace.

# 18.15 oepdev/libutil/diis.h File Reference

```
#include <cstdio>
#include <string>
#include <vector>
#include "psi4/libciomr/libciomr.h"
#include "psi4/libmints/vector.h"
#include "psi4/libmints/matrix.h"
#include "psi4/libqt/qt.h"
#include "psi4/libpsi4util/PsiOutStream.h"
```

## **Classes**

class oepdev::DIISManager
 DIIS manager.

### **Namespaces**

oepdev

# 18.16 oepdev/libutil/integrals\_iter.h File Reference

```
#include <cstdio>
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/integral.h"
#include "../libpsi/integral.h"
```

#### Classes

- class oepdev::ShellCombinationsIterator
   Iterator for Shell Combinations. Abstract Base.
- class oepdev::AOIntegralsIterator
   Iterator for AO Integrals. Abstract Base.
- class oepdev::AllAOShellCombinationsIterator\_4
   Loop over all possible ERI shells in a shell quartet.
- class oepdev::AllAOShellCombinationsIterator\_2
   Loop over all possible ERI shells in a shell doublet.
- class oepdev::AllAOIntegralsIterator\_4
   Loop over all possible ERI within a particular shell quartet.
- class oepdev::AllAOIntegralsIterator\_2
   Loop over all possible ERI within a particular shell doublet.

#### Namespaces

oepdev

OEPDev module namespace.

## **Typedefs**

- using oepdev::SharedIntegralFactory = std::shared\_ptr< IntegralFactory >
- using oepdev::SharedTwoBodyAOInt = std::shared\_ptr< TwoBodyAOInt >
- using oepdev::SharedShellsIterator = std::shared\_ptr< ShellCombinationsIterator >
   Iterator over shells as shared pointer.
- using oepdev::SharedAOIntsIterator = std::shared\_ptr< AOIntegralsIterator >
   Iterator over AO integrals as shared pointer.

# 18.17 oepdev/libutil/scf\_perturb.h File Reference

```
#include "psi4/libmints/vector.h"
#include "psi4/libmints/matrix.h"
#include "psi4/libscf_solver/rhf.h"
```

#### Classes

• struct oepdev::PerturbCharges

Structure to hold perturbing charges.

· class oepdev::RHFPerturbed

RHF theory under electrostatic perturbation.

## **Namespaces**

oepdev

OEPDev module namespace.

# 18.18 oepdev/libutil/unitary\_optimizer.h File Reference

```
#include <string>
#include <complex>
#include "psi4/libmints/vector.h"
#include "psi4/libmints/matrix.h"
```

#### Classes

struct oepdev::ABCD

Simple structure to hold the Fourier series expansion coefficients.

struct oepdev::Fourier9

Simple structure to hold the Fourier series expansion coefficients for N=4.

class oepdev::UnitaryOptimizer

Find the optimim unitary matrix of quadratic matrix equation.

class oepdev::UnitaryOptimizer\_4\_2

Find the optimim unitary matrix for quartic-quadratic matrix equation with trace.

## **Namespaces**

oepdev

#### **Macros**

```
#define IDX(i, j, n) ((n)*(i)+(j))
#define IDX3(i, j, k) (n2_*(i)+n_*(j)+(k))
#define IDX6(i, j, k, l, m, n) (n5_*(i)+n4_*(j)+n3_*(k)+n2_*(l)+n_*(m)+(n))
```

#### **Functions**

- constexpr std::complex< double > oepdev::operator""\_i (unsigned long long d)
- constexpr std::complex< double > oepdev::operator""\_i (long double d)

# 18.19 oepdev/libutil/util.h File Reference

```
#include <cstdio>
#include <string>
#include <cmath>
#include <map>
#include <cassert>
#include "psi4/psi4-dec.h"
#include "psi4/liboptions/liboptions.h"
#include "psi4/libpsio/psio.h"
#include "psi4/libciomr/libciomr.h"
#include "psi4/libpsio/psio.hpp"
#include "psi4/libiwl/iwl.h"
#include "psi4/libqt/qt.h"
#include "psi4/libmints/molecule.h"
#include "psi4/libmints/writer.h"
#include "psi4/libmints/writer_file_prefix.h"
#include "psi4/libmints/wavefunction.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/vector.h"
#include "psi4/libmints/matrix.h"
#include "psi4/libmints/oeprop.h"
#include "psi4/libmints/local.h"
#include "psi4/libfunctional/superfunctional.h"
#include "psi4/libtrans/mospace.h"
#include "psi4/libtrans/integraltransform.h"
#include "psi4/libscf_solver/rhf.h"
#include "psi4/libdpd/dpd.h"
```

## **Namespaces**

oepdev

## **Typedefs**

- using oepdev::SharedMolecule = std::shared\_ptr< Molecule >
- using oepdev::SharedSuperFunctional = std::shared\_ptr< SuperFunctional >
- using oepdev::SharedMOSpace = std::shared\_ptr< MOSpace >
- using oepdev::SharedMOSpaceVector = std::vector < std::shared\_ptr < MOSpace > >
- using oepdev::SharedIntegralTransform = std::shared\_ptr< IntegralTransform >
- using oepdev::SharedLocalizer = std::shared\_ptr< Localizer >

### **Functions**

PSI\_API void oepdev::preambule (void)

Print preambule for module OEPDEV.

template<typename... Args>
 std::string oepdev::string\_sprintf (const char \*format, Args... args)

Format string output. Example: std::string text = oepdev::string\_sprinff("Test %3d, %13.5f", 5, -10.5425);.

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

Set up DFT functional.

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

Extract molecule from dimer.

PSI\_API std::shared\_ptr< Wavefunction > oepdev::solve\_scf (std::shared\_ptr< Molecule > molecule, std::shared\_ptr< BasisSet > primary, std::shared\_ptr< BasisSet > auxiliary, std::shared\_ptr< SuperFunctional > functional, Options & options, std::shared\_ptr< PSIO > psio, bool compute\_mints=false)

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

PSI\_API double oepdev::average\_moment (std::shared\_ptr< psi::Vector > moment)

Compute the scalar magnitude of multipole moment.

# 18.20 oepdev/libutil/wavefunction\_union.h File Reference

```
#include <cstdio>
#include <string>
#include <map>
#include "psi4/psi4-dec.h"
#include "psi4/liboptions/liboptions.h"
#include "psi4/libpsio/psio.h"
#include "psi4/libciomr/libciomr.h"
#include "psi4/libpsio/psio.hpp"
#include "psi4/libpsio/psio.hpp"
#include "psi4/libqt/qt.h"
#include "psi4/libmints/molecule.h"
```

```
#include "psi4/libmints/writer.h"
#include "psi4/libmints/writer_file_prefix.h"
#include "psi4/libmints/wavefunction.h"
#include "psi4/libmints/basisset.h"
#include "psi4/libmints/vector.h"
#include "psi4/libmints/matrix.h"
#include "psi4/libmints/oeprop.h"
#include "psi4/libmints/local.h"
#include "psi4/libfunctional/superfunctional.h"
#include "psi4/libtrans/mospace.h"
#include "psi4/libtrans/integraltransform.h"
#include "psi4/libscf_solver/rhf.h"
#include "psi4/libdpd/dpd.h"
```

### Classes

• class oepdev::WavefunctionUnion

Union of two Wavefunction objects.

## **Namespaces**

oepdev

# Chapter 19

# **Example Documentation**

# 19.1 example\_cphf.cc

Shows how to use the oepdev::CPHF solver to compute molecular and LMO-distributed polarizabilities at RHF level of theory.

```
void example_cphf(std::shared_ptr<psi::Wavefunction> wfn, psi::Options& opt){
    // build the solver
    std::shared_ptr<oepdev::CPHF> solver = std::make_shared<oepdev::CPHF>(wfn, opt);

    // run the solver to converge CPHF equations
    solver->compute();

    // print the LMO-distributed polarizabilities
    for (int i=0; i<solver->nocc(); i++) {
            solver->polarizability(i)->print();
    }

    // print the molecular polarizability
    solver->polarizability()->print();

    // grab 4th LMO-distributed polarizability and its associated LMO centroid
    psi::SharedMatrix pol.4 = solver->polarizability(3);
    psi::SharedVector rmo.4 = solver->lmo_centroid(3);
};
```

# 19.2 example\_integrals\_iter.cc

Iterations over electron repulsion integrals in AO basis. This is an example of how to use

- the oepdev::ShellCombinationsIterator class
- the oepdev::AOIntegralsIterator class.

```
void iterate(std::shared_ptr<oepdev::IntegralFactory> ints)
{
    // Prepare for direct calculation of ERI's (shell by shell)
    std::shared_ptr<psi::TwoBodyAOInt> tei(ints->eri());

    // Grab the buffer where the integrals for a current shell will be placed
    const double* buffer = tei->buffer();
```

```
// Create iterator to go through all shell quartet combinations
oepdev::SharedShellsIterator shellIter =
    oepdev::ShellCombinationsIterator::build(ints, "ALL", 4);
// Iterate over shells, and then over all integrals in each shell quartet
for (shellIter->first(); shellIter->is_done() == false; shellIter->next())
      // Compute all integrals between shells in the current quartet
     shellIter->compute_shell(tei);
     // Create iterator to go through all integrals within a shell quartet
     oepdev::SharedAOIntsIterator intsIter = shellIter->ao_iterator("ALL");
     for (intsIter->first(); intsIter->is_done() == false; intsIter->next())
     {
          // Grab current (ij|kl) indices here
          int i = intsIter->i();
          int j = intsIter->j();
          int k = intsIter->k();
          int l = intsIter->l();
          // Grab the (ij|kl) integral
          double integral = buffer[intsIter->index()];
     }
}
```

# 19.3 example\_scf\_perturb.cc

Perturb HF Hamiltonian with external electrostatic potential. This is an example of how to use the oepdev::RHFPerturbed class.

```
void scf_perturb(std::shared_ptr<psi::Wavefunction> wfn, psi::Options& opt)
   // Set up HF superfunctional
  std::shared.ptr<psi::SuperFunctional> func = oepdev::create_superfunctional
      ("HF", opt);
   // Initialize the perturbed wavefunction
  std::shared_ptr<oepdev::RHFPerturbed> scf = std::make_shared<oepdev::RHFPerturbed>(wfn, func, opt, wfn->
     psio());
   /* Perturb the system with the uniform electric field [Fx, Fy, Fz].
     Then, add two point charges of charge qi placed at [Rxi, Ryi, Rzi].
     Provide all these values in atomic units! \star/
   const double Fx = 0.04, Fy = 0.05, Fz = -0.09;
  const double Rx1=0.00, Rx2=1.30, Rx3=-1.00;
  const double Rx1= 0.10, Rx2=-0.30, Rx3= 3.50;
  const double q1 = 0.30, q2 = -0.09;
   scf->set_perturbation(Fx, Fy, Fz);
                                             /* set it only once, setting it again will overwrite the
      field, not add */
   scf->set_perturbation(Rx1, Ry1, Rz1, q1);
  scf->set_perturbation(Rx2, Ry2, Rz2, q2); /* more charges can be added */
   // Solve perturbed SCF equations
  scf->compute_energy();
   // Grab some data
  double energy = scf->reference_energy();
                                                 // Total energy of the system
  std::shared.ptr<psi::Matrix> Da = scf->Da(); // One-particle density matrix
   /\star Note that the external field and charges perturb only one-electron Hamiltonian.\star/
}
```

# Index

AllAOIntegralsIterator_2	oepdev::TwoBodyAOInt, 206, 207
oepdev::AllAOIntegralsIterator_2, 79, 80	compute_benchmark
AllAOIntegralsIterator_4	oepdev::ChargeTransferEnergySolver, 95
oepdev::AllAOIntegralsIterator_4, 81, 82	oepdev::ElectrostaticEnergySolver, 117
AllAOShellCombinationsIterator_2	oepdev::OEPDevSolver, 165
oepdev::AllAOShellCombinationsIterator_2,	oepdev::RepulsionEnergySolver, 195
83, 84	compute_density_matrix
AllAOShellCombinationsIterator_4	oepdev::GenEffPar, 139, 140
oepdev::AllAOShellCombinationsIterator_4,	compute_oep_based
86, 87	oepdev::ChargeTransferEnergySolver, 95
allocate	oepdev::ElectrostaticEnergySolver, 118
oepdev::GenEffPar, 138	oepdev::OEPDevSolver, 165
ao_iterator	oepdev::RepulsionEnergySolver, 195
oepdev::ShellCombinationsIterator, 200	compute_shell
average_moment	oepdev::AllAOShellCombinationsIterator_2
The OEPDev Utilities, 65	85
	oepdev::AllAOShellCombinationsIterator_4
build	87
oepdev::AOIntegralsIterator, 89	oepdev::ShellCombinationsIterator, 201
oepdev::DMTPole, 107	oepdev::TwoElectronInt, 209
oepdev::Field3D, 131	ConvergenceLevel
oepdev::GenEffParFactory, 144	oepdev::MultipoleConvergence, 160
oepdev::GeneralizedDensityFit, 147	create_superfunctional
oepdev::OEPDevSolver, 164	The OEPDev Utilities, 65
oepdev::OEPotential, 169	
oepdev::Points3DIterator, 175, 176	d_N_n1_n2
oepdev::PointsCollection3D, 178, 179	The Integral Package Library, 54
oepdev::ShellCombinationsIterator, 201	DIISManager
CPHF	oepdev::DIISManager, 101
	DMTPole
oepdev::CPHF, 98	oepdev::DMTPole, 107
Ca_subset	ESPSolver
oepdev::WavefunctionUnion, 229 Cb_subset	oepdev::ESPSolver, 126, 127
	energy
oepdev::WavefunctionUnion, 229	oepdev::DMTPole, 108
compute	extract_monomer
oepdev::DIISManager, 102	The OEPDev Utilities, 66
oepdev::DoubleGeneralizedDensityFit, 112	Field3D
oepdev::GeneralizedDensityFit, 148	oepdev::Field3D, 131
oepdev::MultipoleConvergence, 161	oopacv lolado, lol
oepdev::SingleGeneralizedDensityFit, 203	include/oepdev_files.h, 231

250 INDEX

include/oepdev_options.h, 231	oepdev/libutil/util.h, 244
index	oepdev/libutil/wavefunction_union.h, 245
oepdev::AllAOIntegralsIterator_2, 80	oepdev::ABCD, 77
oepdev::AllAOIntegralsIterator_4, 82	oepdev::AOIntegralsIterator, 88
	build, 89
level	oepdev::AbInitioPolarGEFactory, 77
oepdev::MultipoleConvergence, 161	oepdev::AllAOIntegralsIterator_2, 79
main as 000	AllAOIntegralsIterator_2, 79, 80
main.cc, 232 make_mdh_D1_coeff	index, 80
	oepdev::AllAOIntegralsIterator_4, 80
The Integral Package Library, 54 make_mdh_D2_coeff	AllAOIntegralsIterator_4, 81, 82
	index, 82
The Integral Package Library, 55 make_mdh_D2_coeff_explicit_recursion	oepdev::AllAOShellCombinationsIterator_2, 82
The Integral Package Library, 55	AllAOShellCombinationsIterator_2, 83, 84
make_mdh_D3_coeff	compute_shell, 85
The Integral Package Library, 56	oepdev::AllAOShellCombinationsIterator_4, 85
make_mdh_R_coeff	AllAOShellCombinationsIterator_4, 86, 87
The Integral Package Library, 57	compute_shell, 87
make_oeps3d	oepdev::CAMM, 90
oepdev::OEPotential, 169	oepdev::CPHF, 96
MultipoleConvergence	CPHF, 98
oepdev::DMTPole, 110	oepdev:: Charge Transfer Energy OEP otential,
oepdev::MultipoleConvergence, 160	91
copacttransporcectiveligenes, 100	oepdev::ChargeTransferEnergySolver, 92
OEPDevSolver	compute_benchmark, 95
oepdev::OEPDevSolver, 164	compute_oep_based, 95
OEPotential	oepdev::CubePoints3DIterator, 99
oepdev::OEPotential, 168	oepdev::CubePointsCollection3D, 100
OEPotential3D	oepdev::DIISManager, 101
The Three-Dimensional Vector Fields Li-	compute, 102
brary, 60	DIISManager, 101
oepdev, 69	put, 102
psi, 75	update, 102
oepdev/lib3d/dmtp.h, 232	oepdev::DMTPole, 103
oepdev/lib3d/esp.h, 233	build, 107
oepdev/libgefp/gefp.h, 234	DMTPole, 107
oepdev/libints/eri.h, 235	energy, 108
oepdev/libints/recurr.h, 236	MultipoleConvergence, 110
oepdev/liboep/oep.h, 237	potential, 108
oepdev/liboep/oep_gdf.h, 238	recenter, 109
oepdev/libpsi/integral.h, 239	oepdev::DoubleGeneralizedDensityFit, 110
oepdev/libpsi/potential.h, 239	compute, 112
oepdev/libsolver/solver.h, 240	oepdev::EETCouplingOEPotential, 113
oepdev/libtest/test.h, 240	oepdev::ERI_1_1, 119
oepdev/libutil/diis.h, 241	oepdev::ERI_2_2, 121
oepdev/libutil/integrals_iter.h, 242	oepdev::ERI_3_1, 123
oepdev/libutil/scf_perturb.h, 243	oepdev::ESPSolver, 124
oepdev/libutil/unitary_optimizer.h, 243	ESPSolver, 126, 127

INDEX 251

oepdev::ElectrostaticEnergyOEPotential, 114	make_oeps3d, 169
oepdev::ElectrostaticEnergySolver, 115	OEPotential, 168
compute_benchmark, 117	oepdev::OEPotential3D< T >, 170
compute_oep_based, 118	oepdev::PerturbCharges, 172
oepdev::ElectrostaticPotential3D, 118	oepdev::Points3Dlterator, 173
oepdev::FFAbInitioPolarGEFactory, 127	build, 175, 176
oepdev::Field3D, 128	Points3DIterator, 174
build, 131	oepdev::Points3Dlterator::Point, 172
Field3D, 131	oepdev::PointsCollection3D, 176
oepdev::Fourier9, 132	build, 178, 179
oepdev::GenEffFrag, 133	PointsCollection3D, 178
susceptibility, 134, 135	oepdev::PolarGEFactory, 179
oepdev::GenEffPar, 136	oepdev::PotentialInt, 181
allocate, 138	PotentialInt, 181, 182
compute_density_matrix, 139, 140	set_charge_field, 183
set_susceptibility, 140	oepdev::QuadraticGradientNonUniformEFieldPolarGEFactory
susceptibility, 140, 141	183
oepdev::GenEffParFactory, 142	oepdev::QuadraticNonUniformEFieldPolarGEFactory,
build, 144	185
oepdev::GeneralizedDensityFit, 145	oepdev::QuadraticUniformEFieldPolarGEFactory,
build, 147	186
compute, 148	oepdev::RHFPerturbed, 196
oepdev::GeneralizedPolarGEFactory, 148	oepdev::RandomPoints3DIterator, 187
oepdev::GeneralizedPolarGEFactory::Statistical	•
203	oepdev::RepulsionEnergyOEPotential, 189
	•
oepdev::IntegralFactory, 154	oepdev::RepulsionEnergySolver, 190
oepdev::LinearGradientNonUniformEFieldPolar	•
155	compute_oep_based, 195
oepdev::LinearNonUniformEFieldPolarGEFactor	
156	ao_iterator, 200
oepdev::LinearUniformEFieldPolarGEFactory,	build, 201
157	compute_shell, 201
oepdev::MultipoleConvergence, 158	ShellCombinationsIterator, 200
compute, 161	oepdev::SingleGeneralizedDensityFit, 202
ConvergenceLevel, 160	compute, 203
level, 161	oepdev::TwoBodyAOInt, 206
MultipoleConvergence, 160	compute, 206, 207
Property, 160	oepdev::TwoElectronInt, 207
oepdev::NonUniformEFieldPolarGEFactory,	compute_shell, 209
162	oepdev::UniformEFieldPolarGEFactory, 209
oepdev::OEPDevSolver, 162	oepdev::UnitaryOptimizer, 210
build, 164	UnitaryOptimizer, 215, 216
compute_benchmark, 165	oepdev::UnitaryOptimizer_4_2, 216
compute_oep_based, 165	UnitaryOptimizer_4_2, 221
OEPDevSolver, 164	oepdev:: Unitary Transformed MOP olar GEF actory,
oepdev::OEPType, 171	222
oepdev::OEPotential, 165	oepdev::WavefunctionUnion, 223
build, 169	Ca_subset, 229

252 INDEX

Cb_subset, 229 WavefunctionUnion, 228 oepdev::test::Test, 204	The OEPDev Testing Platform Library, 68 The OEPDev Utilities, 63 average_moment, 65
Points3DIterator oepdev::Points3DIterator, 174 PointsCollection3D oepdev::PointsCollection3D, 178 potential oepdev::DMTPole, 108 PotentialInt oepdev::PotentialInt, 181, 182 Property oepdev::MultipoleConvergence, 160 psi, 74 oepdev, 75	create_superfunctional, 65 extract_monomer, 66 solve_scf, 66 The Three-Dimensional Vector Fields Library 59 OEPotential3D, 60 UnitaryOptimizer oepdev::UnitaryOptimizer, 215, 216 UnitaryOptimizer_4_2 oepdev::UnitaryOptimizer_4_2, 221 update oepdev::DIISManager, 102
read_options, 75 put oepdev::DIISManager, 102	WavefunctionUnion oepdev::WavefunctionUnion, 228
read_options     psi, 75 recenter     oepdev::DMTPole, 109  set_charge_field     oepdev::PotentialInt, 183 set_susceptibility     oepdev::GenEffPar, 140  ShellCombinationsIterator     oepdev::ShellCombinationsIterator, 200 solve_scf     The OEPDev Utilities, 66	
susceptibility oepdev::GenEffFrag, 134, 135 oepdev::GenEffPar, 140, 141	
The Density Functional Theory Library, 62 The Generalized Effective Fragment Potentials Library, 47 The Generalized One-Electron Potentials Library, 45 The Integral Package Library, 49 d_N_n1_n2, 54 make_mdh_D1_coeff, 54 make_mdh_D2_coeff, 55 make_mdh_D2_coeff, 55 make_mdh_D3_coeff, 56 make_mdh_R_coeff, 57	
The OEPDev Solver Library, 46	