Ecosystem

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ii CONTENTS

Contents

1	Intro	duction	2
	1.1	Copyright Statement	2
	1.2	General Information	2
2	Hiera	archical Index	2
	2.1	Class Hierarchy	2
3	Clas	s Index	4
	3.1	Class List	4
4	File	Index	7
	4.1	File List	7
5	Clas	s Documentation	8
	5.1	AdsorptionReaction Class Reference	8
		5.1.1 Detailed Description	12
		5.1.2 Constructor & Destructor Documentation	12
		5.1.3 Member Function Documentation	12
		5.1.4 Member Data Documentation	19
	5.2	ARNOLDI_DATA Struct Reference	21
		5.2.1 Detailed Description	22
		5.2.2 Member Data Documentation	22
	5.3	Atom Class Reference	23
		5.3.1 Detailed Description	25
		5.3.2 Constructor & Destructor Documentation	26
		5.3.3 Member Function Documentation	26
		5.3.4 Member Data Documentation	28
	5.4	BACKTRACK_DATA Struct Reference	29
		5.4.1 Detailed Description	30
		5.4.2 Member Data Documentation	30
	5.5	BiCGSTAB_DATA Struct Reference	31

	5.5.1	Detailed Description	32
	5.5.2	Member Data Documentation	32
5.6	CGS_E	DATA Struct Reference	34
	5.6.1	Detailed Description	36
	5.6.2	Member Data Documentation	36
5.7	Chemis	sorptionReaction Class Reference	38
	5.7.1	Detailed Description	41
	5.7.2	Constructor & Destructor Documentation	41
	5.7.3	Member Function Documentation	41
	5.7.4	Member Data Documentation	47
5.8	Docum	ent Class Reference	47
	5.8.1	Detailed Description	49
	5.8.2	Constructor & Destructor Documentation	50
	5.8.3	Member Function Documentation	50
	5.8.4	Member Data Documentation	53
5.9	DOGFI	SH_DATA Struct Reference	53
	5.9.1	Detailed Description	55
	5.9.2	Member Data Documentation	55
5.10	DOGFI	ISH_PARAM Struct Reference	57
	5.10.1	Detailed Description	57
	5.10.2	Member Data Documentation	57
5.11	FINCH	_DATA Struct Reference	58
	5.11.1	Detailed Description	62
	5.11.2	Member Data Documentation	62
5.12	GCR_E	DATA Struct Reference	71
	5.12.1	Detailed Description	72
	5.12.2	Member Data Documentation	72
5.13	GMRE	SLP_DATA Struct Reference	74
	5.13.1	Detailed Description	75
	5.13.2	Member Data Documentation	75

iv CONTENTS

5.14	GMRESR_DATA Struct Reference	76
	5.14.1 Detailed Description	77
	5.14.2 Member Data Documentation	78
5.15	GMRESRP_DATA Struct Reference	79
	5.15.1 Detailed Description	81
	5.15.2 Member Data Documentation	81
5.16	GPAST_DATA Struct Reference	83
	5.16.1 Detailed Description	84
	5.16.2 Member Data Documentation	84
5.17	GSTA_DATA Struct Reference	85
	5.17.1 Detailed Description	85
	5.17.2 Member Data Documentation	85
5.18	GSTA_OPT_DATA Struct Reference	86
	5.18.1 Detailed Description	86
	5.18.2 Member Data Documentation	87
5.19	Header Class Reference	88
	5.19.1 Detailed Description	90
	5.19.2 Constructor & Destructor Documentation	90
	5.19.3 Member Function Documentation	90
	5.19.4 Member Data Documentation	93
5.20	KeyValueMap Class Reference	93
	5.20.1 Detailed Description	95
	5.20.2 Constructor & Destructor Documentation	95
	5.20.3 Member Function Documentation	96
	5.20.4 Member Data Documentation	98
5.21	KMS_DATA Struct Reference	98
	5.21.1 Detailed Description	99
	5.21.2 Member Data Documentation	99
5.22	MAGPIE_DATA Struct Reference	101
	5.22.1 Detailed Description	101

	5.22.2 Member Data Documentation	101
5.23	MassBalance Class Reference	102
	5.23.1 Detailed Description	104
	5.23.2 Constructor & Destructor Documentation	104
	5.23.3 Member Function Documentation	104
	5.23.4 Member Data Documentation	107
5.24	MasterSpeciesList Class Reference	108
	5.24.1 Detailed Description	109
	5.24.2 Constructor & Destructor Documentation	109
	5.24.3 Member Function Documentation	110
	5.24.4 Member Data Documentation	112
5.25	Matrix < T > Class Template Reference	112
	5.25.1 Detailed Description	115
	5.25.2 Constructor & Destructor Documentation	115
	5.25.3 Member Function Documentation	115
	5.25.4 Member Data Documentation	121
5.26	MIXED_GAS Struct Reference	121
	5.26.1 Detailed Description	122
	5.26.2 Member Data Documentation	122
5.27	Molecule Class Reference	124
	5.27.1 Detailed Description	126
	5.27.2 Constructor & Destructor Documentation	126
	5.27.3 Member Function Documentation	127
	5.27.4 Member Data Documentation	130
5.28	MONKFISH_DATA Struct Reference	131
	5.28.1 Detailed Description	133
	5.28.2 Member Data Documentation	133
5.29	MONKFISH_PARAM Struct Reference	136
	5.29.1 Detailed Description	137
	5.29.2 Member Data Documentation	137

vi CONTENTS

5.30	mSPD_DATA Struct Reference	138
	5.30.1 Detailed Description	139
	5.30.2 Member Data Documentation	139
5.31	MultiligandAdsorption Class Reference	139
	5.31.1 Detailed Description	142
	5.31.2 Constructor & Destructor Documentation	142
	5.31.3 Member Function Documentation	142
	5.31.4 Member Data Documentation	147
5.32	MultiligandChemisorption Class Reference	149
	5.32.1 Detailed Description	151
	5.32.2 Constructor & Destructor Documentation	152
	5.32.3 Member Function Documentation	152
	5.32.4 Member Data Documentation	157
5.33	NUM_JAC_DATA Struct Reference	158
	5.33.1 Detailed Description	159
	5.33.2 Member Data Documentation	159
5.34	OPTRANS_DATA Struct Reference	159
	5.34.1 Detailed Description	159
	5.34.2 Member Data Documentation	160
5.35	PCG_DATA Struct Reference	160
	5.35.1 Detailed Description	161
	5.35.2 Member Data Documentation	161
5.36	PeriodicTable Class Reference	163
	5.36.1 Detailed Description	163
	5.36.2 Constructor & Destructor Documentation	163
	5.36.3 Member Function Documentation	164
	5.36.4 Member Data Documentation	164
5.37	PICARD_DATA Struct Reference	164
	5.37.1 Detailed Description	165
	5.37.2 Member Data Documentation	165

CONTENTS vii

5.38	PJFNK_DATA Struct Reference	166
	5.38.1 Detailed Description	168
	5.38.2 Member Data Documentation	168
5.39	PURE_GAS Struct Reference	171
	5.39.1 Detailed Description	172
	5.39.2 Member Data Documentation	172
5.40	QR_DATA Struct Reference	173
	5.40.1 Detailed Description	173
	5.40.2 Member Data Documentation	174
5.41	Reaction Class Reference	174
	5.41.1 Detailed Description	176
	5.41.2 Constructor & Destructor Documentation	176
	5.41.3 Member Function Documentation	176
	5.41.4 Member Data Documentation	178
5.42	SCOPSOWL_DATA Struct Reference	179
	5.42.1 Detailed Description	181
	5.42.2 Member Data Documentation	181
5.43	SCOPSOWL_OPT_DATA Struct Reference	184
	5.43.1 Detailed Description	186
	5.43.2 Member Data Documentation	186
5.44	SCOPSOWL_PARAM_DATA Struct Reference	189
	5.44.1 Detailed Description	190
	5.44.2 Member Data Documentation	190
5.45	SHARK_DATA Struct Reference	192
	5.45.1 Detailed Description	195
	5.45.2 Member Data Documentation	196
5.46	SKUA_DATA Struct Reference	204
	5.46.1 Detailed Description	205
	5.46.2 Member Data Documentation	205
5.47	SKUA_OPT_DATA Struct Reference	207

viii CONTENTS

	5.47.1 I	Detailed Description	209
	5.47.2 I	Member Data Documentation	209
5.48	SKUA_F	PARAM Struct Reference	211
	5.48.1 I	Detailed Description	212
	5.48.2 I	Member Data Documentation	212
5.49	SubHea	der Class Reference	213
	5.49.1 I	Detailed Description	214
	5.49.2	Constructor & Destructor Documentation	214
	5.49.3 I	Member Function Documentation	215
	5.49.4 I	Member Data Documentation	216
5.50	SYSTEM	M_DATA Struct Reference	217
	5.50.1 I	Detailed Description	218
	5.50.2	Member Data Documentation	218
5.51	TRAJEC	CTORY_DATA Struct Reference	219
	5.51.1	Member Data Documentation	221
5.52	UI_DATA	A Struct Reference	223
	5.52.1 I	Detailed Description	223
	5.52.2	Member Data Documentation	223
5.53	Unstead	lyAdsorption Class Reference	225
	5.53.1 I	Detailed Description	228
	5.53.2	Constructor & Destructor Documentation	228
	5.53.3 I	Member Function Documentation	228
	5.53.4 I	Member Data Documentation	235
5.54	Unstead	lyReaction Class Reference	236
	5.54.1 I	Detailed Description	239
	5.54.2	Constructor & Destructor Documentation	239
	5.54.3 I	Member Function Documentation	239
	5.54.4 I	Member Data Documentation	245
5.55	ValueTyp	pePair Class Reference	247
	5.55.1 I	Detailed Description	248
	5.55.2	Constructor & Destructor Documentation	248
	5.55.3 I	Member Function Documentation	248
	5.55.4 I	Member Data Documentation	249
5.56	yaml_cp	p_class Class Reference	250
	5.56.1 I	Detailed Description	250
	5.56.2	Constructor & Destructor Documentation	251
	5.56.3 I	Member Function Documentation	251
	5.56.4 I	Member Data Documentation	251
5.57	YamlWra	apper Class Reference	252
	5.57.1 I	Detailed Description	253
	5.57.2	Constructor & Destructor Documentation	254
	5.57.3 I	Member Function Documentation	254
	5.57.4 I	Member Data Documentation	256

CONTENTS ix

6	File I	Documentation 25				
	6.1	dogfish.h File Reference				
		6.1.1	Detailed Description	257		
		6.1.2	Function Documentation	258		
	6.2	eel.h F	ille Reference	261		
		6.2.1	Detailed Description	261		
		6.2.2	Function Documentation	262		
	6.3	egret.h	File Reference	262		
		6.3.1	Detailed Description	264		
		6.3.2	Macro Definition Documentation	264		
		6.3.3	Function Documentation	265		
	6.4	error.h	File Reference	266		
		6.4.1	Detailed Description	267		
		6.4.2	Macro Definition Documentation	267		
		6.4.3	Enumeration Type Documentation	267		
		6.4.4	Function Documentation	269		
	6.5	finch.h	File Reference	269		
		6.5.1	Detailed Description	271		
		6.5.2	Enumeration Type Documentation	272		
		6.5.3	Function Documentation	272		
	6.6	flock.h	File Reference	276		
		6.6.1	Detailed Description	276		
	6.7	gsta_o	pt.h File Reference	277		
		6.7.1	Detailed Description	278		
		6.7.2	Macro Definition Documentation	279		
		6.7.3	Function Documentation	279		
	6.8	lark.h I	File Reference	284		
		6.8.1	Detailed Description	286		
		6.8.2	Macro Definition Documentation	288		
		6.8.3	Enumeration Type Documentation	288		

X CONTENTS

	6.8.4	Function Documentation	288
6.9	macaw	v.h File Reference	301
	6.9.1	Detailed Description	302
	6.9.2	Macro Definition Documentation	303
	6.9.3	Function Documentation	303
6.10	magpie	e.h File Reference	303
	6.10.1	Detailed Description	305
	6.10.2	Macro Definition Documentation	305
	6.10.3	Function Documentation	306
6.11	mola.h	File Reference	313
	6.11.1	Detailed Description	314
	6.11.2	Macro Definition Documentation	320
	6.11.3	Enumeration Type Documentation	320
	6.11.4	Function Documentation	321
6.12	monkfis	sh.h File Reference	321
	6.12.1	Detailed Description	322
	6.12.2	Function Documentation	322
6.13	sandbo	ox.h File Reference	325
	6.13.1	Detailed Description	325
	6.13.2	Function Documentation	326
6.14	school.	h File Reference	326
	6.14.1	Detailed Description	326
6.15	scopso	owl.h File Reference	327
	6.15.1	Detailed Description	329
	6.15.2	Macro Definition Documentation	329
	6.15.3	Function Documentation	330
6.16	scopso	owl_opt.h File Reference	338
	6.16.1	Detailed Description	339
	6.16.2	Function Documentation	340
6.17	shark.h	n File Reference	343

	6.17.1	Detailed Description	347
	6.17.2	Macro Definition Documentation	348
	6.17.3	Typedef Documentation	349
	6.17.4	Enumeration Type Documentation	349
	6.17.5	Function Documentation	350
6.18	skua.h	File Reference	367
	6.18.1	Detailed Description	368
	6.18.2	Macro Definition Documentation	369
	6.18.3	Function Documentation	369
6.19	skua_o	ppt.h File Reference	375
	6.19.1	Detailed Description	376
	6.19.2	Function Documentation	377
6.20	Trajecto	ory.h File Reference	380
	6.20.1	Detailed Description	381
	6.20.2	Function Documentation	382
6.21	ui.h File	e Reference	383
	6.21.1	Detailed Description	385
	6.21.2	Macro Definition Documentation	385
	6.21.3	Enumeration Type Documentation	386
	6.21.4	Function Documentation	386
6.22	yaml_w	vrapper.h File Reference	391
	6.22.1	Detailed Description	393
	6.22.2	Typedef Documentation	395
	6.22.3	Enumeration Type Documentation	395
	6.22.4	Function Documentation	396

397

Index

1 Introduction

1.1 Copyright Statement

Copyright

This software was designed and built at the Georgia Institute of Technology by Austin Ladshaw for PhD research in the area of adsorption and surface science. Copyright (c) 2015, all rights reserved.

1.2 General Information

The source code contained within the ecosystem project was designed as a standalone tool set for performing numerical modeling and data analyses associated with adsorption phenomena in both gaseous and aqueous systems. Many of the lower level tools are general enough to be applied to any system you desire to be modeled. Such algorithms included are Krylov subspace methods for linear systems and a Jacobian-Free Newton-Krylov method for non-linear systems. There is also a templated matrix object for generic matrix construction and modification. For specific information on each individual kernel, navigate through the class and file indices or table of contents.

Warning

Many of these algorithms may still be under development. This library is distributed in the hope that it will be useful, but WITHOUT ANY WARRANTY; without even the implied warranty of MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE.

2 Hierarchical Index

2.1 Class Hierarchy

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

AdsorptionReaction	
ChemisorptionReaction	38
UnsteadyAdsorption	225
ARNOLDI_DATA	21
Atom	23
BACKTRACK_DATA	29
BICGSTAB_DATA	31
CGS_DATA	34
DOGFISH_DATA	53
DOGFISH_PARAM	57
FINCH_DATA	58
GCR_DATA	71

ο 4	\sim 1		11:-			
2.1	- Cli	ass	піе	l a	rci	ıv

GMRESLP_DATA	74
GMRESR_DATA	76
GMRESRP_DATA	79
GPAST_DATA	83
GSTA_DATA	85
GSTA_OPT_DATA	86
KeyValueMap	93
KMS_DATA	98
MAGPIE_DATA	101
MassBalance	102
MasterSpeciesList	108
Matrix < T >	112
Matrix< double >	112
Matrix< int >	112
MIXED_GAS	121
Molecule	124
MONKFISH_DATA	131
MONKFISH_PARAM	136
mSPD_DATA	138
MultiligandAdsorption	139
MultiligandChemisorption	149
NUM_JAC_DATA	158
OPTRANS_DATA	159
PCG_DATA	160
PeriodicTable	163
PICARD_DATA	164
PJFNK_DATA	166
PURE_GAS	171
QR_DATA	173
Reaction	174
UnsteadyReaction	236
SCOPSOWL_DATA	179

	SCOPSOWL_OPT_DATA	184
	SCOPSOWL_PARAM_DATA	189
	SHARK_DATA	192
	SKUA_DATA	204
	SKUA_OPT_DATA	207
	SKUA_PARAM	211
	SubHeader	213
	Document	47
	Header	88
	SYSTEM_DATA	217
	TRAJECTORY_DATA	219
	UI_DATA	223
	ValueTypePair	247
	yaml_cpp_class	250
	YamlWrapper	252
3 3.1	Class Index Class List	
He	re are the classes, structs, unions and interfaces with brief descriptions:	
	AdsorptionReaction Adsorption Reaction Object	8
	ARNOLDI_DATA Data structure for the construction of the Krylov subspaces for a linear system	21
	Atom Atom object to hold information about specific atoms in the periodic table (click Atom to go to function definitions)	23
	BACKTRACK_DATA Data structure for the implementation of Backtracking Linesearch	29
	BiCGSTAB_DATA Data structure for the implementation of the BiCGSTAB algorithm for non-symmetric linear systems	31
	CGS_DATA Data structure for the implementation of the CGS algorithm for non-symmetric linear systems	34
	ChemisorptionReaction Chemisorption Reaction Object	38

3.1 Class List 5

Document Object for the various documents in the yaml file	47
DOGFISH_DATA Primary data structure for running the DOGFISH application	53
DOGFISH_PARAM Data structure for species-specific parameters	57
FINCH_DATA Data structure for the FINCH object	58
GCR_DATA Data structure for the implementation of the GCR algorithm for non-symmetric linear systems	71
GMRESLP_DATA Data structure for implementation of the Restarted GMRES algorithm with Left Preconditioning	74
GMRESR_DATA Data structure for the implementation of GCR with Nested GMRES preconditioning (i.e., GM↔ RESR)	76
GMRESRP_DATA Data structure for the Restarted GMRES algorithm with Right Preconditioning	79
GPAST_DATA GPAST Data Structure	83
GSTA_DATA GSTA Data Structure	85
GSTA_OPT_DATA Data structure used in the GSTA optimization routines	86
Header Object for headers in a yaml document (inherits from SubHeader)	88
KeyValueMap Key-Value-Type Map object creating a map of the KeyValuePair objects	93
KMS_DATA Data structure for the implemenation of the Krylov Multi-Space (KMS) Method	98
MAGPIE_DATA MAGPIE Data Structure	101
MassBalance Mass Balance Object	102
MasterSpeciesList Master Species List Object	108
Matrix < T > Templated C++ Matrix Class Object (click Matrix to go to function definitions)	112
MIXED_GAS Data structure holding information necessary for computing mixed gas properties	121
Molecule C++ Molecule Object built from Atom Objects (click Molecule to go to function definitions)	124

MONKFISH_DATA Primary data structure for running MONKFISH	131
MONKFISH_PARAM Data structure for species specific information and parameters	136
mSPD_DATA MSPD Data Structure	138
MultiligandAdsorption Multi-ligand Adsorption Reaction Object	139
MultiligandChemisorption Multi-ligand Chemisorption Reaction Object	149
NUM_JAC_DATA Data structure to form a numerical jacobian matrix with finite differences	158
OPTRANS_DATA Data structure for implementation of linear operator transposition	159
PCG_DATA Data structure for implementation of the PCG algorithms for symmetric linear systems	160
PeriodicTable Class object that store a digitial copy of all Atom objects	163
PICARD_DATA Data structure for the implementation of a Picard or Fixed-Point iteration for non-linear systems	164
PJFNK_DATA Data structure for the implementation of the PJFNK algorithm for non-linear systems	166
PURE_GAS Data structure holding all the parameters for each pure gas spieces	171
QR_DATA Data structure for the implementation of a QR solver given some invertable linear operator	173
Reaction Reaction Object	174
SCOPSOWL_DATA Primary data structure for SCOPSOWL simulations	179
SCOPSOWL_OPT_DATA Data structure for the SCOPSOWL optmization routine	184
SCOPSOWL_PARAM_DATA Data structure for the species' parameters in SCOPSOWL	189
SHARK_DATA Data structure for SHARK simulations	192
SKUA_DATA Data structure for all simulation information in SKUA	204
SKUA_OPT_DATA Data structure for the SKUA Optimization Routine	207
SKUA_PARAM Data structure for species' parameters in SKUA	211

4 File Index 7

SubHeader Object for the Lowest level of Header for the yaml_wrapper	213
SYSTEM_DATA System Data Structure	217
TRAJECTORY_DATA	219
UI_DATA Data structure holding the UI arguments	223
UnsteadyAdsorption Unsteady Adsorption Reaction Object	225
UnsteadyReaction Unsteady Reaction Object (inherits from Reaction)	236
ValueTypePair Value-Type Pair object to recognize data type of a string that was read	247
yaml_cpp_class Primary object used when reading and digitally storing yaml files	250
YamlWrapper Object for the entire yaml file holding all documents, header, sub-headers, keys, an	nd values 252
4 File Index	
4.1 File List	
Here is a list of all files with brief descriptions:	
dogfish.h Diffusion Object Governing Fiber Interior Sorption History	256
eel.h Easy-access Element Library	26 1
egret.h Estimation of Gas-phase pRopErTies	262
error.h All error types are defined here	266
finch.h Flux-limiting Implicit Non-oscillatory Conservative High-resolution scheme	269
flock.h FundamentaL Off-gas Collection of Kernels	276
gsta_opt.h Generalized Statistical Thermodynamic Adsorption (GSTA) Optimization Routine	277
lark.h Linear Algebra Residual Kernels	284
macaw.h MAtrix CAlculation Workspace	30 ⁻

magpie.h Multicomponent Adsorption Generalized Procedure for Isothermal Equilibria	303
mola.h Molecule Object Library from Atoms	313
monkfish.h Multi-fiber wOven Nest Kernel For Interparticle Sorption History	321
sandbox.h Coding Test Area	325
school.h Seawater Codes from a Highly Object-Oriented Library	326
scopsowl.h Simultaneously Coupled Objects for Pore and Surface diffusion Operations With Linear systems	327
scopsowl_opt.h Optimization Routine for Surface Diffusivities in SCOPSOWL	338
shark.h Speciation-object Hierarchy for Adsorption Reactions and Kinetics	343
skua.h Surface Kinetics for Uptake by Adsorption	367
skua_opt.h Optimization Routine for the SKUA Model	375
Trajectory.h Single Particle Trajectory Analysis for Magnetic Filtration	380
ui.h User Interface for Ecosystem	383
yaml_wrapper.h C++ Wrapper for the C-YAML Library	391

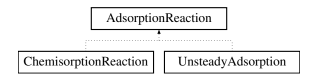
5 Class Documentation

5.1 AdsorptionReaction Class Reference

Adsorption Reaction Object.

#include <shark.h>

Inheritance diagram for AdsorptionReaction:



Public Member Functions

· AdsorptionReaction ()

Default Constructor.

∼AdsorptionReaction ()

Default Destructor.

• void Initialize_Object (MasterSpeciesList &List, int n)

Function to call the initialization of objects sequentially.

· void Display_Info ()

Display the adsorption reaction information (PLACE HOLDER)

void modifyDeltas (MassBalance &mbo)

Modify the Deltas in the MassBalance Object.

• int setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object.

· int checkAqueousIndices ()

Function to check and report errors in the aqueous species indices.

 void setActivityModelInfo (int(*act)(const Matrix< double > &logq, Matrix< double > &activity, const void *data), const void *act_data)

Function to set the surface activity model and data pointer.

void setAqueousIndex (int rxn i, int species i)

Set the primary aqueous species index for the ith reaction.

int setAqueousIndexAuto ()

Automatically sets the primary aqueous species index based on reactions.

· void setActivityEnum (int act)

Set the surface activity enum value.

void setMolarFactor (int rxn_i, double m)

Set the molar factor for the ith reaction (mol/mol)

void setVolumeFactor (int i, double v)

Set the ith volume factor for the species list (cm[^]3/mol)

void setAreaFactor (int i, double a)

Set the ith area factor for the species list (m^2/mol)

void setSpecificArea (double a)

Set the specific area for the adsorbent (m^2/kg)

void setSpecificMolality (double a)

Set the specific molality for the adsorbent (mol/kg)

void setSurfaceCharge (double c)

Set the surface charge of the uncomplexed ligands.

void setTotalMass (double m)

Set the total mass of the adsorbent (kg)

void setTotalVolume (double v)

Set the total volume of the system (L)

void setAreaBasisBool (bool opt)

Set the basis boolean directly.

void setSurfaceChargeBool (bool opt)

Set the boolean for inclusion of surface charging.

void setBasis (std::string option)

Set the basis of the adsorption problem from the given string arg.

void setAdsorbentName (std::string name)

Set the name of the adsorbent to the given string.

• void setChargeDensityValue (double a)

Set the value of the charge density parameter to a (C/m^2)

void setlonicStrengthValue (double a)

Set the value of the ionic strength parameter to a (mol/L)

void setActivities (Matrix< double > &x)

Set the values of activities in the activity matrix.

void calculateAreaFactors ()

Calculates the area factors used from the van der Waals volumes.

void calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

void setChargeDensity (const Matrix < double > &x)

Calculates and sets the current value of charge density.

void setlonicStrength (const Matrix< double > &x)

Calculates and sets the current value of ionic strength.

int callSurfaceActivity (const Matrix < double > &x)

Calls the activity model and returns an int flag for success or failure.

double calculateActiveFraction (const Matrix< double > &x)

Calculates the fraction of the surface that is active and available.

double calculateSurfaceChargeDensity (const Matrix< double > &x)

Function to calculate the surface charge density based on concentrations.

double calculateLangmuirMaxCapacity (int i)

Calculates the theoretical maximum capacity for adsorption in reaction i.

double calculateLangmuirEquParam (const Matrix < double > &x, const Matrix < double > &gama, int i)

Calculates the equivalent Langmuir isotherm equilibrium parameter.

double calculateLangmuirAdsorption (const Matrix< double > &x, const Matrix< double > &gama, int i)

Calculates the equivalent Langmuir adsorption by forming the Langmuir-like parameters.

• double calculatePsi (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

double calculateAqueousChargeExchange (int i)

Function to calculate the net exchange of charges of the ageous species involved in a given reaction.

• double calculateEquilibriumCorrection (double sigma, double T, double I, double rel_epsilon, int i)

Function to calculate the correction term for the equilibrium parameter.

double Eval_Residual (const Matrix< double > &x, const Matrix< double > &gama, double T, double rel_
 perm, int i)

Calculates the residual for the ith reaction in the system.

· Reaction & getReaction (int i)

Return reference to the ith reaction object in the adsorption object.

double getMolarFactor (int i)

Get the ith reaction's molar factor for adsorption (mol/mol)

double getVolumeFactor (int i)

Get the ith volume factor (species not involved return zeros) (cm^3/mol)

double getAreaFactor (int i)

Get the ith area factor (species not involved return zeros) (m²/mol)

double getActivity (int i)

Get the ith activity factor for the surface species.

• double getSpecificArea ()

Get the specific area of the adsorbent (m^2/kg) or (mol/kg)

double getSpecificMolality ()

Get the specific molality of the adsorbent (mol/kg)

double getSurfaceCharge ()

Get the surface charge of the adsorbent.

• double getBulkDensity ()

Calculate and return bulk density of adsorbent in system (kg/L)

• double getTotalMass ()

Get the total mass of adsorbent in the system (kg)

double getTotalVolume ()

Get the total volume of the system (L)

double getChargeDensity ()

Get the value of the surface charge density (C/m^2)

• double getlonicStrength ()

Get the value of the ionic strength of solution (mol/L)

• int getNumberRxns ()

Get the number of reactions involved in the adsorption object.

• int getAdsorbIndex (int i)

Get the index of the adsorbed species in the ith reaction.

int getAqueousIndex (int i)

Get the index of the primary aqueous species in the ith reaction.

int getActivityEnum ()

Return the enum representing the choosen activity function.

· bool isAreaBasis ()

Returns true if we are in the Area Basis. False if in Molar Basis.

bool includeSurfaceCharge ()

Returns true if we are considering surface charging during adsorption.

std::string getAdsorbentName ()

Returns the name of the adsorbent as a string.

Protected Attributes

MasterSpeciesList * List

Pointer to the MasterSpeciesList object.

int(* surface_activity)(const Matrix< double > &logq, Matrix< double > &activity, const void *data)

Pointer to a surface activity model.

const void * activity_data

Pointer to the data structure needed for surface activities.

int act_fun

Enumeration of the activity function being used for the surface phase.

std::vector< double > area_factors

List of the van der Waals areas associated with surface species (m[^]2/mol)

• std::vector< double > volume_factors

List of the van der Waals volumes of each surface species (cm[^]3/mol)

std::vector< int > adsorb_index

List of the indices for the adsorbed species in the reactions.

• std::vector< int > aqueous_index

List of the indices for the primary aqueous species in the reactions.

std::vector< double > molar_factor

List of the number of ligands needed to form one mole of adsorption in each reaction.

• Matrix< double > activities

List of the activities calculated by the activity model.

double specific_area

Specific surface area of the adsorbent (m^2/kg)

· double specific molality

Specific molality of the adsorbent - moles of ligand per kg sorbent (mol/kg)

double surface_charge

Charge of the uncomplexed surface ligand species.

· double total mass

Total mass of the adsorbent in the system (kg)

· double total volume

Total volume of the system (L)

· double ionic_strength

Ionic Strength of the system used to adjust equilibria constants (mol/L)

· double charge_density

Surface charge density of the adsorbent used to adjust equilbria (C/m²)

· int num rxns

Number of reactions involved in the adsorption equilibria.

· bool AreaBasis

True = Adsorption on an area basis, False = Adsorption on a ligand basis.

bool IncludeSurfCharge

True = Includes surface charging corrections, False = Does not consider surface charge.

std::string adsorbent_name

Name of the adsorbent for this object.

Private Attributes

std::vector< Reaction > ads rxn

List of reactions involved with adsorption.

5.1.1 Detailed Description

Adsorption Reaction Object.

C++ Object to handle data and functions associated with forumlating adsorption equilibrium reactions in a aqueous mixture. Each unique surface in a system will require an instance of this structure.

5.1.2 Constructor & Destructor Documentation

5.1.2.1 AdsorptionReaction::AdsorptionReaction ()

Default Constructor.

5.1.2.2 AdsorptionReaction:: \sim AdsorptionReaction ()

Default Destructor.

5.1.3 Member Function Documentation

5.1.3.1 void AdsorptionReaction::Initialize_Object (MasterSpeciesList & List, int n)

Function to call the initialization of objects sequentially.

5.1.3.2 void AdsorptionReaction::Display_Info()

Display the adsorption reaction information (PLACE HOLDER)

5.1.3.3 void AdsorptionReaction::modifyDeltas (MassBalance & mbo)

Modify the Deltas in the MassBalance Object.

This function will take a mass balance object as an argument and modify the deltas in that object to correct for how adsorption affects that particular mass balance. Since adsorption can effect multiple mass balances, this function must be called for each mass balance in the system.

Parameters

mbo reference to the MassBalance Object the adsorption is acting on

5.1.3.4 int AdsorptionReaction::setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object.

This function searches through the Reaction objects in AdsorptionReaction to find the solid species and their indices to set that information in the adsorb_index structure. That information will be used later to approximate maximum capacities and equilibrium parameters for use in a modified extended Langmuir type expression. Function will return 0 if successful and -1 on a failure.

5.1.3.5 int AdsorptionReaction::checkAqueousIndices ()

Function to check and report errors in the aqueous species indices.

5.1.3.6 void AdsorptionReaction::setActivityModelInfo (int(*)(const Matrix< double > &logq, Matrix< double > &activity, const void *data) act, const void * act_data)

Function to set the surface activity model and data pointer.

This function will setup the surface activity model based on the given pointer arguments. If no arguments are given, or are given as NULL, then the activity model will default to ideal solution assumption.

5.1.3.7 void AdsorptionReaction::setAqueousIndex (int rxn_i, int species_i)

Set the primary aqueous species index for the ith reaction.

5.1.3.8 int AdsorptionReaction::setAqueousIndexAuto ()

Automatically sets the primary aqueous species index based on reactions.

This function will go through all species and all reactions in the adsorption object and automatically set the primary aqueous species index based on the stoicheometry of the reaction. It will also check and make sure that the primary aqueous index species appears opposite of the adsorbed species in the reactions. Note: This function assumes that the adsorbed indices have already been set.

5.1.3.9 void AdsorptionReaction::setActivityEnum (int act)

Set the surface activity enum value.

5.1.3.10 void AdsorptionReaction::setMolarFactor (int rxn_i , double m)

Set the molar factor for the ith reaction (mol/mol)

5.1.3.11 void AdsorptionReaction::setVolumeFactor (int i, double v)

Set the ith volume factor for the species list (cm³/mol)

```
5.1.3.12 void AdsorptionReaction::setAreaFactor (int i, double a)
Set the ith area factor for the species list (m<sup>2</sup>/mol)
5.1.3.13 void AdsorptionReaction::setSpecificArea ( double a )
Set the specific area for the adsorbent (m<sup>2</sup>/kg)
5.1.3.14 void AdsorptionReaction::setSpecificMolality ( double a )
Set the specific molality for the adsorbent (mol/kg)
5.1.3.15 void AdsorptionReaction::setSurfaceCharge ( double c )
Set the surface charge of the uncomplexed ligands.
5.1.3.16 void AdsorptionReaction::setTotalMass ( double m )
Set the total mass of the adsorbent (kg)
5.1.3.17 void AdsorptionReaction::setTotalVolume ( double v )
Set the total volume of the system (L)
5.1.3.18 void AdsorptionReaction::setAreaBasisBool (bool opt)
Set the basis boolean directly.
5.1.3.19 void AdsorptionReaction::setSurfaceChargeBool ( bool opt )
Set the boolean for inclusion of surface charging.
5.1.3.20 void AdsorptionReaction::setBasis ( std::string option )
Set the basis of the adsorption problem from the given string arg.
5.1.3.21 void AdsorptionReaction::setAdsorbentName ( std::string name )
Set the name of the adsorbent to the given string.
5.1.3.22 void AdsorptionReaction::setChargeDensityValue ( double a )
Set the value of the charge density parameter to a (C/m<sup>2</sup>)
5.1.3.23 void AdsorptionReaction::setlonicStrengthValue ( double a )
Set the value of the ionic strength parameter to a (mol/L)
```

5.1.3.24 void AdsorptionReaction::setActivities (Matrix < double > & x)

Set the values of activities in the activity matrix.

5.1.3.25 void AdsorptionReaction::calculateAreaFactors ()

Calculates the area factors used from the van der Waals volumes.

5.1.3.26 void AdsorptionReaction::calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

5.1.3.27 void AdsorptionReaction::setChargeDensity (const Matrix < double > & x)

Calculates and sets the current value of charge density.

5.1.3.28 void AdsorptionReaction::setlonicStrength (const Matrix < double > & x)

Calculates and sets the current value of ionic strength.

5.1.3.29 int AdsorptionReaction::callSurfaceActivity (const Matrix < double > & x)

Calls the activity model and returns an int flag for success or failure.

5.1.3.30 double AdsorptionReaction::calculateActiveFraction (const Matrix< double > & x)

Calculates the fraction of the surface that is active and available.

5.1.3.31 double AdsorptionReaction::calculateSurfaceChargeDensity (const Matrix < double > & x)

Function to calculate the surface charge density based on concentrations.

This function is used to calculate the surface charge density of the adsorbed species based on the charges and concentrations of the adsorbed species. The calculation is used to correct the adsorption equilibria constant based on a localized surface charge balance. This requires that you know the molality of the uncomplexed ligand species on the surface, as well as the specific surface area for the adsorbent.

Parameters

x matrix of the log(C) concentration values at the current non-linear step

5.1.3.32 double AdsorptionReaction::calculateLangmuirMaxCapacity (int i)

Calculates the theoretical maximum capacity for adsorption in reaction i.

This function is used to calculate the current maximum capacity of a species for a given adsorption reaction using the concentrations and activities of other species in the system. You must pass the index of the reaction of interest. The index of the species of interest is determined from the adsorb_index object. Note: This is only true if the stoicheometry for the adsorbed species is 1.

Parameters

i index of the reaction of interest for the adsorption object

5.1.3.33 double AdsorptionReaction::calculateLangmuirEquParam (const Matrix< double > & x, const Matrix< double > & gama, int i)

Calculates the equivalent Langmuir isotherm equilibrium parameter.

This function will take in the current aqueous activities and calculate an effective Langmuir adsorption parameter for use in determining the adsorption in the system. It uses the system temperature as well to calculate equilibrium. Note: This is only true if the stoicheometry for the adsorbed species is 1.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
i	index of the reaction of interest for the adsorption object

5.1.3.34 double AdsorptionReaction::calculateLangmuirAdsorption (const Matrix< double > & x, const Matrix< double > & gama, int i)

Calculates the equivalent Langmuir adsorption by forming the Langmuir-like parameters.

This function will use the calculateLangmuirMaxCapacity and calculateLangmuirEquParam functions to approximate the adsorption of the ith reaction given the concentration of aqueous species, activities, and temperature. Note: This is only true if the stoicheometry for the adsorbed species is 1.

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
i	index of the reaction of interest for the adsorption object

5.1.3.35 double AdsorptionReaction::calculatePsi (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

This function will calculate the electric surface potential of the adsorbent under the current conditions of charge density, temperature, ionic strength, and relative permittivity.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)

5.1.3.36 double AdsorptionReaction::calculateAqueousChargeExchange (int i)

Function to calculate the net exchange of charges of the ageous species involved in a given reaction.

This function will look at all aqueous species involved in the ith adsorption reaction and sum up their stoicheometries and charges to see what the net change in charge is caused by the adsorption of charged species in solution. It is then used to adjust or correct the equilibrium constant for the given adsorption reaction.

Parameters

i index of the reaction of interest for the adsorption object

5.1.3.37 double AdsorptionReaction::calculateEquilibriumCorrection (double *sigma*, double *T*, double *I*, double *rel_epsilon*, int *i*)

Function to calculate the correction term for the equilibrium parameter.

This function calculates the correction term that gets applied to the equilibrium parameter to correct for surface charge and charge accumulation/depletion effects. It will call the psi approximation and charge exchange functions, therefore it needs to have those functions arguments passed to it as well.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)
i	index of the reaction of interest for the adsorption object

5.1.3.38 double AdsorptionReaction::Eval_Residual (const Matrix< double > & x, const Matrix< double > & gama, double T, double rel_perm, int i)

Calculates the residual for the ith reaction in the system.

This function will provide a system residual for the ith reaction object involved in the Adsorption Reaction. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously. This function will also adjust the equilibrium parameter for the reaction

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
i	index of the reaction of interest for the adsorption object

5.1.3.39 Reaction& AdsorptionReaction::getReaction (int i)

Return reference to the ith reaction object in the adsorption object.

```
5.1.3.40 double AdsorptionReaction::getMolarFactor ( int i )
Get the ith reaction's molar factor for adsorption (mol/mol)
5.1.3.41 double AdsorptionReaction::getVolumeFactor ( int i )
Get the ith volume factor (species not involved return zeros) (cm<sup>^3</sup>/mol)
5.1.3.42 double AdsorptionReaction::getAreaFactor ( int i )
Get the ith area factor (species not involved return zeros) (m<sup>2</sup>/mol)
5.1.3.43 double AdsorptionReaction::getActivity ( int i )
Get the ith activity factor for the surface species.
5.1.3.44 double AdsorptionReaction::getSpecificArea ( )
Get the specific area of the adsorbent (m<sup>2</sup>/kg) or (mol/kg)
5.1.3.45 double AdsorptionReaction::getSpecificMolality ( )
Get the specific molality of the adsorbent (mol/kg)
5.1.3.46 double AdsorptionReaction::getSurfaceCharge ( )
Get the surface charge of the adsorbent.
5.1.3.47 double AdsorptionReaction::getBulkDensity ( )
Calculate and return bulk density of adsorbent in system (kg/L)
5.1.3.48 double AdsorptionReaction::getTotalMass ( )
Get the total mass of adsorbent in the system (kg)
5.1.3.49 double AdsorptionReaction::getTotalVolume ( )
Get the total volume of the system (L)
5.1.3.50 double AdsorptionReaction::getChargeDensity ( )
Get the value of the surface charge density (C/m<sup>2</sup>)
5.1.3.51 double AdsorptionReaction::getlonicStrength ( )
Get the value of the ionic strength of solution (mol/L)
```

5.1.3.52 int AdsorptionReaction::getNumberRxns ()

Get the number of reactions involved in the adsorption object.

5.1.3.53 int AdsorptionReaction::getAdsorbIndex (int i)

Get the index of the adsorbed species in the ith reaction.

5.1.3.54 int AdsorptionReaction::getAqueousIndex (int i)

Get the index of the primary aqueous species in the ith reaction.

5.1.3.55 int AdsorptionReaction::getActivityEnum ()

Return the enum representing the choosen activity function.

5.1.3.56 bool AdsorptionReaction::isAreaBasis ()

Returns true if we are in the Area Basis, False if in Molar Basis.

5.1.3.57 bool AdsorptionReaction::includeSurfaceCharge ()

Returns true if we are considering surface charging during adsorption.

5.1.3.58 std::string AdsorptionReaction::getAdsorbentName ()

Returns the name of the adsorbent as a string.

5.1.4 Member Data Documentation

5.1.4.1 MasterSpeciesList* AdsorptionReaction::List [protected]

Pointer to the MasterSpeciesList object.

5.1.4.2 int(* AdsorptionReaction::surface_activity) (const Matrix< double > &logq, Matrix< double > &activity, const void *data) [protected]

Pointer to a surface activity model.

This is a function pointer for a surface activity model. The function must accept the log of the surface concentrations as an argument (logq) and provide the activities for each species (activity). The pointer data is used to pass any additional arguments needed.

Parameters

logq	matrix of the log (base 10) of surface concentrations of all species
activity	matrix of activity coefficients for all surface species (must be overriden)
data	pointer to a data structure needed to calculate activities

```
5.1.4.3 const void* AdsorptionReaction::activity_data [protected]
Pointer to the data structure needed for surface activities.
5.1.4.4 int AdsorptionReaction::act_fun [protected]
Enumeration of the activity function being used for the surface phase.
5.1.4.5 std::vector<double> AdsorptionReaction::area_factors [protected]
List of the van der Waals areas associated with surface species (m<sup>2</sup>/mol)
5.1.4.6 std::vector<double> AdsorptionReaction::volume_factors [protected]
List of the van der Waals volumes of each surface species (cm<sup>^</sup>3/mol)
5.1.4.7 std::vector<int> AdsorptionReaction::adsorb_index [protected]
List of the indices for the adsorbed species in the reactions.
5.1.4.8 std::vector<int> AdsorptionReaction::aqueous_index [protected]
List of the indices for the primary aqueous species in the reactions.
5.1.4.9 std::vector<double> AdsorptionReaction::molar_factor [protected]
List of the number of ligands needed to form one mole of adsorption in each reaction.
5.1.4.10 Matrix<double> AdsorptionReaction::activities [protected]
List of the activities calculated by the activity model.
5.1.4.11 double AdsorptionReaction::specific_area [protected]
Specific surface area of the adsorbent (m<sup>2</sup>/kg)
5.1.4.12 double AdsorptionReaction::specific_molality [protected]
Specific molality of the adsorbent - moles of ligand per kg sorbent (mol/kg)
5.1.4.13 double AdsorptionReaction::surface_charge [protected]
Charge of the uncomplexed surface ligand species.
5.1.4.14 double AdsorptionReaction::total_mass [protected]
Total mass of the adsorbent in the system (kg)
```

5.1.4.15 double AdsorptionReaction::total_volume [protected] Total volume of the system (L) **5.1.4.16** double AdsorptionReaction::ionic_strength [protected] Ionic Strength of the system used to adjust equilibria constants (mol/L) **5.1.4.17** double AdsorptionReaction::charge_density [protected] Surface charge density of the adsorbent used to adjust equilbria (C/m²) **5.1.4.18** int AdsorptionReaction::num_rxns [protected] Number of reactions involved in the adsorption equilibria. **5.1.4.19 bool AdsorptionReaction::AreaBasis** [protected]

True = Adsorption on an area basis, False = Adsorption on a ligand basis.

5.1.4.20 bool AdsorptionReaction::IncludeSurfCharge [protected]

True = Includes surface charging corrections, False = Does not consider surface charge.

5.1.4.21 std::string AdsorptionReaction::adsorbent_name [protected]

Name of the adsorbent for this object.

5.1.4.22 std::vector<Reaction> AdsorptionReaction::ads_rxn [private]

List of reactions involved with adsorption.

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

· shark.h

5.2 ARNOLDI_DATA Struct Reference

Data structure for the construction of the Krylov subspaces for a linear system.

#include <lark.h>

Public Attributes

int k

Desired size of the Krylov subspace.

· int iter

Actual size of the Krylov subspace.

· double beta

Normalization parameter.

double hp1

Additional row element of H (separate storage for holding)

• bool Output = true

True = print messages to console.

std::vector< Matrix< double >> Vk

(N) x (k) orthonormal vector basis stored as a vector of column matrices

Matrix< double > Hkp1

(k+1) x (k) upper Hessenberg matrix

Matrix< double > yk

(k) x (1) vector search direction

Matrix< double > e1

(k) x (1) orthonormal vector with 1 in first position

Matrix< double > w

(N) x (1) interim result of the matrix_vector multiplication

Matrix< double > v

(N) x (1) holding cell for the column entries of Vk and other interims

Matrix< double > sum

(N) x (1) running sum of subspace vectors for use in altering w

5.2.1 Detailed Description

Data structure for the construction of the Krylov subspaces for a linear system.

C-style object used in conjunction with the Arnoldi algorithm to construct an orthonormal basis and upper Hessenberg representation of a given linear operator. This is used to solve a linear system both iteratively (i.e., in conjunction with GMRESLP) and directly (i.e., in conjunction with FOM). Alternatively, you can just store the factorized components for later use in another routine.

5.2.2 Member Data Documentation

5.2.2.1 int ARNOLDI DATA::k

Desired size of the Krylov subspace.

5.2.2.2 int ARNOLDI_DATA::iter

Actual size of the Krylov subspace.

5.2.2.3 double ARNOLDI_DATA::beta

Normalization parameter.

5.3 Atom Class Reference 23

5.2.2.4 double ARNOLDI_DATA::hp1

Additional row element of H (separate storage for holding)

5.2.2.5 bool ARNOLDI_DATA::Output = true

True = print messages to console.

5.2.2.6 std::vector< Matrix<double>> ARNOLDI_DATA::Vk

(N) x (k) orthonormal vector basis stored as a vector of column matrices

5.2.2.7 Matrix<double> ARNOLDI_DATA::Hkp1

(k+1) x (k) upper Hessenberg matrix

5.2.2.8 Matrix<double> ARNOLDI_DATA::yk

(k) x (1) vector search direction

5.2.2.9 Matrix<double> ARNOLDI_DATA::e1

(k) x (1) orthonormal vector with 1 in first position

5.2.2.10 Matrix<double> ARNOLDI_DATA::w

(N) x (1) interim result of the matrix_vector multiplication

5.2.2.11 Matrix<double> ARNOLDI_DATA::v

(N) x (1) holding cell for the column entries of Vk and other interims

5.2.2.12 Matrix < double > ARNOLDI_DATA::sum

(N) x (1) running sum of subspace vectors for use in altering w

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

· lark.h

5.3 Atom Class Reference

Atom object to hold information about specific atoms in the periodic table (click Atom to go to function definitions)

#include <eel.h>

Public Member Functions

• Atom ()

Default Constructor.

• ~Atom ()

Default Destructor.

Atom (std::string Name)

Constructor by Atom Name.

• Atom (int number)

Constructor by Atomic number.

void Register (std::string Symbol)

Register an atom object by symbol.

· void Register (int number)

Register an atom object by number.

void editAtomicWeight (double AW)

Manually changes the atomic weight.

void editOxidationState (int state)

Manually changes the oxidation state.

• void editProtons (int proton)

Manually changes the number of protons.

void editNeutrons (int neutron)

Manually changes the number of neutrons.

• void editElectrons (int electron)

Manually changes the number of electrons.

void editValence (int val)

Manually changes the number of valence electrons.

• void editRadii (double r)

Manually changes the van der Waals radii.

void removeProton ()

Manually removes 1 proton and adjusts weight.

• void removeNeutron ()

Manually removes 1 neutron and adjusts weight.

void removeElectron ()

Manually removes 1 electron from valence.

double AtomicWeight ()

Returns the current atomic weight (g/mol)

• int OxidationState ()

Returns the current oxidation state.

• int Protons ()

Returns the current number of protons.

• int Neutrons ()

Returns the current number of neutrons.

• int Electrons ()

Returns the current number of electrons.

• int BondingElectrons ()

Returns the number of electrons available for bonding.

double AtomicRadii ()

Returns the current van der Waals radii (in angstroms)

std::string AtomName ()

Returns the name of the atom.

std::string AtomSymbol ()

Returns the symbol of the atom.

std::string AtomCategory ()

Returns the category of the atom.

std::string AtomState ()

Returns the state of the atom.

int AtomicNumber ()

Returns the atomic number of the atom.

void DisplayInfo ()

Displays Atom information to console.

Protected Attributes

· double atomic_weight

Holds the atomic weight of the atom.

int oxidation_state

Holds the oxidation state of the atom.

· int protons

Holds the number of protons in the atom.

· int neutrons

Holds the number of neutrons in the atom.

· int electrons

Holds the number of electrons in the atom.

• int valence_e

Holds the number of valence electrons in the atom.

· double atomic_radii

Holds the van der Waals radii of the element (in angstroms)

Private Attributes

• std::string Name

Holds the name of the atom.

• std::string Symbol

Holds the atomic symbol for the atom.

• std::string Category

Holds the category of the atom (e.g., Alkali Metal)

std::string NaturalState

Holds the natural state of the atom (e.g., Gas)

• int atomic_number

Holds the atomic number of the atom.

5.3.1 Detailed Description

Atom object to hold information about specific atoms in the periodic table (click Atom to go to function definitions)

C++ class object holding data and functions associated with atoms. Objects can be registered at the time of object construction, or after declaring an Atom object. Registration can be done via the atomic symbol or atomic number. Valid atoms go from Hydrogen (1) to Ununoctium (118).

```
5.3.2 Constructor & Destructor Documentation
5.3.2.1 Atom::Atom()
Default Constructor.
5.3.2.2 Atom::∼Atom ( )
Default Destructor.
5.3.2.3 Atom::Atom ( std::string Name )
Constructor by Atom Name.
5.3.2.4 Atom::Atom ( int number )
Constructor by Atomic number.
5.3.3 Member Function Documentation
5.3.3.1 void Atom::Register ( std::string Symbol )
Register an atom object by symbol.
5.3.3.2 void Atom::Register (int number)
Register an atom object by number.
5.3.3.3 void Atom::editAtomicWeight ( double AW )
Manually changes the atomic weight.
5.3.3.4 void Atom::editOxidationState (int state)
Manually changes the oxidation state.
5.3.3.5 void Atom::editProtons (int proton)
Manually changes the number of protons.
5.3.3.6 void Atom::editNeutrons (int neutron)
Manually changes the number of neutrons.
5.3.3.7 void Atom::editElectrons (int electron)
Manually changes the number of electrons.
```

5.3 Atom Class Reference

```
5.3.3.8 void Atom::editValence (int val)
Manually changes the number of valence electrons.
5.3.3.9 void Atom::editRadii (double r)
Manually changes the van der Waals radii.
5.3.3.10 void Atom::removeProton()
Manually removes 1 proton and adjusts weight.
5.3.3.11 void Atom::removeNeutron()
Manually removes 1 neutron and adjusts weight.
5.3.3.12 void Atom::removeElectron ( )
Manually removes 1 electron from valence.
5.3.3.13 double Atom::AtomicWeight ( )
Returns the current atomic weight (g/mol)
5.3.3.14 int Atom::OxidationState ( )
Returns the current oxidation state.
5.3.3.15 int Atom::Protons ( )
Returns the current number of protons.
5.3.3.16 int Atom::Neutrons ( )
Returns the current number of neutrons.
5.3.3.17 int Atom::Electrons ( )
Returns the current number of electrons.
5.3.3.18 int Atom::BondingElectrons ( )
Returns the number of electrons available for bonding.
5.3.3.19 double Atom::AtomicRadii ( )
Returns the current van der Waals radii (in angstroms)
```

```
5.3.3.20 std::string Atom::AtomName ( )
Returns the name of the atom.
5.3.3.21 std::string Atom::AtomSymbol ( )
Returns the symbol of the atom.
5.3.3.22 std::string Atom::AtomCategory ( )
Returns the category of the atom.
5.3.3.23 std::string Atom::AtomState ( )
Returns the state of the atom.
5.3.3.24 int Atom::AtomicNumber ( )
Returns the atomic number of the atom.
5.3.3.25 void Atom::DisplayInfo ( )
Displays Atom information to console.
5.3.4 Member Data Documentation
5.3.4.1 double Atom::atomic_weight [protected]
Holds the atomic weight of the atom.
5.3.4.2 int Atom::oxidation_state [protected]
Holds the oxidation state of the atom.
5.3.4.3 int Atom::protons [protected]
Holds the number of protons in the atom.
5.3.4.4 int Atom::neutrons [protected]
Holds the number of neutrons in the atom.
5.3.4.5 int Atom::electrons [protected]
Holds the number of electrons in the atom.
5.3.4.6 int Atom::valence_e [protected]
Holds the number of valence electrons in the atom.
```

```
5.3.4.7 double Atom::atomic_radii [protected]
```

Holds the van der Waals radii of the element (in angstroms)

```
5.3.4.8 std::string Atom::Name [private]
```

Holds the name of the atom.

```
5.3.4.9 std::string Atom::Symbol [private]
```

Holds the atomic symbol for the atom.

```
5.3.4.10 std::string Atom::Category [private]
```

Holds the category of the atom (e.g., Alkali Metal)

```
5.3.4.11 std::string Atom::NaturalState [private]
```

Holds the natural state of the atom (e.g., Gas)

```
5.3.4.12 int Atom::atomic_number [private]
```

Holds the atomic number of the atom.

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

· eel.h

5.4 BACKTRACK DATA Struct Reference

Data structure for the implementation of Backtracking Linesearch.

```
#include <lark.h>
```

Public Attributes

• int fun_call = 0

Number of function calls made during line search.

• double alpha = 1e-4

Scaling parameter for determination of search step size.

• double rho = 0.1

Scaling parameter for to change step size by.

• double lambdaMin =DBL_EPSILON

Smallest allowable step length.

double normFkp1

New residual norm of the Newton step.

• bool constRho = false

True = use a constant value for rho.

Matrix< double > Fk

Old residual vector of the Newton step.

Matrix< double > xk

Old solution vector of the Newton step.

5.4.1 Detailed Description

Data structure for the implementation of Backtracking Linesearch.

C-style object used in conjunction with the Backtracking Linesearch algorithm to smooth out convergence of Netwon based iterative methods for non-linear systems of equations. The actual algorithm has been separated from the interior of the Newton method so that it can be included in any future Newton based iterative methods being developed.

5.4.2 Member Data Documentation

5.4.2.1 int BACKTRACK_DATA::fun_call = 0

Number of function calls made during line search.

5.4.2.2 double BACKTRACK_DATA::alpha = 1e-4

Scaling parameter for determination of search step size.

5.4.2.3 double BACKTRACK_DATA::rho = 0.1

Scaling parameter for to change step size by.

5.4.2.4 double BACKTRACK_DATA::lambdaMin = DBL_EPSILON

Smallest allowable step length.

5.4.2.5 double BACKTRACK_DATA::normFkp1

New residual norm of the Newton step.

5.4.2.6 bool BACKTRACK_DATA::constRho = false

True = use a constant value for rho.

5.4.2.7 Matrix < double > BACKTRACK_DATA::Fk

Old residual vector of the Newton step.

5.4.2.8 Matrix < double > BACKTRACK_DATA::xk

Old solution vector of the Newton step.

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

· lark.h

5.5 BiCGSTAB_DATA Struct Reference

Data structure for the implementation of the BiCGSTAB algorithm for non-symmetric linear systems.

```
#include <lark.h>
```

Public Attributes

• int maxit = 0

Maximum allowable iterations - default = min(2*vector_size,1000)

• int iter = 0

Actual number of iterations.

· bool breakdown

Boolean to determine if the method broke down.

· double alpha

Step size parameter for next solution.

· double beta

Step size parameter for search direction.

• double rho

Scaling parameter for alpha and beta.

· double rho_old

Previous scaling parameter for alpha and beta.

· double omega

Scaling parameter and additional step length.

• double omega_old

Previous scaling parameter and step length.

double tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

• double tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

double res

Absolute residual norm.

· double relres

Relative residual norm.

double relres_base

Initial residual norm.

· double bestres

Best found residual norm.

• bool Output = true

True = print messages to console.

Matrix< double > x

Current solution to the linear system.

• Matrix< double > bestx

Best found solution to the linear system.

• Matrix < double > r

Residual vector for the linear system.

Matrix< double > r0

Initial residual vector.

Matrix< double > v

Search direction for p.

Matrix< double > p

Search direction for updating.

Matrix< double > y

Preconditioned search direction.

Matrix< double > s

Residual updating vector.

Matrix< double > z

Preconditioned residual updating vector.

Matrix< double > t

Search direction for resdidual updates.

5.5.1 Detailed Description

Data structure for the implementation of the BiCGSTAB algorithm for non-symmetric linear systems.

C-style object used in conjunction with the Bi-Conjugate Gradient STABalized (BiCGSTAB) algorithm to solve a linear system of equations. This algorithm is generally more efficient than any GMRES or GCR variant, but may not always reduce the residual at each step. However, if used with preconditioning, then this algorithm is very efficient, especially when used for solving grid-based linear systems.

5.5.2 Member Data Documentation

5.5.2.1 int BiCGSTAB_DATA::maxit = 0

Maximum allowable iterations - default = min(2*vector_size,1000)

5.5.2.2 int BiCGSTAB_DATA::iter = 0

Actual number of iterations.

5.5.2.3 bool BiCGSTAB DATA::breakdown

Boolean to determine if the method broke down.

5.5.2.4 double BiCGSTAB_DATA::alpha

Step size parameter for next solution.

5.5.2.5 double BiCGSTAB_DATA::beta

Step size parameter for search direction.

5.5.2.6 double BiCGSTAB_DATA::rho

Scaling parameter for alpha and beta.

5.5.2.7 double BiCGSTAB_DATA::rho_old

Previous scaling parameter for alpha and beta.

5.5.2.8 double BiCGSTAB_DATA::omega

Scaling parameter and additional step length.

5.5.2.9 double BiCGSTAB_DATA::omega_old

Previous scaling parameter and step length.

5.5.2.10 double BiCGSTAB_DATA::tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

5.5.2.11 double BiCGSTAB_DATA::tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

5.5.2.12 double BiCGSTAB_DATA::res

Absolute residual norm.

5.5.2.13 double BiCGSTAB_DATA::relres

Relative residual norm.

5.5.2.14 double BiCGSTAB_DATA::relres_base

Initial residual norm.

5.5.2.15 double BiCGSTAB_DATA::bestres

Best found residual norm.

5.5.2.16 bool BiCGSTAB_DATA::Output = true

True = print messages to console.

5.5.2.17 Matrix<double> BiCGSTAB_DATA::x

Current solution to the linear system.

 $\textbf{5.5.2.18} \quad \textbf{Matrix}{<} \textbf{double}{>} \, \textbf{BiCGSTAB_DATA}{::} \textbf{bestx}$

Best found solution to the linear system.

 $\textbf{5.5.2.19} \quad \textbf{Matrix}{<} \textbf{double}{>} \, \textbf{BiCGSTAB_DATA}{::r}$

Residual vector for the linear system.

5.5.2.20 Matrix<double> BiCGSTAB_DATA::r0 Initial residual vector. 5.5.2.21 Matrix<double> BiCGSTAB_DATA::v Search direction for p. 5.5.2.22 Matrix<double> BiCGSTAB_DATA::p Search direction for updating. 5.5.2.23 Matrix<double> BiCGSTAB_DATA::y Preconditioned search direction. 5.5.2.24 Matrix<double> BiCGSTAB_DATA::s Residual updating vector. 5.5.2.25 Matrix<double> BiCGSTAB_DATA::z Preconditioned residual updating vector. 5.5.2.26 Matrix<double> BiCGSTAB_DATA::t Search direction for resdidual updates. The documentation for this struct was generated from the following file: · lark.h 5.6 CGS_DATA Struct Reference

Data structure for the implementation of the CGS algorithm for non-symmetric linear systems.

#include <lark.h>

Generated by Doxygen

Public Attributes

• int maxit = 0

Maximum allowable iterations - default = min(2*vector_size,1000)

• int iter = 0

Actual number of iterations.

· bool breakdown

Boolean to determine if the method broke down.

· double alpha

Step size parameter for next solution.

· double beta

Step size parameter for search direction.

· double rho

Scaling parameter for alpha and beta.

· double sigma

Scaling parameter and additional step length.

• double tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

double tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

double res

Absolute residual norm.

· double relres

Relative residual norm.

· double relres_base

Initial residual norm.

· double bestres

Best found residual norm.

• bool Output = true

True = print messages to console.

Matrix< double > x

Current solution to the linear system.

• Matrix< double > bestx

Best found solution to the linear system.

Matrix< double > r

Residual vector for the linear system.

• Matrix< double > r0

Initial residual vector.

Matrix< double > u

Search direction for v.

Matrix< double > w

Updates sigma and u.

Matrix< double > v

Search direction for x.

• Matrix< double > p

Preconditioning result for w, z, and matvec for Ax.

Matrix< double > c

Holds the matvec result between A and p.

Matrix< double > z

Full search direction for x.

5.6.1 Detailed Description

Data structure for the implementation of the CGS algorithm for non-symmetric linear systems.

C-style object to be used in conjunction with the Conjugate Gradient Squared (CGS) algorithm to solve linear systems of equations. This algorithm is slightly less computational work than BiCGSTAB, but is much less stable. As a result, I do not recommend using this algorithm unless you also use some form of preconditioning.

5.6.2 Member Data Documentation

5.6.2.1 int CGS_DATA::maxit = 0

Maximum allowable iterations - default = min(2*vector_size,1000)

5.6.2.2 int CGS_DATA::iter = 0

Actual number of iterations.

5.6.2.3 bool CGS_DATA::breakdown

Boolean to determine if the method broke down.

5.6.2.4 double CGS_DATA::alpha

Step size parameter for next solution.

5.6.2.5 double CGS_DATA::beta

Step size parameter for search direction.

5.6.2.6 double CGS_DATA::rho

Scaling parameter for alpha and beta.

5.6.2.7 double CGS_DATA::sigma

Scaling parameter and additional step length.

5.6.2.8 double CGS_DATA::tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

5.6.2.9 double CGS_DATA::tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

5.6.2.10 double CGS_DATA::res

Absolute residual norm.

5.6.2.11 double CGS_DATA::relres

Relative residual norm.

5.6.2.12 double CGS_DATA::relres_base

Initial residual norm.

5.6.2.13 double CGS_DATA::bestres

Best found residual norm.

5.6.2.14 bool CGS_DATA::Output = true

True = print messages to console.

 $\textbf{5.6.2.15} \quad \textbf{Matrix}{<} \textbf{double}{>} \textbf{CGS_DATA}{::} \textbf{x}$

Current solution to the linear system.

5.6.2.16 Matrix<double> CGS_DATA::bestx

Best found solution to the linear system.

5.6.2.17 Matrix<double> CGS_DATA::r

Residual vector for the linear system.

5.6.2.18 Matrix<double> CGS_DATA::r0

Initial residual vector.

 $\textbf{5.6.2.19} \quad \textbf{Matrix}{<} \textbf{double}{>} \textbf{CGS_DATA}{::} \textbf{u}$

Search direction for v.

5.6.2.20 Matrix<double> CGS_DATA::w

Updates sigma and u.

 $\textbf{5.6.2.21} \quad \textbf{Matrix}{<} \textbf{double}{>} \textbf{CGS_DATA}{::} \textbf{v}$

Search direction for x.

5.6.2.22 Matrix<double> CGS_DATA::p

Preconditioning result for w, z, and matvec for Ax.

```
5.6.2.23 Matrix<double> CGS_DATA::c
```

Holds the matvec result between A and p.

```
5.6.2.24 Matrix<double> CGS_DATA::z
```

Full search direction for x.

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

· lark.h

5.7 ChemisorptionReaction Class Reference

Chemisorption Reaction Object.

```
#include <shark.h>
```

Inheritance diagram for ChemisorptionReaction:



Public Member Functions

ChemisorptionReaction ()

Default Constructor.

∼ChemisorptionReaction ()

Default Destructor.

void Initialize_Object (MasterSpeciesList &List, int n)

Function to call the initialization of objects sequentially.

• void Display_Info ()

Display the adsorption reaction information.

• void modifyMBEdeltas (MassBalance &mbo)

Modify the Deltas in the MassBalance Object.

• int setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object.

• int setLigandIndex ()

Find and set the ligand species index.

• int setDeltas ()

Find and set all the delta values for the site balance.

 void setActivityModelInfo (int(*act)(const Matrix< double > &logq, Matrix< double > &activity, const void *data), const void *act_data)

Function to set the surface activity model and data pointer.

void setActivityEnum (int act)

Set the surface activity enum value.

void setDelta (int i, double v)

Set the ith delta factor for the site balance.

void setVolumeFactor (int i, double v)

Set the ith volume factor for the species list (cm^{\(\circ\)}3/mol)

void setAreaFactor (int i, double a)

Set the ith area factor for the species list (m^2/mol)

void setSpecificArea (double a)

Set the specific area for the adsorbent (m²/kg)

void setSpecificMolality (double a)

Set the specific molality for the adsorbent (mol/kg)

void setTotalMass (double m)

Set the total mass of the adsorbent (kg)

void setTotalVolume (double v)

Set the total volume of the system (L)

void setSurfaceChargeBool (bool opt)

Set the boolean for inclusion of surface charging.

void setAdsorbentName (std::string name)

Set the name of the adsorbent to the given string.

void setChargeDensityValue (double a)

Set the value of the charge density parameter to a (C/m^2)

• void setlonicStrengthValue (double a)

Set the value of the ionic strength parameter to a (mol/L)

void setActivities (Matrix< double > &x)

Set the values of activities in the activity matrix.

void calculateAreaFactors ()

Calculates the area factors used from the van der Waals volumes.

• void calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

void setChargeDensity (const Matrix< double > &x)

Calculates and sets the current value of charge density.

void setlonicStrength (const Matrix< double > &x)

Calculates and sets the current value of ionic strength.

int callSurfaceActivity (const Matrix< double > &x)

Calls the activity model and returns an int flag for success or failure.

- double calculateSurfaceChargeDensity (const Matrix < double > &x)

Function to calculate the surface charge density based on concentrations.

• double calculateElecticPotential (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

double calculateAqueousChargeExchange (int i)

Function to calculate the net exchange of charges of the aqeous species involved in a given reaction.

double calculateEquilibriumCorrection (double sigma, double T, double I, double rel_epsilon, int i)

Function to calculate the correction term for the equilibrium parameter.

double Eval_RxnResidual (const Matrix< double > &x, const Matrix< double > &gama, double T, double rel perm, int i)

Calculates the residual for the ith reaction in the system.

double Eval SiteBalanceResidual (const Matrix< double > &x)

Calculates the residual for the overall site balance.

Reaction & getReaction (int i)

Return reference to the ith reaction object in the adsorption object.

double getDelta (int i)

Get the ith delta factor for the site balance.

• double getVolumeFactor (int i)

Get the ith volume factor (species not involved return zeros) (cm[^]3/mol)

double getAreaFactor (int i)

Get the ith area factor (species not involved return zeros) (m²/mol)

double getActivity (int i)

Get the ith activity factor for the surface species.

double getSpecificArea ()

Get the specific area of the adsorbent (m^2/kg) or (mol/kg)

double getSpecificMolality ()

Get the specific molality of the adsorbent (mol/kg)

• double getBulkDensity ()

Calculate and return bulk density of adsorbent in system (kg/L)

double getTotalMass ()

Get the total mass of adsorbent in the system (kg)

• double getTotalVolume ()

Get the total volume of the system (L)

double getChargeDensity ()

Get the value of the surface charge density (C/m^2)

• double getlonicStrength ()

Get the value of the ionic strength of solution (mol/L)

• int getNumberRxns ()

Get the number of reactions involved in the adsorption object.

• int getAdsorbIndex (int i)

Get the index of the adsorbed species in the ith reaction.

• int getLigandIndex ()

Get the index of the ligand species.

• int getActivityEnum ()

Return the enum representing the choosen activity function.

· bool includeSurfaceCharge ()

Returns true if we are considering surface charging during adsorption.

• std::string getAdsorbentName ()

Returns the name of the adsorbent as a string.

Protected Attributes

· int ligand index

Index of the ligand for all reactions.

std::vector< double > Delta

Vector of weights (i.e., deltas) used in the site balance.

Private Attributes

• std::vector< Reaction > ads rxn

List of reactions involved with adsorption.

Additional Inherited Members

5.7.1 Detailed Description

Chemisorption Reaction Object.

C++ Object to handle data and functions associated with forumlating adsorption equilibrium reactions in a aqueous mixture based on chemisorption mechanisms. Each unique surface in a system will require an instance of this structure. This is very similar to AdsorptionReaction, however, this will include a site balance residual that will allow us to consider protonation and deprotonation of the ligands.

5.7.2 Constructor & Destructor Documentation

5.7.2.1 ChemisorptionReaction::ChemisorptionReaction ()

Default Constructor.

5.7.2.2 ChemisorptionReaction:: ∼ ChemisorptionReaction ()

Default Destructor.

5.7.3 Member Function Documentation

5.7.3.1 void ChemisorptionReaction::Initialize_Object (MasterSpeciesList & List, int n)

Function to call the initialization of objects sequentially.

5.7.3.2 void ChemisorptionReaction::Display_Info ()

Display the adsorption reaction information.

5.7.3.3 void ChemisorptionReaction::modifyMBEdeltas (MassBalance & mbo)

Modify the Deltas in the MassBalance Object.

This function will take a mass balance object as an argument and modify the deltas in that object to correct for how adsorption affects that particular mass balance. Since adsorption can effect multiple mass balances, this function must be called for each mass balance in the system.

Parameters

mbo reference to the MassBalance Object the adsorption is acting on

5.7.3.4 int ChemisorptionReaction::setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object.

This function searches through the Reaction objects in ChemisorptionReaction to find the adsorbed species and their indices to set that information in the adsorb_index structure. Function will return 0 if successful and -1 on a failure.

```
5.7.3.5 int ChemisorptionReaction::setLigandIndex ( )
```

Find and set the ligand species index.

This function searches through the Reaction objects in ChemisorptionReaction to find the ligand species and its index to set that information in the ligand index structure. Function will return 0 if successful and -1 on a failure.

```
5.7.3.6 int ChemisorptionReaction::setDeltas ( )
```

Find and set all the delta values for the site balance.

This function searches through all reaction object instances for the stoicheometry of the ligand in each adsorption reaction. That stoicheometry serves as the basis for determining the site balance. NOTE: the delta for the ligand is set automatically in the setLigandIndex() function, so we can ignore that species. In addition, this function must be called after setLigandIndex() and setAdsorbIndices() are called and after the stoicheometry of each reaction has been determined.

```
5.7.3.7 void ChemisorptionReaction::setActivityModelInfo ( int(*)(const Matrix< double > &logq, Matrix< double > &activity, const void *data) act, const void * act_data )
```

Function to set the surface activity model and data pointer.

This function will setup the surface activity model based on the given pointer arguments. If no arguments are given, or are given as NULL, then the activity model will default to ideal solution assumption.

```
5.7.3.8 void ChemisorptionReaction::setActivityEnum (int act)
```

Set the surface activity enum value.

```
5.7.3.9 void ChemisorptionReaction::setDelta (int i, double v)
```

Set the ith delta factor for the site balance.

```
5.7.3.10 void ChemisorptionReaction::setVolumeFactor (int i, double v)
```

Set the ith volume factor for the species list (cm³/mol)

5.7.3.11 void ChemisorptionReaction::setAreaFactor (int i, double a)

Set the ith area factor for the species list (m²/mol)

5.7.3.12 void ChemisorptionReaction::setSpecificArea (double a)

Set the specific area for the adsorbent (m²/kg)

5.7.3.13 void ChemisorptionReaction::setSpecificMolality (double a)

Set the specific molality for the adsorbent (mol/kg)

```
5.7.3.14 void ChemisorptionReaction::setTotalMass ( double m )
Set the total mass of the adsorbent (kg)
5.7.3.15 void ChemisorptionReaction::setTotalVolume ( double v )
Set the total volume of the system (L)
5.7.3.16 void ChemisorptionReaction::setSurfaceChargeBool (bool opt)
Set the boolean for inclusion of surface charging.
5.7.3.17 void ChemisorptionReaction::setAdsorbentName ( std::string name )
Set the name of the adsorbent to the given string.
5.7.3.18 void ChemisorptionReaction::setChargeDensityValue ( double a )
Set the value of the charge density parameter to a (C/m^2)
5.7.3.19 void ChemisorptionReaction::setlonicStrengthValue ( double a )
Set the value of the ionic strength parameter to a (mol/L)
5.7.3.20 void ChemisorptionReaction::setActivities ( Matrix< double > & x )
Set the values of activities in the activity matrix.
5.7.3.21 void ChemisorptionReaction::calculateAreaFactors ( )
Calculates the area factors used from the van der Waals volumes.
5.7.3.22 void ChemisorptionReaction::calculateEquilibria ( double T )
Calculates all equilibrium parameters as a function of temperature.
5.7.3.23 void ChemisorptionReaction::setChargeDensity (const Matrix < double > & x)
Calculates and sets the current value of charge density.
5.7.3.24 void ChemisorptionReaction::setlonicStrength (const Matrix < double > & x)
Calculates and sets the current value of ionic strength.
5.7.3.25 int ChemisorptionReaction::callSurfaceActivity ( const Matrix < double > & x )
Calls the activity model and returns an int flag for success or failure.
5.7.3.26 double ChemisorptionReaction::calculateSurfaceChargeDensity ( const Matrix < double > & x )
Function to calculate the surface charge density based on concentrations.
```

This function is used to calculate the surface charge density of the adsorbed species based on the charges and concentrations of the adsorbed species. The calculation is used to correct the adsorption equilibria constant based on a localized surface charge balance. This requires that you know the molality of the uncomplexed ligand species on the surface, as well as the specific surface area for the adsorbent.

Parameters

x matrix of the log(C) concentration values at the current non-linear step

5.7.3.27 double ChemisorptionReaction::calculateElecticPotential (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

This function will calculate the electric surface potential of the adsorbent under the current conditions of charge density, temperature, ionic strength, and relative permittivity.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)

5.7.3.28 double ChemisorptionReaction::calculateAqueousChargeExchange (int i)

Function to calculate the net exchange of charges of the ageous species involved in a given reaction.

This function will look at all aqueous species involved in the ith adsorption reaction and sum up their stoicheometries and charges to see what the net change in charge is caused by the adsorption of charged species in solution. It is then used to adjust or correct the equilibrium constant for the given adsorption reaction.

Parameters

i index of the reaction of interest for the adsorption object

5.7.3.29 double ChemisorptionReaction::calculateEquilibriumCorrection (double *sigma*, double *T*, double *I*, double *rel_epsilon*, int *i*)

Function to calculate the correction term for the equilibrium parameter.

This function calculates the correction term that gets applied to the equilibrium parameter to correct for surface charge and charge accumulation/depletion effects. It will call the psi approximation and charge exchange functions, therefore it needs to have those functions arguments passed to it as well.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)
i	index of the reaction of interest for the adsorption object

5.7.3.30 double ChemisorptionReaction::Eval_RxnResidual (const Matrix< double > & x, const Matrix< double > & gama, double T, double rel_perm, int i)

Calculates the residual for the ith reaction in the system.

This function will provide a system residual for the ith reaction object involved in the Adsorption Reaction. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously. This function will also adjust the equilibrium parameter for the reaction

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
i	index of the reaction of interest for the adsorption object

5.7.3.31 double ChemisorptionReaction::Eval SiteBalanceResidual (const Matrix < double > & x)

Calculates the residual for the overall site balance.

This function will provide a system residual for the site/ligand balance for the Chemisorption Reaction object. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously.

Parameters

x matrix of the log(C) concentration values at the current non-linear step

5.7.3.32 Reaction& ChemisorptionReaction::getReaction (int i)

Return reference to the ith reaction object in the adsorption object.

5.7.3.33 double ChemisorptionReaction::getDelta (int i)

Get the ith delta factor for the site balance.

5.7.3.34 double ChemisorptionReaction::getVolumeFactor (int i)

Get the ith volume factor (species not involved return zeros) (cm^{\(\chi\)}3/mol)

5.7.3.35 double ChemisorptionReaction::getAreaFactor (int i)

Get the ith area factor (species not involved return zeros) (m²/mol)

5.7.3.36 double ChemisorptionReaction::getActivity (int i)

Get the ith activity factor for the surface species.

```
5.7.3.37 double ChemisorptionReaction::getSpecificArea ( )
Get the specific area of the adsorbent (m<sup>2</sup>/kg) or (mol/kg)
5.7.3.38 double ChemisorptionReaction::getSpecificMolality ( )
Get the specific molality of the adsorbent (mol/kg)
5.7.3.39 double ChemisorptionReaction::getBulkDensity ( )
Calculate and return bulk density of adsorbent in system (kg/L)
5.7.3.40 double ChemisorptionReaction::getTotalMass ( )
Get the total mass of adsorbent in the system (kg)
5.7.3.41 double ChemisorptionReaction::getTotalVolume ( )
Get the total volume of the system (L)
5.7.3.42 double ChemisorptionReaction::getChargeDensity ( )
Get the value of the surface charge density (C/m^2)
5.7.3.43 double ChemisorptionReaction::getlonicStrength ( )
Get the value of the ionic strength of solution (mol/L)
5.7.3.44 int ChemisorptionReaction::getNumberRxns ( )
Get the number of reactions involved in the adsorption object.
5.7.3.45 int ChemisorptionReaction::getAdsorbIndex (int i)
Get the index of the adsorbed species in the ith reaction.
5.7.3.46 int ChemisorptionReaction::getLigandIndex ( )
Get the index of the ligand species.
5.7.3.47 int ChemisorptionReaction::getActivityEnum ( )
Return the enum representing the choosen activity function.
5.7.3.48 bool ChemisorptionReaction::includeSurfaceCharge ( )
Returns true if we are considering surface charging during adsorption.
```

5.7.3.49 std::string ChemisorptionReaction::getAdsorbentName ()

Returns the name of the adsorbent as a string.

5.7.4 Member Data Documentation

5.7.4.1 int ChemisorptionReaction::ligand_index [protected]

Index of the ligand for all reactions.

5.7.4.2 std::vector<**double**> **ChemisorptionReaction::Delta** [protected]

Vector of weights (i.e., deltas) used in the site balance.

5.7.4.3 std::vector<Reaction> ChemisorptionReaction::ads_rxn [private]

List of reactions involved with adsorption.

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

· shark.h

5.8 Document Class Reference

Object for the various documents in the yaml file.

```
#include <yaml_wrapper.h>
```

Inheritance diagram for Document:



Public Member Functions

Document ()

Default constructor.

∼Document ()

Default destructor.

Document (const Document &doc)

Copy constructor.

Document (std::string name)

Constructor by name.

• Document (const KeyValueMap &map)

Constructor by existing map.

Document (std::string name, const KeyValueMap &map)

Constructor by name and map.

• Document (std::string key, const Header &head)

Constructor by single header.

• Document & operator= (const Document &doc)

Equals overload.

ValueTypePair & operator[] (const std::string key)

Return the ValueType reference at the given key.

ValueTypePair operator[] (const std::string key) const

Return the ValueType at the given key.

Header & operator() (const std::string key)

Return the Header reference at the given key.

Header operator() (const std::string key) const

Return the Header at the given key.

std::map< std::string, Header > & getHeadMap ()

Return the reference to the Header Map.

KeyValueMap & getDataMap ()

Return the reference to the KeyValueMap.

Header & getHeader (std::string key)

Return reference to the Header in map at the key.

std::map< std::string, Header >::const_iterator end () const

Returns a const iterator pointing to the end of the list.

std::map< std::string, Header >::iterator end ()

Returns an iterator pointing to the end of the list.

• std::map< std::string, Header >::const_iterator begin () const

Returns a const iterator pointing to the begining of the list.

std::map< std::string, Header >::iterator begin ()

Returns an iterator pointing to the begining of the list.

• void clear ()

Clear out info in the Document.

· void resetKeys ()

Set all keys in the map to match names of the headers.

void changeKey (std::string oldKey, std::string newKey)

Change a given oldKey in the header map to the newKey given.

void revalidateAllKeys ()

Resets and validates keys in header and subheader maps.

void addPair (std::string key, std::string val)

Adds a pair object to the map (with only strings)

void addPair (std::string key, std::string val, int t)

Adds a pair object and asserts a type.

void setName (std::string name)

Set the name of the Document.

void setAlias (std::string alias)

Set the alias of the Document.

void setNameAliasPair (std::string n, std::string a, int s)

Set the name, alias, and state of the document.

void setState (int state)

Set the state of the Document.

• void DisplayContents ()

Display the contents of the Document.

void addHeadKey (std::string key)

Add a key to the Header without a header object.

void copyAnchor2Alias (std::string alias, Header &ref)

Find the anchor in the map, and copy to the Header reference given.

• int size ()

Return the size of the header map.

std::string getName ()

Return the name of the document.

• std::string getAlias ()

Return the alias of the document.

• int getState ()

Return the state of the document.

· bool isAlias ()

Returns true if the document is an alias.

· bool isAnchor ()

Returns true if the document is an anchor.

Header & getAnchoredHeader (std::string alias)

Returns reference to the anchored header, if any.

Header & getHeadFromSubAlias (std::string alias)

Returns reference to the Header that contains a Sub with the given alias.

Private Attributes

std::map< std::string, Header > Head_Map
 Map of headers contained within the document.

Additional Inherited Members

5.8.1 Detailed Description

Object for the various documents in the yaml file.

C++ Object for the documents in a yaml input file as denoted by a Key: followed by — (three dashes) and ending with a ... (three dots). A single yaml file can have multiple document structures and each document structure can have multiple headers (which have sub-headers and key-values) and key-value-pairs. This is the larges single object in the yaml file itself.

Just like Header, this object also inherits from SubHeader and therefore has access to its protected members. You can use access to those members to establish the KeyValuePairs in the Document, name the Document, and give the Document an alias or anchor value.

```
5.8.2 Constructor & Destructor Documentation
5.8.2.1 Document::Document ( )
Default constructor.
5.8.2.2 Document::~Document ( )
Default destructor.
5.8.2.3 Document::Document ( const Document & doc )
Copy constructor.
5.8.2.4 Document::Document ( std::string name )
Constructor by name.
5.8.2.5 Document::Document ( const KeyValueMap & map )
Constructor by existing map.
5.8.2.6 Document::Document ( std::string name, const KeyValueMap & map )
Constructor by name and map.
5.8.2.7 Document::Document ( std::string key, const Header & head )
Constructor by single header.
5.8.3 Member Function Documentation
5.8.3.1 Document& Document::operator= ( const Document & doc )
Equals overload.
5.8.3.2 ValueTypePair& Document::operator[]( const std::string key )
Return the ValueType reference at the given key.
5.8.3.3 ValueTypePair Document::operator[]( const std::string key ) const
Return the ValueType at the given key.
5.8.3.4 Header& Document::operator() ( const std::string key )
Return the Header reference at the given key.
```

```
5.8.3.5 Header Document::operator() ( const std::string key ) const
Return the Header at the given key.
5.8.3.6 std::map<std::string, Header>& Document::getHeadMap ( )
Return the reference to the Header Map.
5.8.3.7 KeyValueMap& Document::getDataMap ( )
Return the reference to the KeyValueMap.
5.8.3.8 Header& Document::getHeader ( std::string key )
Return reference to the Header in map at the key.
5.8.3.9 std::map<std::string, Header>::const_iterator Document::end ( ) const
Returns a const iterator pointing to the end of the list.
5.8.3.10 std::map<std::string, Header>::iterator Document::end ( )
Returns an iterator pointing to the end of the list.
5.8.3.11 std::map<std::string, Header>::const_iterator Document::begin ( ) const
Returns a const iterator pointing to the begining of the list.
5.8.3.12 std::map<std::string, Header>::iterator Document::begin ( )
Returns an iterator pointing to the begining of the list.
5.8.3.13 void Document::clear ( )
Clear out info in the Document.
5.8.3.14 void Document::resetKeys ( )
Set all keys in the map to match names of the headers.
5.8.3.15 void Document::changeKey ( std::string oldKey, std::string newKey )
Change a given oldKey in the header map to the newKey given.
5.8.3.16 void Document::revalidateAllKeys ( )
Resets and validates keys in header and subheader maps.
```

```
5.8.3.17 void Document::addPair ( std::string key, std::string val )
Adds a pair object to the map (with only strings)
5.8.3.18 void Document::addPair ( std::string key, std::string val, int t )
Adds a pair object and asserts a type.
5.8.3.19 void Document::setName ( std::string name )
Set the name of the Document.
5.8.3.20 void Document::setAlias ( std::string alias )
Set the alias of the Document.
5.8.3.21 void Document::setNameAliasPair ( std::string n, std::string a, int s )
Set the name, alias, and state of the document.
5.8.3.22 void Document::setState ( int state )
Set the state of the Document.
5.8.3.23 void Document::DisplayContents ( )
Display the contents of the Document.
5.8.3.24 void Document::addHeadKey ( std::string key )
Add a key to the Header without a header object.
5.8.3.25 void Document::copyAnchor2Alias ( std::string alias, Header & ref )
Find the anchor in the map, and copy to the Header reference given.
5.8.3.26 int Document::size ( )
Return the size of the header map.
5.8.3.27 std::string Document::getName ( )
Return the name of the document.
5.8.3.28 std::string Document::getAlias ( )
Return the alias of the document.
```

```
5.8.3.29 int Document::getState ( )
Return the state of the document.
5.8.3.30 bool Document::isAlias ( )
Returns true if the document is an alias.
5.8.3.31 bool Document::isAnchor ( )
Returns true if the document is an anchor.
5.8.3.32 Header & Document::getAnchoredHeader ( std::string alias )
Returns reference to the anchored header, if any.
5.8.3.33 Header& Document::getHeadFromSubAlias ( std::string alias )
Returns reference to the Header that contains a Sub with the given alias.
5.8.4 Member Data Documentation
5.8.4.1 std::map<std::string, Header> Document::Head_Map [private]
Map of headers contained within the document.
The documentation for this class was generated from the following file:
    • yaml_wrapper.h
5.9 DOGFISH_DATA Struct Reference
Primary data structure for running the DOGFISH application.
#include <dogfish.h>
```

Public Attributes

• unsigned long int total_steps = 0

Total number of solver steps taken.

• double time old = 0.0

Old value of time (hrs)

• double time = 0.0

Current value of time (hrs)

• bool Print2File = true

True = results to .txt; False = no printing.

• bool Print2Console = true

True = results to console; False = no printing.

• bool DirichletBC = false

False = uses film mass transfer for BC, True = Dirichlet BC.

bool NonLinear = false

False = Solve directly, True = Solve iteratively.

• double t_counter = 0.0

Counter for the time output.

double t_print

Print output at every t_print time (hrs)

int NumComp

Number of species to track.

• double end_time

Units: hours.

· double total sorption old

Per mass or volume of single fiber.

• double total_sorption

Per mass or volume of single fiber.

· double fiber_length

Units: um.

· double fiber diameter

Units: um.

· double fiber_specific_area

Units: m^2/kg .

FILE * OutputFile

Output file pointer to the output file for postprocesses and results.

• double(* eval_R)(int i, int I, const void *data)

Function pointer to evaluate retardation coefficient.

double(* eval_DI)(int i, int I, const void *data)

Function pointer to evaluate intraparticle diffusivity.

double(* eval_kf)(int i, const void *data)

Function pointer to evaluate film mass transfer coefficient.

• double(* eval_qs)(int i, const void *data)

Function pointer to evaluate fiber surface concentration.

const void * user_data

Data structure for users info to calculate all parameters.

std::vector< FINCH_DATA > finch_dat

Data structure for FINCH_DATA objects.

• std::vector< DOGFISH_PARAM > param_dat

Data structure for DOGFISH_PARAM objects.

5.9.1 Detailed Description

Primary data structure for running the DOGFISH application.

C-style object to hold information for the adsorption simulations. Contains function pointers and other data structures. This information is passed around to other functions used to simulate the fiber diffusion physics.

5.9.2 Member Data Documentation

5.9.2.1 unsigned long int DOGFISH_DATA::total_steps = 0

Total number of solver steps taken.

5.9.2.2 double DOGFISH_DATA::time_old = 0.0

Old value of time (hrs)

5.9.2.3 double DOGFISH_DATA::time = 0.0

Current value of time (hrs)

5.9.2.4 bool DOGFISH_DATA::Print2File = true

True = results to .txt; False = no printing.

5.9.2.5 bool DOGFISH_DATA::Print2Console = true

True = results to console; False = no printing.

5.9.2.6 bool DOGFISH_DATA::DirichletBC = false

False = uses film mass transfer for BC, True = Dirichlet BC.

5.9.2.7 bool DOGFISH_DATA::NonLinear = false

False = Solve directly, True = Solve iteratively.

5.9.2.8 double DOGFISH_DATA::t_counter = 0.0

Counter for the time output.

5.9.2.9 double DOGFISH_DATA::t_print

Print output at every t_print time (hrs)

5.9.2.10 int DOGFISH_DATA::NumComp

Number of species to track.

5.9.2.11 double DOGFISH_DATA::end_time

Units: hours.

5.9.2.12 double DOGFISH_DATA::total_sorption_old

Per mass or volume of single fiber.

5.9.2.13 double DOGFISH_DATA::total_sorption

Per mass or volume of single fiber.

5.9.2.14 double DOGFISH_DATA::fiber_length

Units: um.

5.9.2.15 double DOGFISH_DATA::fiber_diameter

Units: um.

5.9.2.16 double DOGFISH_DATA::fiber_specific_area

Units: m^2/kg.

5.9.2.17 FILE* DOGFISH_DATA::OutputFile

Output file pointer to the output file for postprocesses and results.

5.9.2.18 double(* DOGFISH_DATA::eval_R) (int i, int I, const void *data)

Function pointer to evaluate retardation coefficient.

5.9.2.19 double(* DOGFISH_DATA::eval_DI) (int i, int I, const void *data)

Function pointer to evaluate intraparticle diffusivity.

5.9.2.20 double(* DOGFISH_DATA::eval_kf) (int i, const void *data)

Function pointer to evaluate film mass transfer coefficient.

5.9.2.21 double(* DOGFISH_DATA::eval_qs) (int i, const void *data)

Function pointer to evaluate fiber surface concentration.

5.9.2.22 const void* DOGFISH_DATA::user_data

Data structure for users info to calculate all parameters.

5.9.2.23 std::vector<FINCH_DATA> DOGFISH_DATA::finch_dat

Data structure for FINCH_DATA objects.

5.9.2.24 std::vector<DOGFISH_PARAM> DOGFISH_DATA::param_dat

Data structure for DOGFISH_PARAM objects.

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

· dogfish.h

5.10 DOGFISH_PARAM Struct Reference

Data structure for species-specific parameters.

```
#include <dogfish.h>
```

Public Attributes

· double intraparticle_diffusion

Units: um^2/hr.

• double film_transfer_coeff

Units: um/hr.

• double surface_concentration

Units: mol/kg.

• double initial_sorption

Units: mol/kg.

• double sorbed_molefraction

Molefraction of sorbed species.

· Molecule species

Adsorbed species Molecule Object.

5.10.1 Detailed Description

Data structure for species-specific parameters.

C-style object to hold information on all adsorbing species. Parameters are given descriptive names to indicate what each is for.

5.10.2 Member Data Documentation

5.10.2.1 double DOGFISH_PARAM::intraparticle_diffusion

Units: um²/hr.

```
5.10.2.2 double DOGFISH_PARAM::film_transfer_coeff
Units: um/hr.
5.10.2.3 double DOGFISH_PARAM::surface_concentration
Units: mol/kg.
5.10.2.4 double DOGFISH_PARAM::initial_sorption
Units: mol/kg.
5.10.2.5 double DOGFISH_PARAM::sorbed_molefraction
Molefraction of sorbed species.
5.10.2.6 Molecule DOGFISH_PARAM::species
Adsorbed species Molecule Object.
The documentation for this struct was generated from the following file:
    · dogfish.h
5.11 FINCH_DATA Struct Reference
Data structure for the FINCH object.
#include <finch.h>
Public Attributes
    • int d = 0
          Dimension of the problem: 0 = cartesian, 1 = cylindrical, 2 = spherical.

    double dt = 0.0125

          Time step.
    • double dt_old = 0.0125
          Previous time step.
    • double T = 1.0
          Total time.
    • double dz = 0.1
          Space step.
    • double L = 1.0
          Total space.
    • double s = 1.0
          Char quantity (spherical = 1, cylindrical = length, cartesian = area)
    • double t = 0.0
          Current Time.

    double t_old = 0.0
```

5.11 FINCH DATA Struct Reference Previous Time. • double uT = 0.0Total amount of conserved quantity in domain. • double uT old = 0.0Old Total amount of conserved quantity. double uAvg = 0.0 Average amount of conserved quantity in domain. • double uAvg_old = 0.0 Old Average amount of conserved quantity. • double uIC = 0.0 Initial condition of Conserved Quantity (if constant) double vIC = 1.0 Initial condition of Velocity (if constant) • double **DIC** = 1.0 Initial condition of Dispersion (if constant) double kIC = 1.0 Initial condition of Reaction (if constant) • double RIC = 1.0 Initial condition of the Time Coefficient (if constant) • double <u>uo</u> = 1.0 Boundary Value of Conserved Quantity. • double vo = 1.0 Boundary Value of Velocity. • double **Do** = 1.0 Boundary Value of Dispersion.

• double **ko** = 1.0

Boundary Value of Reaction.

• double Ro = 1.0

Boundary Value of Time Coefficient.

• double kfn = 1.0

Film mass transfer coefficient Old.

• double kfnp1 = 1.0

Film mass transfer coefficient New.

· double lambda I

Boundary Coefficient for Implicit Neumann (Calculated at Runtime)

double lambda_E

Boundary Coefficient for Explicit Neumann (Calculated at Runtime)

• int LN = 10

Number of nodes.

bool CN = true

True if Crank-Nicholson, false if Implicit, never use explicit.

bool Update = false

Flag to check if the system needs updating.

bool Dirichlet = false

Flag to indicate use of Dirichlet or Neumann starting boundary.

• bool CheckMass = false

Flag to indicate whether or not mass is to be checked.

• bool ExplicitFlux = false

Flag to indicate whether or not to use fully explicit flux limiters.

• bool Iterative = true

Flag to indicate whether to solve directly, or iteratively.

• bool SteadyState = false Flag to determine whether or not to solve the steady-state problem. bool NormTrack = true Flag to determine whether or not to track the norms during simulation. • double beta = 0.5 Scheme type indicator: 0.5=CN & 1.0=Implicit; all else NULL. • double tol rel = 1e-6 Relative Tolerance for Convergence. double tol abs = 1e-6 Absolute Tolerance for Convergence. • int max iter = 20 Maximum number of iterations allowed. int total iter = 0 Total number of iterations made. int nl method = FINCH Picard Non-linear solution method - default = FINCH_Picard. std::vector< double > CL_I Left side, implicit coefficients (Calculated at Runtime) std::vector< double > CL E Left side, explicit coefficients (Calculated at Runtime) std::vector< double > CC_I Centered, implicit coefficients (Calculated at Runtime) std::vector< double > CC E Centered, explicit coefficients (Calculated at Runtime) std::vector< double > CR_I Right side, implicit coefficients (Calculated at Runtime) std::vector< double > CR E Right side, explicit coefficients (Calculated at Runtime) std::vector< double > fL_I Left side, implicit fluxes (Calculated at Runtime) std::vector< double > fL E Left side, explicit fluxes (Calculated at Runtime) std::vector< double > fC_I Centered, implicit fluxes (Calculated at Runtime) std::vector< double > fC E Centered, explicit fluxes (Calculated at Runtime) std::vector< double > fR | I Right side, implicit fluxes (Calculated at Runtime) std::vector< double > fR_E Right side, explicit fluxes (Calculated at Runtime) std::vector< double > OI Implicit upper diagonal matrix elements (Calculated at Runtime) std::vector< double > OE Explicit upper diagonal matrix elements (Calculated at Runtime) std::vector< double > NI Implicit diagonal matrix elements (Calculated at Runtime) std::vector< double > NE Explicit diagonal matrix elements (Calculated at Runtime) std::vector< double > MI Implicit lower diagonal matrix elements (Calculated at Runtime)

std::vector< double > ME

```
Explicit lower diagonal matrix elements (Calculated at Runtime)
std::vector< double > uz_l_l

    std::vector< double > uz_lm1_l

    std::vector< double > uz lp1 l

      Implicit local slopes (Calculated at Runtime)
std::vector< double > uz_I_E

    std::vector< double > uz lm1 E

    std::vector< double > uz_lp1_E

      Explicit local slopes (Calculated at Runtime)

    Matrix< double > unm1

      Conserved Quantity Older.

    Matrix< double > un

      Conserved Quantity Old.

    Matrix< double > unp1

      Conserved Quantity New.
• Matrix< double > u star
      Conserved Quantity Projected New.

    Matrix< double > ubest

     Best found solution if solving iteratively.

    Matrix< double > vn

      Velocity Old.

    Matrix< double > vnp1

      Velocity New.

    Matrix< double > Dn

      Dispersion Old.

    Matrix< double > Dnp1

     Dispersion New.

    Matrix< double > kn

      Reaction Old.

    Matrix< double > knp1

      Reaction New.
• Matrix< double > Sn
      Forcing Function Old.
• Matrix< double > Snp1
      Forcing Function New.
• Matrix< double > Rn
      Time Coeff Old.

    Matrix< double > Rnp1

      Time Coeff New.

    Matrix< double > Fn

      Flux Limiter Old.

    Matrix< double > Fnp1

     Flux Limiter New.

    Matrix< double > gl

      Implicit Side Boundary Conditions.

    Matrix< double > gE

      Explicit Side Boundary Conditions.
```

```
    Matrix< double > gE
        Explicit Side Boundary Conditions
    Matrix< double > res
        Current residual.
    Matrix< double > pres
        Current search direction.
```

int(* callroutine)(const void *user_data)

Function pointer to executioner (DEFAULT = default_execution)

int(* setic)(const void *user data)

Function pointer to initial conditions (DEFAULT = default_ic)

int(* settime)(const void *user data)

Function pointer to set time step (DEFAULT = default_timestep)

int(* setpreprocess)(const void *user_data)

Function pointer to preprocesses (DEFAULT = default_preprocess)

int(* solve)(const void *user data)

Function pointer to the solver (DEFAULT = default_solve)

int(* setparams)(const void *user_data)

Function pointer to set parameters (DEFAULT = default_params)

int(* discretize)(const void *user_data)

Function pointer to discretization (DEFAULT = ospre_discretization)

- int(* setbcs)(const void *user data)
- int(* evalres)(const Matrix< double > &x, Matrix< double > &res, const void *user_data)

Function pointer to the residual function (DEFAULT = default_res)

int(* evalprecon)(const Matrix < double > &b, Matrix < double > &p, const void *user_data)

Function pointer to the preconditioning function (DEFAULT = default_precon)

int(* setpostprocess)(const void *user_data)

Function pointer to the postprocesses (DEFAULT = default_postprocess)

int(* resettime)(const void *user_data)

Function pointer to reset time (DEFAULT = default_reset)

• PICARD_DATA picard_dat

Data structure for PICARD method (no need to use this)

PJFNK_DATA pjfnk_dat

Data structure for PJFNK method (more rigours method)

const void * param_data

User's data structure used to evaluate the parameter function (Must override if setparams is overriden)

5.11.1 Detailed Description

Data structure for the FINCH object.

C-style object that holds data, functions, and other structures necessary to discretize and solve a FINCH problem. All of this information must be overriden or initialized prior to running a FINCH simulation. Many, many default functions are provided to make it easier to incorporate FINCH into other problems. The main function to override will be the setparams function. This will be a function that the user provides to tell the FINCH simulation how the parameters of the problem vary in time and space and whether or not they are coupled the the variable u. All functions are overridable and several can be skipped entirely, or called directly at different times in the execution of a particular routine. This make FINCH extremely flexible to the user.

Note

All parameters and dimensions do not carry any units with them. The user is required to keep track of all their own units in their particular problem and ensure that units will cancel and be consistent in their own physical model.

5.11.2 Member Data Documentation

5.11.2.1 int FINCH_DATA::d = 0

Dimension of the problem: 0 = cartesian, 1 = cylindrical, 2 = spherical.

5.11.2.2 double FINCH_DATA::dt = 0.0125 Time step. 5.11.2.3 double FINCH_DATA::dt_old = 0.0125 Previous time step. 5.11.2.4 double FINCH_DATA::T = 1.0 Total time. 5.11.2.5 double FINCH_DATA::dz = 0.1 Space step. 5.11.2.6 double FINCH_DATA::L = 1.0 Total space. 5.11.2.7 double FINCH_DATA::s = 1.0 Char quantity (spherical = 1, cylindrical = length, cartesian = area) 5.11.2.8 double FINCH_DATA::t = 0.0 Current Time. 5.11.2.9 double FINCH_DATA::t_old = 0.0 Previous Time. 5.11.2.10 double FINCH_DATA::uT = 0.0 Total amount of conserved quantity in domain. 5.11.2.11 double FINCH_DATA::uT_old = 0.0 Old Total amount of conserved quantity. 5.11.2.12 double FINCH_DATA::uAvg = 0.0 Average amount of conserved quantity in domain.

Generated by Doxygen

5.11.2.13 double FINCH_DATA::uAvg_old = 0.0

Old Average amount of conserved quantity.

5.11.2.14 double FINCH_DATA::uIC = 0.0

Initial condition of Conserved Quantity (if constant)

5.11.2.15 double FINCH_DATA::vIC = 1.0

Initial condition of Velocity (if constant)

5.11.2.16 double FINCH_DATA::DIC = 1.0

Initial condition of Dispersion (if constant)

5.11.2.17 double FINCH_DATA::kIC = 1.0

Initial condition of Reaction (if constant)

5.11.2.18 double FINCH_DATA::RIC = 1.0

Initial condition of the Time Coefficient (if constant)

5.11.2.19 double FINCH_DATA::uo = 1.0

Boundary Value of Conserved Quantity.

5.11.2.20 double FINCH_DATA::vo = 1.0

Boundary Value of Velocity.

5.11.2.21 double FINCH_DATA::Do = 1.0

Boundary Value of Dispersion.

5.11.2.22 double FINCH_DATA::ko = 1.0

Boundary Value of Reaction.

5.11.2.23 double FINCH_DATA::Ro = 1.0

Boundary Value of Time Coefficient.

5.11.2.24 double FINCH_DATA::kfn = 1.0

Film mass transfer coefficient Old.

5.11.2.25 double FINCH_DATA::kfnp1 = 1.0

Film mass transfer coefficient New.

5.11.2.26 double FINCH_DATA::lambda_I

Boundary Coefficient for Implicit Neumann (Calculated at Runtime)

5.11.2.27 double FINCH_DATA::lambda_E

Boundary Coefficient for Explicit Neumann (Calculated at Runtime)

5.11.2.28 int FINCH_DATA::LN = 10

Number of nodes.

5.11.2.29 bool FINCH_DATA::CN = true

True if Crank-Nicholson, false if Implicit, never use explicit.

5.11.2.30 bool FINCH_DATA::Update = false

Flag to check if the system needs updating.

5.11.2.31 bool FINCH_DATA::Dirichlet = false

Flag to indicate use of Dirichlet or Neumann starting boundary.

5.11.2.32 bool FINCH_DATA::CheckMass = false

Flag to indicate whether or not mass is to be checked.

5.11.2.33 bool FINCH_DATA::ExplicitFlux = false

Flag to indicate whether or not to use fully explicit flux limiters.

5.11.2.34 bool FINCH_DATA::Iterative = true

Flag to indicate whether to solve directly, or iteratively.

5.11.2.35 bool FINCH_DATA::SteadyState = false

Flag to determine whether or not to solve the steady-state problem.

5.11.2.36 bool FINCH_DATA::NormTrack = true

Flag to determine whether or not to track the norms during simulation.

5.11.2.37 double FINCH_DATA::beta = 0.5

Scheme type indicator: 0.5=CN & 1.0=Implicit; all else NULL.

5.11.2.38 double FINCH_DATA::tol_rel = 1e-6

Relative Tolerance for Convergence.

5.11.2.39 double FINCH_DATA::tol_abs = 1e-6

Absolute Tolerance for Convergence.

5.11.2.40 int FINCH_DATA::max_iter = 20

Maximum number of iterations allowed.

5.11.2.41 int FINCH_DATA::total_iter = 0

Total number of iterations made.

5.11.2.42 int FINCH_DATA::nl_method = FINCH_Picard

Non-linear solution method - default = FINCH Picard.

5.11.2.43 std::vector<double> FINCH_DATA::CL_I

Left side, implicit coefficients (Calculated at Runtime)

5.11.2.44 std::vector<double> FINCH_DATA::CL_E

Left side, explicit coefficients (Calculated at Runtime)

5.11.2.45 std::vector<double> FINCH_DATA::CC_I

Centered, implicit coefficients (Calculated at Runtime)

 $5.11.2.46 \quad std::vector{<}double{>} FINCH_DATA::CC_E$

Centered, explicit coefficients (Calculated at Runtime)

 $5.11.2.47 \quad std::vector{<}double{>} FINCH_DATA::CR_I$

Right side, implicit coefficients (Calculated at Runtime)

5.11.2.48 std::vector<double> FINCH_DATA::CR_E

Right side, explicit coefficients (Calculated at Runtime)

 $5.11.2.49 \quad std::vector{<}double{>} FINCH_DATA::fL_I$

Left side, implicit fluxes (Calculated at Runtime)

5.11.2.50 std::vector<double> FINCH_DATA::fL_E Left side, explicit fluxes (Calculated at Runtime) $5.11.2.51 \quad std::vector{<}double{>} FINCH_DATA::fC_I$ Centered, implicit fluxes (Calculated at Runtime) 5.11.2.52 std::vector<double> FINCH_DATA::fC_E Centered, explicit fluxes (Calculated at Runtime) 5.11.2.53 std::vector<double> FINCH_DATA::fR_I Right side, implicit fluxes (Calculated at Runtime) 5.11.2.54 std::vector<double> FINCH_DATA::fR_E Right side, explicit fluxes (Calculated at Runtime) 5.11.2.55 std::vector<double> FINCH_DATA::OI Implicit upper diagonal matrix elements (Calculated at Runtime) $5.11.2.56 \quad std::vector{<}double{>} FINCH_DATA::OE$ Explicit upper diagonal matrix elements (Calculated at Runtime) 5.11.2.57 std::vector<double> FINCH_DATA::NI Implicit diagonal matrix elements (Calculated at Runtime) 5.11.2.58 std::vector<double> FINCH_DATA::NE Explicit diagonal matrix elements (Calculated at Runtime) 5.11.2.59 std::vector<double> FINCH_DATA::MI Implicit lower diagonal matrix elements (Calculated at Runtime) 5.11.2.60 std::vector<double> FINCH_DATA::ME Explicit lower diagonal matrix elements (Calculated at Runtime) 5.11.2.61 std::vector<double> FINCH_DATA::uz_I_I

Implicit local slopes (Calculated at Runtime)

5.11.2.62 std::vector<double> FINCH_DATA::uz_lm1_l

5.11.2.63 std::vector<double> FINCH_DATA::uz_lp1_l

5.11.2.64 std::vector<double> FINCH_DATA::uz_I_E 5.11.2.65 std::vector<double> FINCH_DATA::uz_lm1_E 5.11.2.66 std::vector<double> FINCH_DATA::uz_lp1_E Explicit local slopes (Calculated at Runtime) 5.11.2.67 Matrix < double > FINCH_DATA::unm1 Conserved Quantity Older. 5.11.2.68 Matrix < double > FINCH_DATA::un Conserved Quantity Old. 5.11.2.69 Matrix < double > FINCH_DATA::unp1 Conserved Quantity New. 5.11.2.70 Matrix<double> FINCH_DATA::u_star Conserved Quantity Projected New. 5.11.2.71 Matrix<double> FINCH_DATA::ubest Best found solution if solving iteratively. 5.11.2.72 Matrix<double> FINCH_DATA::vn Velocity Old. 5.11.2.73 Matrix < double > FINCH_DATA::vnp1 Velocity New. 5.11.2.74 Matrix < double > FINCH_DATA::Dn Dispersion Old. 5.11.2.75 Matrix<double> FINCH_DATA::Dnp1 Dispersion New. 5.11.2.76 Matrix<double> FINCH_DATA::kn

Reaction Old.

5.11.2.77 Matrix < double > FINCH_DATA::knp1

Reaction New.

5.11.2.78 Matrix < double > FINCH_DATA::Sn

Forcing Function Old.

 $5.11.2.79 \quad Matrix{<} double{>} FINCH_DATA::Snp1$

Forcing Function New.

5.11.2.80 Matrix < double > FINCH_DATA::Rn

Time Coeff Old.

5.11.2.81 Matrix < double > FINCH_DATA::Rnp1

Time Coeff New.

5.11.2.82 Matrix < double > FINCH_DATA::Fn

Flux Limiter Old.

5.11.2.83 Matrix < double > FINCH_DATA::Fnp1

Flux Limiter New.

5.11.2.84 Matrix<double> FINCH_DATA::gl

Implicit Side Boundary Conditions.

5.11.2.85 Matrix < double > FINCH_DATA::gE

Explicit Side Boundary Conditions.

5.11.2.86 Matrix < double > FINCH_DATA::res

Current residual.

5.11.2.87 Matrix<double> FINCH_DATA::pres

Current search direction.

5.11.2.88 int(* FINCH_DATA::callroutine) (const void *user_data)

Function pointer to executioner (DEFAULT = default_execution)

```
5.11.2.89 int(* FINCH_DATA::setic) (const void *user_data)
Function pointer to initial conditions (DEFAULT = default_ic)
5.11.2.90 int(* FINCH_DATA::settime) (const void *user_data)
Function pointer to set time step (DEFAULT = default_timestep)
5.11.2.91 int(* FINCH_DATA::setpreprocess) (const void *user_data)
Function pointer to preprocesses (DEFAULT = default_preprocess)
5.11.2.92 int(* FINCH_DATA::solve) (const void *user_data)
Function pointer to the solver (DEFAULT = default_solve)
5.11.2.93 int(* FINCH_DATA::setparams) (const void *user_data)
Function pointer to set parameters (DEFAULT = default params)
5.11.2.94 int(* FINCH_DATA::discretize) (const void *user_data)
Function pointer to discretization (DEFAULT = ospre_discretization)
5.11.2.95 int(* FINCH_DATA::setbcs) (const void *user_data)
Function pointer to set boundary conditions (DEFAULT = default_bcs)
5.11.2.96 int(* FINCH_DATA::evalres) (const Matrix < double > &x, Matrix < double > &res, const void *user_data)
Function pointer to the residual function (DEFAULT = default_res)
5.11.2.97 int(* FINCH_DATA::evalprecon) (const Matrix < double > &b, Matrix < double > &p, const void *user_data)
Function pointer to the preconditioning function (DEFAULT = default_precon)
5.11.2.98 int(* FINCH_DATA::setpostprocess) (const void *user_data)
Function pointer to the postprocesses (DEFAULT = default_postprocess)
5.11.2.99 int(* FINCH_DATA::resettime) (const void *user_data)
Function pointer to reset time (DEFAULT = default_reset)
5.11.2.100 PICARD_DATA FINCH_DATA::picard_dat
Data structure for PICARD method (no need to use this)
```

```
5.11.2.101 PJFNK_DATA FINCH_DATA::pjfnk_dat
```

Data structure for PJFNK method (more rigours method)

```
5.11.2.102 const void* FINCH_DATA::param_data
```

User's data structure used to evaluate the parameter function (Must override if setparams is overriden)

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

· finch.h

5.12 GCR_DATA Struct Reference

Data structure for the implementation of the GCR algorithm for non-symmetric linear systems.

```
#include <lark.h>
```

Public Attributes

• int restart = -1

Restart parameter for outer iterations - default = 20.

• int maxit = 0

Maximum allowable outer iterations.

• int iter_outer = 0

Number of outer iterations taken.

• int iter_inner = 0

Number of inner iterations taken.

• int total iter = 0

Total number of iterations taken.

bool breakdown = false

Boolean to determine if a step has failed.

· double alpha

Inner iteration step size.

• double beta

Outer iteration step size.

• double tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

• double tol_abs = 1e-6

Absolute tolerance for convergence - default = 1e-6.

• double res

Absolute residual norm for linear system.

· double relres

Relative residual norm for linear system.

double relres_base

Initial residual norm of the linear system.

· double bestres

Best found residual norm of the linear system.

bool Output = true

True = print messages to the console.

Matrix< double > x

Current solution to the linear system.

Matrix< double > bestx

Best found solution to the linear system.

• Matrix< double > r

Residual Vector.

Matrix< double > c temp

Temporary c vector to be updated.

Matrix< double > u_temp

Temporary u vector to be updated.

std::vector< Matrix< double > > u

Vector span for updating x.

std::vector< Matrix< double > > c

Vector span for updating r.

OPTRANS_DATA transpose_dat

Data structure for Operator Transposition.

5.12.1 Detailed Description

Data structure for the implementation of the GCR algorithm for non-symmetric linear systems.

C-style object used in conjunction with the Generalized Conjugate Residual (GCR) algorithm for solving a non-symmetric linear system of equations. When the linear system in question has a positive-definite-symmetric component to it, then this algorithm is equivalent to GMRESRP. However, it is generally less efficient than GMRESRP and can suffer breakdowns.

```
5.12.2 Member Data Documentation
```

5.12.2.1 int GCR_DATA::restart = -1

Restart parameter for outer iterations - default = 20.

5.12.2.2 int GCR_DATA::maxit = 0

Maximum allowable outer iterations.

5.12.2.3 int GCR_DATA::iter_outer = 0

Number of outer iterations taken.

5.12.2.4 int GCR_DATA::iter_inner = 0

Number of inner iterations taken.

5.12.2.5 int GCR_DATA::total_iter = 0

Total number of iterations taken.

5.12.2.6 bool GCR_DATA::breakdown = false

Boolean to determine if a step has failed.

5.12.2.7 double GCR_DATA::alpha

Inner iteration step size.

5.12.2.8 double GCR_DATA::beta

Outer iteration step size.

5.12.2.9 double GCR_DATA::tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

5.12.2.10 double GCR_DATA::tol_abs = 1e-6

Absolute tolerance for convergence - default = 1e-6.

5.12.2.11 double GCR_DATA::res

Absolute residual norm for linear system.

5.12.2.12 double GCR_DATA::relres

Relative residual norm for linear system.

5.12.2.13 double GCR_DATA::relres_base

Initial residual norm of the linear system.

5.12.2.14 double GCR_DATA::bestres

Best found residual norm of the linear system.

5.12.2.15 bool GCR_DATA::Output = true

True = print messages to the console.

5.12.2.16 Matrix<double> GCR_DATA::x

Current solution to the linear system.

5.12.2.17 Matrix < double > GCR_DATA::bestx

Best found solution to the linear system.

```
5.12.2.18 Matrix<double> GCR_DATA::r
```

Residual Vector.

```
5.12.2.19 Matrix<double> GCR_DATA::c_temp
```

Temporary c vector to be updated.

```
5.12.2.20 Matrix < double > GCR_DATA::u_temp
```

Temporary u vector to be updated.

```
5.12.2.21 std::vector<Matrix<double>> GCR_DATA::u
```

Vector span for updating x.

```
5.12.2.22 std::vector<Matrix<double>> GCR_DATA::c
```

Vector span for updating r.

```
5.12.2.23 OPTRANS_DATA GCR_DATA::transpose_dat
```

Data structure for Operator Transposition.

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

· lark.h

5.13 GMRESLP_DATA Struct Reference

Data structure for implementation of the Restarted GMRES algorithm with Left Preconditioning.

```
#include <lark.h>
```

Public Attributes

```
• int restart = -1
```

Restart parameter - default = min(vector_size,20)

• int maxit = 0

Maximum allowable iterations - default = min(vector_size,1000)

• int iter = 0

Number of iterations needed for convergence.

• int steps = 0

Total number of gmres iterations and krylov iterations.

• double tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

• double tol abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

double res

Absolution redisual norm of the linear system.

· double relres

Relative residual norm of the linear system.

· double relres base

Initial residual norm of the linear system.

· double bestres

Best found residual norm of the linear system.

bool Output = true

True = print messages to console.

Matrix< double > x

Current solution to the linear system.

Matrix< double > bestx

Best found solution to the linear system.

• Matrix< double > r

Residual vector for the linear system.

ARNOLDI_DATA arnoldi_dat

Data structure for the kyrlov subspace.

5.13.1 Detailed Description

Data structure for implementation of the Restarted GMRES algorithm with Left Preconditioning.

C-style object used in conjunction with Generalized Minimum RESidual Left-Precondtioned (GMRESLP) and Full Orthogonalization Method (FOM) algorithms to iteratively or directly solve a linear system of equations. When using with GMRESLP, you can only check/observe the linear residuals before a restart or after the Arnoldi space is constructed. This is because this object uses Left-side Preconditioning. A faster routine may be GMRESRP, which is able to construct residuals after each Arnoldi iteration.

```
5.13.2 Member Data Documentation
```

```
5.13.2.1 int GMRESLP DATA::restart = -1
```

Restart parameter - default = min(vector_size,20)

5.13.2.2 int GMRESLP_DATA::maxit = 0

Maximum allowable iterations - default = min(vector_size,1000)

5.13.2.3 int GMRESLP_DATA::iter = 0

Number of iterations needed for convergence.

5.13.2.4 int GMRESLP_DATA::steps = 0

Total number of gmres iterations and krylov iterations.

5.13.2.5 double GMRESLP_DATA::tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

5.13.2.6 double GMRESLP_DATA::tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

5.13.2.7 double GMRESLP_DATA::res

Absolution redisual norm of the linear system.

5.13.2.8 double GMRESLP_DATA::relres

Relative residual norm of the linear system.

5.13.2.9 double GMRESLP_DATA::relres_base

Initial residual norm of the linear system.

5.13.2.10 double GMRESLP_DATA::bestres

Best found residual norm of the linear system.

5.13.2.11 bool GMRESLP_DATA::Output = true

True = print messages to console.

5.13.2.12 Matrix < double > GMRESLP_DATA::x

Current solution to the linear system.

 $5.13.2.13 \quad \textbf{Matrix}{<} \textbf{double}{>} \textbf{GMRESLP_DATA}{::} \textbf{bestx}$

Best found solution to the linear system.

5.13.2.14 Matrix < double > GMRESLP_DATA::r

Residual vector for the linear system.

5.13.2.15 ARNOLDI_DATA GMRESLP_DATA::arnoldi_dat

Data structure for the kyrlov subspace.

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

• lark.h

5.14 GMRESR_DATA Struct Reference

Data structure for the implementation of GCR with Nested GMRES preconditioning (i.e., GMRESR)

#include <lark.h>

Public Attributes

```
int gcr_restart = -1
```

Number of GCR restarts (default = 20, max = N)

• int gcr maxit = 0

Number of GCR iterations.

• int gmres_restart = -1

Number of GMRES restarts (max = 20)

• int gmres maxit = 1

Number of GMRES iterations (max = 5, default = 1)

int N

Dimension of the linear system.

· int total iter

Total GMRES and GCR iterations.

· int iter outer

Total GCR iterations.

· int iter inner

Total GMRES iterations.

• bool GCR_Output = true

True = print GCR messages.

• bool GMRES_Output = false

True = print GMRES messages.

• double gmres_tol = 0.1

Tolerance relative to GCR iterations.

• double gcr_rel_tol = 1e-6

Relative outer residual tolerance.

• double gcr_abs_tol = 1e-6

Absolute outer residual tolerance.

Matrix< double > arg

Argument matrix passed between preconditioner and iterator.

GCR_DATA gcr_dat

Data structure for the outer GCR steps.

• GMRESRP_DATA gmres_dat

Data structure for the inner GMRES steps.

int(* matvec)(const Matrix < double > &x, Matrix < double > &Ax, const void *matvec_data)

User supplied matrix-vector product function.

int(* terminal_precon)(const Matrix< double > &r, Matrix< double > &p, const void *precon_data)

Optional user supplied terminal preconditioner.

const void * matvec_data

Data structure for the user's matvec function.

const void * term_precon

Data structure for the user's terminal preconditioner.

5.14.1 Detailed Description

Data structure for the implementation of GCR with Nested GMRES preconditioning (i.e., GMRESR)

C-style object to be used in conjunction with the Generalized Minimum RESidual Recurive (GMRESR) algorithm. Although the name suggests that this method used GMRES recursively, what it is actually doing is nesting GMRE \leftarrow SRP iterations inside the GCR method to form a preconditioner for GCR. The name GMRESR came from literature (Vorst and Vuik, "GMRESR: A family of nested GMRES methods", 1991).

5.14.2 Member Data Documentation

5.14.2.1 int GMRESR_DATA::gcr_restart = -1

Number of GCR restarts (default = 20, max = N)

5.14.2.2 int GMRESR_DATA::gcr_maxit = 0

Number of GCR iterations.

5.14.2.3 int GMRESR_DATA::gmres_restart = -1

Number of GMRES restarts (max = 20)

5.14.2.4 int GMRESR_DATA::gmres_maxit = 1

Number of GMRES iterations (max = 5, default = 1)

5.14.2.5 int GMRESR_DATA::N

Dimension of the linear system.

5.14.2.6 int GMRESR_DATA::total_iter

Total GMRES and GCR iterations.

5.14.2.7 int GMRESR_DATA::iter_outer

Total GCR iterations.

5.14.2.8 int GMRESR_DATA::iter_inner

Total GMRES iterations.

5.14.2.9 bool GMRESR_DATA::GCR_Output = true

True = print GCR messages.

5.14.2.10 bool GMRESR_DATA::GMRES_Output = false

True = print GMRES messages.

5.14.2.11 double GMRESR_DATA::gmres_tol = 0.1

Tolerance relative to GCR iterations.

5.14.2.12 double GMRESR_DATA::gcr_rel_tol = 1e-6

Relative outer residual tolerance.

5.14.2.13 double GMRESR_DATA::gcr_abs_tol = 1e-6

Absolute outer residual tolerance.

5.14.2.14 Matrix < double > GMRESR_DATA::arg

Argument matrix passed between preconditioner and iterator.

5.14.2.15 GCR_DATA GMRESR_DATA::gcr_dat

Data structure for the outer GCR steps.

5.14.2.16 GMRESRP_DATA GMRESR_DATA::gmres_dat

Data structure for the inner GMRES steps.

 $5.14.2.17 \quad \text{int} (* \text{ GMRESR_DATA}:: \text{matvec}) \ (\text{const Matrix} < \text{double} > \&x, \\ \text{Matrix} < \text{double} > \&Ax, \\ \text{const void } * \text{matvec_data})$

User supplied matrix-vector product function.

5.14.2.18 int(* GMRESR_DATA::terminal_precon) (const Matrix < double > &r, Matrix < double > &p, const void *precon_data)

Optional user supplied terminal preconditioner.

5.14.2.19 const void* GMRESR_DATA::matvec_data

Data structure for the user's matvec function.

5.14.2.20 const void* GMRESR_DATA::term_precon

Data structure for the user's terminal preconditioner.

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

· lark.h

5.15 GMRESRP_DATA Struct Reference

Data structure for the Restarted GMRES algorithm with Right Preconditioning.

#include <lark.h>

Public Attributes

• int restart = -1

Restart parameter - default = min(20,vector_size)

• int maxit = 0

Maximum allowable outer iterations.

• int iter outer = 0

Total number of outer iterations.

• int iter inner = 0

Total number of inner iterations.

• int iter_total = 0

Total number of overall iterations.

• double tol rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

double tol_abs = 1e-6

Absolute tolerance for convergence - default = 1e-6.

• double res

Absolute residual norm for linear system.

· double relres

Relative residual norm for linear system.

· double relres_base

Initial residual norm of the linear system.

· double bestres

Best found residual norm of the linear system.

• bool Output = true

True = print messages to console.

Matrix< double > x

Current solution to the linear system.

Matrix< double > bestx

Best found solution to the linear system.

Matrix< double > r

Residual vector for the linear system.

std::vector< Matrix< double > > Vk

(N x k) orthonormal vector basis

std::vector< Matrix< double >> Zk

(N x k) preconditioned vector set

std::vector< std::vector< double > > H

(k+1 x k) upper Hessenberg storage matrix

• $std::vector < std::vector < double >> H_bar$

(k+1 x k) Factorized matrix

std::vector< double > y

(k x 1) Vector search direction

std::vector< double > e0

(k+1 x 1) Normalized vector with residual info

• $std::vector < double > e0_bar$

(k+1 x 1) Factorized normal vector

Matrix< double > w

(N) x (1) interim result of the matrix_vector multiplication

Matrix< double > v

(N) x (1) holding cell for the column entries of Vk and other interims

Matrix< double > sum

(N) x (1) running sum of subspace vectors for use in altering w

5.15.1 Detailed Description

Data structure for the Restarted GMRES algorithm with Right Preconditioning.

C-style object used in conjunction with Generalized Minimum RESidual Right Preconditioned (GMRESRP) algorithm to iteratively solve a linear system of equations. Unlike GMRESLP, the GMRESRP method is capable of checking linear residuals at both the inner and outer steps. As a result, this algorithm may terminate earlier than GMRESLP if it has found a suitable solution during one of the inner steps.

5.15.2 Member Data Documentation

5.15.2.1 int GMRESRP_DATA::restart = -1

Restart parameter - default = min(20,vector_size)

5.15.2.2 int GMRESRP_DATA::maxit = 0

Maximum allowable outer iterations.

5.15.2.3 int GMRESRP_DATA::iter_outer = 0

Total number of outer iterations.

5.15.2.4 int GMRESRP_DATA::iter_inner = 0

Total number of inner iterations.

5.15.2.5 int GMRESRP_DATA::iter_total = 0

Total number of overall iterations.

5.15.2.6 double GMRESRP_DATA::tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

5.15.2.7 double GMRESRP_DATA::tol_abs = 1e-6

Absolute tolerance for convergence - default = 1e-6.

5.15.2.8 double GMRESRP_DATA::res

Absolute residual norm for linear system.

5.15.2.9 double GMRESRP_DATA::relres

Relative residual norm for linear system.

5.15.2.10 double GMRESRP_DATA::relres_base

Initial residual norm of the linear system.

5.15.2.11 double GMRESRP_DATA::bestres

Best found residual norm of the linear system.

5.15.2.12 bool GMRESRP_DATA::Output = true

True = print messages to console.

5.15.2.13 Matrix < double > GMRESRP_DATA::x

Current solution to the linear system.

5.15.2.14 Matrix < double > GMRESRP_DATA::bestx

Best found solution to the linear system.

5.15.2.15 Matrix < double > GMRESRP_DATA::r

Residual vector for the linear system.

5.15.2.16 std::vector< Matrix<double> > GMRESRP_DATA::Vk

(N x k) orthonormal vector basis

5.15.2.17 std::vector< Matrix<double>> GMRESRP_DATA::Zk

 $(N \times k)$ preconditioned vector set

5.15.2.18 std::vector< std::vector< double >> GMRESRP_DATA::H

(k+1 x k) upper Hessenberg storage matrix

 ${\tt 5.15.2.19} \quad {\tt std::vector}{< \tt std::vector}{< \tt double} > > {\tt GMRESRP_DATA::H_bar}$

(k+1 x k) Factorized matrix

5.15.2.20 std::vector< double > GMRESRP_DATA::y

(k x 1) Vector search direction

 ${\it 5.15.2.21 \quad std::} vector < double > {\it GMRESRP_DATA::e0}$

(k+1 x 1) Normalized vector with residual info

```
5.15.2.22 std::vector< double > GMRESRP_DATA::e0_bar

(k+1 x 1) Factorized normal vector

5.15.2.23 Matrix<double> GMRESRP_DATA::w

(N) x (1) interim result of the matrix_vector multiplication

5.15.2.24 Matrix<double> GMRESRP_DATA::v

(N) x (1) holding cell for the column entries of Vk and other interims

5.15.2.25 Matrix<double> GMRESRP_DATA::sum
```

(N) x (1) running sum of subspace vectors for use in altering w

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

· lark.h

5.16 GPAST_DATA Struct Reference

GPAST Data Structure.

```
#include <magpie.h>
```

Public Attributes

• double x

Adsorbed mole fraction.

· double y

Gas phase mole fraction.

double He

Henry's Coefficient (mol/kg/kPa)

• double q

Amount adsorbed for each component (mol/kg)

std::vector< double > gama_inf

Infinite dilution activities.

• double qo

Pure component capacities (mol/kg)

• double Plo

Pure component spreading pressures (mol/kg)

std::vector< double > po

Pure component reference state pressures (kPa)

double poi

Reference state pressures solved for using Recover eval GPAST.

· bool present

If true, then the component is present; if false, then the component is not present.

5.16.1 Detailed Description

GPAST Data Structure.

C-style object holding all parameter information associated with the Generalized Predictive Adsorbed Solution Theory (GPAST) system of equations. Each species in the gas phase will have one of these objects.

5.16.2 Member Data Documentation

5.16.2.1 double GPAST_DATA::x

Adsorbed mole fraction.

5.16.2.2 double GPAST_DATA::y

Gas phase mole fraction.

5.16.2.3 double GPAST_DATA::He

Henry's Coefficient (mol/kg/kPa)

5.16.2.4 double GPAST_DATA::q

Amount adsorbed for each component (mol/kg)

5.16.2.5 std::vector<double> GPAST_DATA::gama_inf

Infinite dilution activities.

5.16.2.6 double GPAST_DATA::qo

Pure component capacities (mol/kg)

5.16.2.7 double GPAST_DATA::Plo

Pure component spreading pressures (mol/kg)

 $5.16.2.8 \quad std::vector < double > GPAST_DATA::po$

Pure component reference state pressures (kPa)

5.16.2.9 double GPAST_DATA::poi

Reference state pressures solved for using Recover eval GPAST.

5.16.2.10 bool GPAST_DATA::present

If true, then the component is present; if false, then the component is not present.

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

• magpie.h

5.17 GSTA_DATA Struct Reference

GSTA Data Structure.

```
#include <magpie.h>
```

Public Attributes

· double qmax

Theoretical maximum capacity of adsorbate-adsorbent pair (mol/kg)

• int m

Number of parameters in the GSTA isotherm.

std::vector< double > dHo

Enthalpies for each site (J/mol)

std::vector< double > dSo

Entropies for each site (J/(K*mol))

5.17.1 Detailed Description

GSTA Data Structure.

C-style object holding all parameter information associated with the Generalized Statistical Thermodynamic Adsorption (GSTA) isotherm model. Each species in the gas phase will have one of these objects.

5.17.2 Member Data Documentation

5.17.2.1 double GSTA_DATA::qmax

Theoretical maximum capacity of adsorbate-adsorbent pair (mol/kg)

5.17.2.2 int GSTA_DATA::m

Number of parameters in the GSTA isotherm.

5.17.2.3 std::vector<double> GSTA_DATA::dHo

Enthalpies for each site (J/mol)

```
5.17.2.4 std::vector<double> GSTA_DATA::dSo
```

Entropies for each site (J/(K*mol))

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

· magpie.h

5.18 GSTA_OPT_DATA Struct Reference

Data structure used in the GSTA optimization routines.

```
#include <gsta_opt.h>
```

Public Attributes

· int total_eval

Keeps track of the total number of function evaluations.

int n par

Number of parameters being optimized for.

· double qmax

Maximum theoretical adsorption capacity (M/M) (0 if unknown)

• int iso

 ${\it Keeps is otherm that is currently being optimized.}$

std::vector< std::vector< double > > Fobj

Creates a dynamic array to store all Fobj values.

- std::vector< std::vector< double > > q
- std::vector< std::vector< double >> P

Creates a dynamic array for q and P data pairs.

std::vector< std::vector< double > > best_par

Used to store the values of the parameters of best fit.

std::vector< std::vector< double > > Kno

Dimensionless parameters determined from best_par.

std::vector< std::vector< double >> > all_pars

Used to create a ragged array of all parameters.

• std::vector < std::vector < double >> norms

Used to store the values of all the calculated norms.

• $std::vector < double > opt_qmax$

If qmax is unknown, this vector holds it's optimized values.

5.18.1 Detailed Description

Data structure used in the GSTA optimization routines.

C-style structure that keeps track of all infomation during the optimization routine. All solutions and parameters to the GSTA isotherm are held in order to find the best solution with the fewest parameters.

5.18.2 Member Data Documentation

5.18.2.1 int GSTA_OPT_DATA::total_eval

Keeps track of the total number of function evaluations.

5.18.2.2 int GSTA_OPT_DATA::n_par

Number of parameters being optimized for.

5.18.2.3 double GSTA_OPT_DATA::qmax

Maximum theoretical adsorption capacity (M/M) (0 if unknown)

5.18.2.4 int GSTA_OPT_DATA::iso

Keeps isotherm that is currently being optimized.

5.18.2.5 std::vector<std::vector<double>> GSTA_OPT_DATA::Fobj

Creates a dynamic array to store all Fobj values.

5.18.2.6 std::vector<std::vector<double>> GSTA_OPT_DATA::q

5.18.2.7 std::vector<std::vector<double>> GSTA_OPT_DATA::P

Creates a dynamic array for q and P data pairs.

 ${\tt 5.18.2.8 \quad std::vector}{<} {\tt std::vector}{<} {\tt double}{\gt} {\gt} {\tt GSTA_OPT_DATA::best_par}$

Used to store the values of the parameters of best fit.

 ${\tt 5.18.2.9 \quad std::vector}{<} {\tt std::vector}{<} {\tt double}{\gt} {\gt} {\tt GSTA_OPT_DATA::Kno}$

Dimensionless parameters determined from best_par.

 $5.18.2.10 \quad std::vector < std::vector < double > > GSTA_OPT_DATA::all_pars$

Used to create a ragged array of all parameters.

5.18.2.11 std::vector<std::vector<double>> GSTA_OPT_DATA::norms

Used to store the values of all the calculated norms.

5.18.2.12 std::vector<double> GSTA_OPT_DATA::opt_qmax

If qmax is unknown, this vector holds it's optimized values.

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

• gsta_opt.h

5.19 Header Class Reference

Object for headers in a yaml document (inherits from SubHeader)

#include <yaml_wrapper.h>

Inheritance diagram for Header:



Public Member Functions

· Header ()

Default Constructor.

∼Header ()

Default Destructor.

• Header (const Header &head)

Copy constructor.

Header (std::string name)

Constructor by header name.

Header (const KeyValueMap &map)

Constructor by existing map.

Header (std::string name, const KeyValueMap &map)

Constructor by name and map.

• Header (std::string key, const SubHeader &sub)

Constructor by single subheader object.

Header & operator= (const Header &head)

Equals overload.

ValueTypePair & operator[] (const std::string key)

Return the ValueType reference at the given key.

ValueTypePair operator[] (const std::string key) const

Return the ValueType at the given key.

SubHeader & operator() (const std::string key)

Return the SubHeader reference at the given key.

• SubHeader operator() (const std::string key) const

Return the SubHeader at the given key.

std::map< std::string, SubHeader > & getSubMap ()

Return the reference to the SubHeader Map.

KeyValueMap & getDataMap ()

Return the reference to the KeyValueMap.

• SubHeader & getSubHeader (std::string key)

Return the subheader at the given key.

std::map< std::string, SubHeader >::const_iterator end () const

Returns a const iterator pointing to the end of the list.

std::map< std::string, SubHeader >::iterator end ()

Returns an iterator pointing to the end of the list.

• std::map< std::string, SubHeader >::const_iterator begin () const

Returns a const iterator pointing to the begining of the list.

• std::map< std::string, SubHeader >::iterator begin ()

Returns an iterator pointing to the begining of the list.

• void clear ()

Clear out the SubMap, KeyValueMap, and other info.

void resetKeys ()

Reset the keys of the SubMap to the names of each SubHeader.

void changeKey (std::string oldKey, std::string newKey)

Change one of the keys in the map.

void addPair (std::string key, std::string val)

Adds a pair object to the map (with only strings)

void addPair (std::string key, std::string val, int t)

Adds a pair object and asserts a type.

void setName (std::string name)

Set the name of the Header.

· void setAlias (std::string alias)

Set the alias of the header, if any.

• void setNameAliasPair (std::string n, std::string a, int s)

Set the name, alias, and state for the header.

void setState (int state)

Set the state of the header, if any.

• void DisplayContents ()

Display the contents of the header object.

void addSubKey (std::string key)

Adds a key to the SubHeader Map.

void copyAnchor2Alias (std::string alias, SubHeader &ref)

Find the anchor in the map, and copy to the Header reference given.

• int size ()

Return the size of the Sub_Map.

• std::string getName ()

Return the name of the header.

• std::string getAlias ()

Return the alias of the header.

• int getState ()

Return the state of the header.

• bool isAlias ()

Returns true if the header is an alias.

• bool isAnchor ()

Returns true if the header is an anchor.

SubHeader & getAnchoredSub (std::string alias)

Returns reference to the anchored subheader, if any.

Private Attributes

std::map< std::string, SubHeader > Sub Map

Map of the contained subheaders in the main header.

Additional Inherited Members

5.19.1 Detailed Description

Equals overload.

Object for headers in a yaml document (inherits from SubHeader)

C++ Object for headers in a yaml document that is built from the SubHeader object already created. The chain of inheritance works in this direction because a Header can have both a map of SubHeaders and a map of KeyValue ← Pairs. Therefore, the SubHeader object is actually the more generic form of a header.

Since this object inherits from SubHeader, it has access to all it's protected members, including the alias, state, name, and KeyValueMap. Operator overloads and other functions are provided to allow the user to query both the KeyValueMap and SubHeader for specific information. The names of the SubHeaders are also used as it's keys. Make sure all SubHeader keys are unique to this header.

```
5.19.2 Constructor & Destructor Documentation
5.19.2.1 Header::Header ( )
Default Constructor.
5.19.2.2 Header:: ∼ Header ( )
Default Destructor.
5.19.2.3 Header::Header ( const Header & head )
Copy constructor.
5.19.2.4 Header::Header ( std::string name )
Constructor by header name.
5.19.2.5 Header::Header (const KeyValueMap & map)
Constructor by existing map.
5.19.2.6 Header::Header ( std::string name, const KeyValueMap & map )
Constructor by name and map.
5.19.2.7 Header::Header ( std::string key, const SubHeader & sub )
Constructor by single subheader object.
5.19.3 Member Function Documentation
5.19.3.1 Header & Header::operator= ( const Header & head )
```

```
5.19.3.2 ValueTypePair& Header::operator[]( const std::string key )
Return the ValueType reference at the given key.
5.19.3.3 ValueTypePair Header::operator[]( const std::string key ) const
Return the ValueType at the given key.
5.19.3.4 SubHeader& Header::operator() ( const std::string key )
Return the SubHeader reference at the given key.
5.19.3.5 SubHeader Header::operator() ( const std::string key ) const
Return the SubHeader at the given key.
5.19.3.6 std::map<std::string, SubHeader>& Header::getSubMap ( )
Return the reference to the SubHeader Map.
5.19.3.7 KeyValueMap& Header::getDataMap ( )
Return the reference to the KeyValueMap.
5.19.3.8 SubHeader& Header::getSubHeader ( std::string key )
Return the subheader at the given key.
5.19.3.9 std::map<std::string, SubHeader>::const_iterator Header::end ( ) const
Returns a const iterator pointing to the end of the list.
5.19.3.10 std::map<std::string, SubHeader>::iterator Header::end ( )
Returns an iterator pointing to the end of the list.
5.19.3.11 std::map<std::string, SubHeader>::const_iterator Header::begin ( ) const
Returns a const iterator pointing to the begining of the list.
5.19.3.12 std::map<std::string, SubHeader>::iterator Header::begin ( )
Returns an iterator pointing to the begining of the list.
5.19.3.13 void Header::clear ( )
Clear out the SubMap, KeyValueMap, and other info.
```

```
5.19.3.14 void Header::resetKeys ( )
Reset the keys of the SubMap to the names of each SubHeader.
5.19.3.15 void Header::changeKey ( std::string oldKey, std::string newKey )
Change one of the keys in the map.
5.19.3.16 void Header::addPair ( std::string key, std::string val )
Adds a pair object to the map (with only strings)
5.19.3.17 void Header::addPair ( std::string key, std::string val, int t )
Adds a pair object and asserts a type.
5.19.3.18 void Header::setName ( std::string name )
Set the name of the Header.
5.19.3.19 void Header::setAlias ( std::string alias )
Set the alias of the header, if any.
5.19.3.20 void Header::setNameAliasPair ( std::string n, std::string a, int s )
Set the name, alias, and state for the header.
5.19.3.21 void Header::setState (int state)
Set the state of the header, if any.
5.19.3.22 void Header::DisplayContents ( )
Display the contents of the header object.
5.19.3.23 void Header::addSubKey ( std::string key )
Adds a key to the SubHeader Map.
5.19.3.24 void Header::copyAnchor2Alias ( std::string alias, SubHeader & ref )
Find the anchor in the map, and copy to the Header reference given.
5.19.3.25 int Header::size ( )
Return the size of the Sub_Map.
```

```
5.19.3.26 std::string Header::getName ( )
Return the name of the header.
5.19.3.27 std::string Header::getAlias ( )
Return the alias of the header.
5.19.3.28 int Header::getState ( )
Return the state of the header.
5.19.3.29 bool Header::isAlias ( )
Returns true if the header is an alias.
5.19.3.30 bool Header::isAnchor()
Returns true if the header is an anchor.
5.19.3.31 SubHeader& Header::getAnchoredSub ( std::string alias )
Returns reference to the anchored subheader, if any.
5.19.4 Member Data Documentation
5.19.4.1 std::map<std::string, SubHeader> Header::Sub_Map [private]
Map of the contained subheaders in the main header.
The documentation for this class was generated from the following file:
    • yaml_wrapper.h
5.20 KeyValueMap Class Reference
Key-Value-Type Map object creating a map of the KeyValuePair objects.
#include <yaml_wrapper.h>
```

Public Member Functions

KeyValueMap ()

Default constructor.

∼KeyValueMap ()

Default destructor.

KeyValueMap (const std::map< std::string, std::string > &map)

Construct from a map of strings.

KeyValueMap (std::string key, std::string value)

Construct one element in the map.

KeyValueMap (const KeyValueMap &map)

Copy constructor.

KeyValueMap & operator= (const KeyValueMap &map)

Equals overload.

ValueTypePair & operator[] (const std::string key)

Return the ValueType reference at the given key.

ValueTypePair operator[] (const std::string key) const

Return the ValueType at the give key.

std::map< std::string, ValueTypePair > & getMap ()

Return a reference to the Key_Value map object.

std::map< std::string, ValueTypePair >::const_iterator end () const

Returns a const iterator pointing to the end of the list.

• std::map< std::string, ValueTypePair >::iterator end ()

Returns an iterator pointing to the end of the list.

• std::map< std::string, ValueTypePair >::const_iterator begin () const

Returns a const iterator pointing to the beginning of the list.

std::map< std::string, ValueTypePair >::iterator begin ()

Returns an iterator pointing to the beginning of the list.

• void clear ()

Clears the map.

void addKey (std::string key)

Adds a key to the object with a default value.

void editValue4Key (std::string val, std::string key)

Edits a given value for a pre-existing key.

void editValue4Key (std::string val, int type, std::string key)

Edits a value for a pre-existing key and asserts type.

void addPair (std::string key, ValueTypePair val)

Adds a pair object to the map.

void addPair (std::string key, std::string val)

Adds a pair object to the map (with only strings)

void addPair (std::string key, std::string val, int type)

Adds a pair object and asserts a type.

void findType (std::string key)

Find what data type the value at the key is.

void assertType (std::string key, int type)

Assert the given type at the given key.

void findAllTypes ()

Find all types for all data in map.

void DisplayMap ()

Print out the map to console.

int size ()

Returns the size of the map.

std::string getString (std::string key)

Retrieve the string at the key.

bool getBool (std::string key)

Retrieve the boolean at the key.

• double getDouble (std::string key)

Retrieve the double at the key.

• int getInt (std::string key)

Retrieve the int at the key.

• std::string getValue (std::string key)

Retrieve the value at the key.

int getType (std::string key)

Retrieve the type at the key.

ValueTypePair & getPair (std::string key)

Retrieve the pair at the key.

Private Attributes

std::map< std::string, ValueTypePair > Key_Value
 Map of Keys and Values paired with types.

5.20.1 Detailed Description

Key-Value-Type Map object creating a map of the KeyValuePair objects.

C++ Object that creates a map of the KeyValuePair objects. Functions defined here allow the user to iterate through this map, access specific keys in the map, edit values associated with those keys, find the data types for the values in those keys, ect. The keys are used as an access operator for their corresponding value. As such, each key in the map is required to be unique, but the values are allowed to be duplicated.

```
5.20.2 Constructor & Destructor Documentation
```

```
5.20.2.1 KeyValueMap::KeyValueMap ( )
```

Default constructor.

```
5.20.2.2 KeyValueMap::\simKeyValueMap ( )
```

Default destructor.

5.20.2.3 KeyValueMap::KeyValueMap (const std::map < std::string, std::string > & map)

Construct from a map of strings.

 $5.20.2.4 \quad \text{KeyValueMap::KeyValueMap (std::string \textit{key,} \text{ std::string \textit{value})} \\$

Construct one element in the map.

```
5.20.2.5 KeyValueMap::KeyValueMap ( const KeyValueMap & map )
Copy constructor.
5.20.3 Member Function Documentation
5.20.3.1 KeyValueMap& KeyValueMap::operator= ( const KeyValueMap & map )
Equals overload.
5.20.3.2 ValueTypePair& KeyValueMap::operator[] ( const std::string key )
Return the ValueType reference at the given key.
5.20.3.3 ValueTypePair KeyValueMap::operator[]( const std::string key ) const
Return the ValueType at the give key.
5.20.3.4 std::map<std::string, ValueTypePair > & KeyValueMap::getMap ( )
Return a reference to the Key_Value map object.
5.20.3.5 std::map<std::string, ValueTypePair>::const_iterator KeyValueMap::end(_) const_
Returns a const iterator pointing to the end of the list.
5.20.3.6 std::map<std::string, ValueTypePair>::iterator KeyValueMap::end ( )
Returns an iterator pointing to the end of the list.
5.20.3.7 std::map<std::string, ValueTypePair>::const_iterator KeyValueMap::begin ( ) const
Returns a const iterator pointing to the beginning of the list.
5.20.3.8 std::map<std::string, ValueTypePair>::iterator KeyValueMap::begin ( )
Returns an iterator pointing to the beginning of the list.
5.20.3.9 void KeyValueMap::clear ( )
Clears the map.
5.20.3.10 void KeyValueMap::addKey ( std::string key )
Adds a key to the object with a default value.
5.20.3.11 void KeyValueMap::editValue4Key ( std::string val, std::string key )
Edits a given value for a pre-existing key.
```

```
5.20.3.12 void KeyValueMap::editValue4Key ( std::string val, int type, std::string key )
Edits a value for a pre-existing key and asserts type.
5.20.3.13 void KeyValueMap::addPair ( std::string key, ValueTypePair val )
Adds a pair object to the map.
5.20.3.14 void KeyValueMap::addPair ( std::string key, std::string val )
Adds a pair object to the map (with only strings)
5.20.3.15 void KeyValueMap::addPair ( std::string key, std::string val, int type )
Adds a pair object and asserts a type.
5.20.3.16 void KeyValueMap::findType ( std::string key )
Find what data type the value at the key is.
5.20.3.17 void KeyValueMap::assertType ( std::string key, int type )
Assert the given type at the given key.
5.20.3.18 void KeyValueMap::findAllTypes ( )
Find all types for all data in map.
5.20.3.19 void KeyValueMap::DisplayMap ( )
Print out the map to console.
5.20.3.20 int KeyValueMap::size ( )
Returns the size of the map.
5.20.3.21 std::string KeyValueMap::getString ( std::string key )
Retrieve the string at the key.
5.20.3.22 bool KeyValueMap::getBool ( std::string key )
Retrieve the boolean at the key.
5.20.3.23 double KeyValueMap::getDouble ( std::string key )
Retrieve the double at the key.
```

```
int KeyValueMap::getInt ( std::string key )
Retrieve the int at the key.
5.20.3.25 std::string KeyValueMap::getValue ( std::string key )
Retrieve the value at the key.
5.20.3.26 int KeyValueMap::getType ( std::string key )
Retrieve the type at the key.
5.20.3.27 ValueTypePair& KeyValueMap::getPair ( std::string key )
Retrieve the pair at the key.
5.20.4 Member Data Documentation
5.20.4.1 std::map<std::string, ValueTypePair > KeyValueMap::Key_Value [private]
Map of Keys and Values paired with types.
The documentation for this class was generated from the following file:
    · yaml_wrapper.h
       KMS_DATA Struct Reference
Data structure for the implemenation of the Krylov Multi-Space (KMS) Method.
#include <lark.h>
Public Attributes
    • int level = 0
          Current level in the recursion.
    • int max level = 0
          Maximum allowable recursion levels (Default = 0 -> GMRES, Max = 5)
    • int restart = -1
          Restart parameter for the outer iterates (Default = 20, Max = N)
    • int maxit = 0
          Maximum allowable iterations for the outer steps.
    • int inner_iter = 0
          Number of inner steps taken.
    • int outer_iter = 0
          Number of outer steps taken.
    • int total iter = 0
          Total number of iterations in all steps.
```

• double outer_reltol = 1e-6

Relative residual tolerance for outer steps (Default = 1e-6)

• double outer abstol = 1e-6

Absolute residual tolerance for outer steps (Default = 1e-6)

• double inner reltol = 0.1

Residual tolerance for inner steps made relative to outer steps (Default = 0.1)

bool Output_outer = true

True = Print the outer steps residuals.

• bool Output_inner = false

True = Print the inner steps residuals.

GMRESRP_DATA gmres_out

Data structure for the outer steps.

std::vector< GMRESRP DATA > gmres in

Data structures for each recursion level.

int(* matvec)(const Matrix< double > &x, Matrix< double > &Ax, const void *matvec_data)

User supplied matrix-vector product function.

• int(* terminal precon)(const Matrix< double > &r, Matrix< double > &p, const void *precon data)

Optional user supplied terminal preconditioner.

const void * matvec data

Data structure for the user's matvec function.

const void * term_precon

Data structure for the user's terminal preconditioner.

5.21.1 Detailed Description

Data structure for the implemenation of the Krylov Multi-Space (KMS) Method.

C-style object to be used in conjunction with the Krylov Multi-Space (KMS) Algorithm to iteratively solve non-symmetric, indefinite linear systems. This method was inspired by the Flexible GMRES (FGMRES) and Recursive GMRES (GMRESR) methods proposed by Saad (1993) and Vorst and Vuik (1991), respectively. The idea behind this method is to recursively call FGMRES to solve a linear system with pregressively smaller Krylov Subspaces built by a Right-Preconditioned GMRES algorithm. Thus creating a "V-cycle" of iteration similar to that seen in Multi-Grid algorithms.

5.21.2 Member Data Documentation

5.21.2.1 int KMS_DATA::level = 0

Current level in the recursion.

5.21.2.2 int KMS_DATA::max_level = 0

Maximum allowable recursion levels (Default = 0 -> GMRES, Max = 5)

5.21.2.3 int KMS_DATA::restart = -1

Restart parameter for the outer iterates (Default = 20, Max = N)

5.21.2.4 int KMS_DATA::maxit = 0

Maximum allowable iterations for the outer steps.

5.21.2.5 int KMS_DATA::inner_iter = 0

Number of inner steps taken.

5.21.2.6 int KMS_DATA::outer_iter = 0

Number of outer steps taken.

5.21.2.7 int KMS_DATA::total_iter = 0

Total number of iterations in all steps.

5.21.2.8 double KMS_DATA::outer_reltol = 1e-6

Relative residual tolerance for outer steps (Default = 1e-6)

5.21.2.9 double KMS_DATA::outer_abstol = 1e-6

Absolute residual tolerance for outer steps (Default = 1e-6)

5.21.2.10 double KMS_DATA::inner_reltol = 0.1

Residual tolerance for inner steps made relative to outer steps (Default = 0.1)

5.21.2.11 bool KMS_DATA::Output_outer = true

True = Print the outer steps residuals.

5.21.2.12 bool KMS_DATA::Output_inner = false

True = Print the inner steps residuals.

5.21.2.13 GMRESRP_DATA KMS_DATA::gmres_out

Data structure for the outer steps.

5.21.2.14 std::vector < GMRESRP_DATA > KMS_DATA::gmres_in

Data structures for each recursion level.

 $\textbf{5.21.2.15} \quad \text{int} (* \text{ KMS_DATA::matvec}) \text{ (const Matrix} < \text{double} > \&x, \text{Matrix} < \text{double} > \&Ax, \text{const void } * \text{matvec_data})$

User supplied matrix-vector product function.

5.21.2.16 int(* KMS_DATA::terminal_precon) (const Matrix < double > &r, Matrix < double > &p, const void *precon_data)

Optional user supplied terminal preconditioner.

5.21.2.17 const void* KMS_DATA::matvec_data

Data structure for the user's matvec function.

5.21.2.18 const void* KMS_DATA::term_precon

Data structure for the user's terminal preconditioner.

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

· lark.h

5.22 MAGPIE_DATA Struct Reference

MAGPIE Data Structure.

```
#include <magpie.h>
```

Public Attributes

- std::vector< GSTA_DATA > gsta_dat
- std::vector< mSPD_DATA > mspd_dat
- std::vector < GPAST_DATA > gpast_dat
- SYSTEM_DATA sys_dat

5.22.1 Detailed Description

MAGPIE Data Structure.

C-style object holding all information necessary to run a MAGPIE simulation. This is the data structure that will be used in other sub-routines when a mixed gas adsorption simulation needs to be run.

5.22.2 Member Data Documentation

5.22.2.1 std::vector < GSTA_DATA > MAGPIE_DATA::gsta_dat

5.22.2.2 std::vector<mSPD_DATA> MAGPIE_DATA::mspd_dat

5.22.2.3 std::vector < GPAST_DATA > MAGPIE_DATA::gpast_dat

5.22.2.4 SYSTEM_DATA MAGPIE_DATA::sys_dat

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

· magpie.h

5.23 MassBalance Class Reference

```
Mass Balance Object.
```

```
#include <shark.h>
```

Public Member Functions

• MassBalance ()

Default Constructor.

∼MassBalance ()

Default Destructor.

void Initialize_Object (MasterSpeciesList &List)

Function to initialize the MassBalance object from the MasterSpeciesList.

• void Display Info ()

Display the mass balance information.

void Set_Delta (int i, double v)

Function to set the ith weight (delta) for the mass balance.

void Set TotalConcentration (double v)

Set the total concentration of the mass balance to v (mol/L)

void Set_Type (int type)

Set the Mass Balance type to BATCH, CSTR, or PFR.

void Set Volume (double v)

Set the volume of the reactor.

void Set_FlowRate (double v)

Set the flow rate for the CSTR or PFR.

void Set_Area (double v)

Set the cross sectional area for the PFR.

• void Set_TimeStep (double v)

Set the time step for the CSTR or PFR.

void Set_InitialConcentration (double v)

Set the initial concentration for the mass balance.

void Set_InletConcentration (double v)

Set the inlet concentration for the CSTR or PFR.

void Set_SteadyState (bool ss)

Set the boolean for Steady-State simulation.

void Set_ZeroInitialSolids (bool solids)

Set the boolean for initial solids in solution.

void Set_Name (std::string name)

Set the name of the mass balance (i.e., Uranium, Carbonate, etc.)

• double Get Delta (int i)

Fetch the ith weight (i.e., delta) value.

double Sum_Delta ()

Sums up the delta values and returns the total (should never be zero)

double Get_TotalConcentration ()

Fetch the total concentration (mol/L)

• int Get_Type ()

Fetch the reactor type.

• double Get_Volume ()

Fetch the reactor volume.

• double Get_FlowRate ()

Fetch the reactor flow rate.

• double Get Area ()

Fetch the reactor cross section area.

double Get_TimeStep ()

Fetch the time step.

double Get_InitialConcentration ()

Fetch the initial concentration.

double Get_InletConcentration ()

Fetch the inlet concentration.

• bool isSteadyState ()

Fetch the steady-state condition.

bool isZeroInitialSolids ()

Fetch the initial solids condition.

std::string Get Name ()

Return name of mass balance object.

double Eval_Residual (const Matrix< double > &x_new, const Matrix< double > &x_old)

Evaluate the residual for the mass balance object given the log(C) concentrations.

double Eval_IC_Residual (const Matrix< double > &x)

Evaluate the initial residual for the unsteady mass balance object given the log(C) concentrations.

Protected Attributes

MasterSpeciesList * List

Pointer to a master species object.

• std::vector< double > Delta

Vector of weights (i.e., deltas) used in the mass balance.

• double TotalConcentration

Total concentration of specific object (mol/L)

int Type

Type of mass balance object (default = BATCH)

· double volume

Volume of the reactor (L)

· double flow_rate

Volumetric flow rate in reactor (L/hr)

· double xsec_area

Cross sectional area in PFR configuration (m^2 2)

· double dt

Time step for non-batch case (hrs)

· double InitialConcentration

Concentration initially in the domain (mol/L)

· double InletConcentration

Concentration in the inlet of the domain (mol/L)

bool SteadyState

True if running steady-state simulation.

bool ZeroInitialSolids

True if zero solids present for initial condition.

Private Attributes

· std::string Name

Name designation used in mass balance.

5.23.1 Detailed Description

Mass Balance Object.

C++ style object that holds data and functions associated with mass balances of primary species in a system. The mass balances involve a total concentration (in mol/L) and a vector of weighted contributions to that total concentration from each species in the MasterSpeciesList. This object only considers mass balances in a batch type of system (i.e., not input or output of mass). However, one could inherit from this object to create mass balances for flow systems as well.

```
5.23.2 Constructor & Destructor Documentation
```

```
5.23.2.1 MassBalance::MassBalance()
```

Default Constructor.

5.23.2.2 MassBalance::~MassBalance()

Default Destructor.

5.23.3 Member Function Documentation

5.23.3.1 void MassBalance::Initialize_Object (MasterSpeciesList & List)

Function to initialize the MassBalance object from the MasterSpeciesList.

```
5.23.3.2 void MassBalance::Display_Info()
```

Display the mass balance information.

```
5.23.3.3 void MassBalance::Set_Delta (int i, double v)
```

Function to set the ith weight (delta) for the mass balance.

This function sets the weight (i.e., delta value) of the ith species in the list to the value of v. That value represents the weighting of that species in the determination of the total mass for the primary species set.

Parameters

i	index of the species in the MasterSpeciesList
V	value of the weigth (or delta) applied to the mass balance

```
5.23.3.4 void MassBalance::Set_TotalConcentration ( double v )
Set the total concentration of the mass balance to v (mol/L)
5.23.3.5 void MassBalance::Set_Type ( int type )
Set the Mass Balance type to BATCH, CSTR, or PFR.
5.23.3.6 void MassBalance::Set_Volume ( double v )
Set the volume of the reactor.
5.23.3.7 void MassBalance::Set_FlowRate ( double v )
Set the flow rate for the CSTR or PFR.
5.23.3.8 void MassBalance::Set_Area ( double v )
Set the cross sectional area for the PFR.
5.23.3.9 void MassBalance::Set_TimeStep ( double v )
Set the time step for the CSTR or PFR.
5.23.3.10 void MassBalance::Set_InitialConcentration ( double v )
Set the initial concentration for the mass balance.
5.23.3.11 void MassBalance::Set_InletConcentration ( double v )
Set the inlet concentration for the CSTR or PFR.
5.23.3.12 void MassBalance::Set_SteadyState (bool ss)
Set the boolean for Steady-State simulation.
5.23.3.13 void MassBalance::Set_ZeroInitialSolids ( bool solids )
Set the boolean for initial solids in solution.
5.23.3.14 void MassBalance::Set_Name ( std::string name )
Set the name of the mass balance (i.e., Uranium, Carbonate, etc.)
5.23.3.15 double MassBalance::Get_Delta (int i)
Fetch the ith weight (i.e., delta) value.
```

```
5.23.3.16 double MassBalance::Sum_Delta ( )
Sums up the delta values and returns the total (should never be zero)
5.23.3.17 double MassBalance::Get_TotalConcentration ( )
Fetch the total concentration (mol/L)
5.23.3.18 int MassBalance::Get_Type ( )
Fetch the reactor type.
5.23.3.19 double MassBalance::Get_Volume ( )
Fetch the reactor volume.
5.23.3.20 double MassBalance::Get_FlowRate ( )
Fetch the reactor flow rate.
5.23.3.21 double MassBalance::Get_Area ( )
Fetch the reactor cross section area.
5.23.3.22 double MassBalance::Get_TimeStep ( )
Fetch the time step.
5.23.3.23 double MassBalance::Get_InitialConcentration ( )
Fetch the initial concentration.
5.23.3.24 double MassBalance::Get_InletConcentration ( )
Fetch the inlet concentration.
5.23.3.25 bool MassBalance::isSteadyState ( )
Fetch the steady-state condition.
5.23.3.26 bool MassBalance::isZeroInitialSolids ( )
Fetch the initial solids condition.
5.23.3.27 std::string MassBalance::Get_Name ( )
Return name of mass balance object.
5.23.3.28 double MassBalance::Eval_Residual ( const Matrix < double > & x_new, const Matrix < double > & x_old )
```

This function calculates and provides the residual for this mass balance object based on the total concentration in the system and the weighted contributions from each species. Concentrations are given as the log(C) values.

Evaluate the residual for the mass balance object given the log(C) concentrations.

Parameters

x_new	matrix of the log(C) concentration values at the current non-linear step
x_old	matrix of the old log(C) concentration values for transient simulations

5.23.3.29 double MassBalance::Eval_IC_Residual (const Matrix < double > & x)

Evaluate the initial residual for the unsteady mass balance object given the log(C) concentrations.

This function calculates and provides the initial residual for this mass balance object based on the initial concentration in the system and the weighted contributions from each species. Concentrations are given as the log(C) values.

Parameters

x matrix of the log(C) concentration values at the current non-linear step

5.23.4 Member Data Documentation

5.23.4.1 MasterSpeciesList* MassBalance::List [protected]

Pointer to a master species object.

5.23.4.2 std::vector<double> MassBalance::Delta [protected]

Vector of weights (i.e., deltas) used in the mass balance.

5.23.4.3 double MassBalance::TotalConcentration [protected]

Total concentration of specific object (mol/L)

5.23.4.4 int MassBalance::Type [protected]

Type of mass balance object (default = BATCH)

5.23.4.5 double MassBalance::volume [protected]

Volume of the reactor (L)

5.23.4.6 double MassBalance::flow_rate [protected]

Volumetric flow rate in reactor (L/hr)

5.23.4.7 double MassBalance::xsec_area [protected]

Cross sectional area in PFR configuration (m^2)

```
5.23.4.8 double MassBalance::dt [protected]
Time step for non-batch case (hrs)
5.23.4.9 double MassBalance::InitialConcentration [protected]
Concentration initially in the domain (mol/L)
5.23.4.10 double MassBalance::InletConcentration [protected]
Concentration in the inlet of the domain (mol/L)
5.23.4.11 bool MassBalance::SteadyState [protected]
True if running steady-state simulation.
5.23.4.12 bool MassBalance::ZeroInitialSolids [protected]
True if zero solids present for initial condition.
5.23.4.13 std::string MassBalance::Name [private]
Name designation used in mass balance.
The documentation for this class was generated from the following file:
    · shark.h
5.24
      MasterSpeciesList Class Reference
Master Species List Object.
#include <shark.h>
Public Member Functions

    MasterSpeciesList ()

         Default constructor.

    ∼MasterSpeciesList ()

         Default destructor.

    MasterSpeciesList (const MasterSpeciesList &msl)

         Copy Constructor.

    MasterSpeciesList & operator= (const MasterSpeciesList &msl)

          Equals operator.

    void set_list_size (int i)
```

Function to register the ith species in the list based on a registered molecular formula (see mola.h)

Function to initialize the size of the list.

• void set_species (int i, std::string formula)

 void set_species (int i, int charge, double enthalpy, double entropy, double energy, bool HS, bool G, std::string Phase, std::string Name, std::string Formula, std::string lin_formula)

Function to register the ith species in the list based on custom molecule information (see mola.h)

void DisplayInfo (int i)

Function to display information of ith object.

void DisplayAll ()

Function to display all information of list.

void DisplayConcentrations (Matrix< double > &C)

Function to display the concentrations of species in list.

void set_alkalinity (double alk)

Set the alkalinity of the solution (Default = 0 M)

• int list_size ()

Returns size of list.

Molecule & get_species (int i)

Returns a reference to the ith species in master list.

int get_index (std::string name)

Returns an integer representing location of the named species in the list.

• double charge (int i)

Fetch and return charge of ith species in list.

· double alkalinity ()

Fetch the value of alkalinity of the solution (mol/L)

• std::string speciesName (int i)

Function to return the name of the ith species.

double Eval_ChargeResidual (const Matrix< double > &x)

Calculate charge balance residual for the electroneutrality constraint.

Protected Attributes

· int size

Size of the list.

• std::vector< Molecule > species

List of Molecule Objects.

· double residual_alkalinity

Conc of strong base - conc of strong acid in solution (mol/L)

5.24.1 Detailed Description

Master Species List Object.

C++ style object that holds data and function associated with solving multi-species problems. This object contains a vector of Molecule objects from mola.h and uses those objects to help setup speciation problems that need to be solved. One of the primary functions in this object is the contribution of electroneutrality (Eval_ChargeResidual). However, we only need this constraint if the pH of our aqueous system is unknown.

5.24.2 Constructor & Destructor Documentation

5.24.2.1 MasterSpeciesList::MasterSpeciesList ()

Default constructor.

```
5.24.2.2 MasterSpeciesList:: ~ MasterSpeciesList ( )
Default destructor.
5.24.2.3 MasterSpeciesList::MasterSpeciesList ( const MasterSpeciesList & msl )
Copy Constructor.
5.24.3 Member Function Documentation
5.24.3.1 MasterSpeciesList& MasterSpeciesList::operator= ( const MasterSpeciesList & msl )
Equals operator.
5.24.3.2 void MasterSpeciesList::set_list_size ( int i )
Function to initialize the size of the list.
5.24.3.3 void MasterSpeciesList::set_species ( int i, std::string formula )
Function to register the ith species in the list based on a registered molecular formula (see mola.h)
5.24.3.4 void MasterSpeciesList::set species (int i, int charge, double enthalpy, double entropy, double energy, bool HS,
         bool G, std::string Phase, std::string Name, std::string Formula, std::string lin_formula)
Function to register the ith species in the list based on custom molecule information (see mola.h)
5.24.3.5 void MasterSpeciesList::DisplayInfo (int i)
Function to display information of ith object.
5.24.3.6 void MasterSpeciesList::DisplayAll ( )
Function to display all information of list.
5.24.3.7 void MasterSpeciesList::DisplayConcentrations ( Matrix < double > \& C )
Function to display the concentrations of species in list.
This function will print to the console the species list in order with each species associated concentration from the
matrix C. The concentrations and species list MUST be in the same order and the units of C are assumed to be
mol/L.
Parameters
```

matrix of concentrations of species in the list in mol/L

5.24.3.8 void MasterSpeciesList::set_alkalinity (double alk)

Set the alkalinity of the solution (Default = 0 M)

This function is used to set the value of residual alkalinity used in the electroneutrality calculations. Typically, this value will be 0 M (mol/L) if all species in the system are present as variables. However, occasionally, one may want to set the alkalinity of the solution to a constant in order to restrict the pH of the solution.

Parameters

alk Residual alkalinity in M (mol/L)

5.24.3.9 int MasterSpeciesList::list_size ()

Returns size of list.

5.24.3.10 Molecule& MasterSpeciesList::get_species (int i)

Returns a reference to the ith species in master list.

This function will return a Molecule object for the ith species in the list of molecules. Once returned, the user then can operate on that molecule using the functions define in mola.h.

5.24.3.11 int MasterSpeciesList::get_index (std::string name)

Returns an integer representing location of the named species in the list.

5.24.3.12 double MasterSpeciesList::charge (int i)

Fetch and return charge of ith species in list.

5.24.3.13 double MasterSpeciesList::alkalinity ()

Fetch the value of alkalinity of the solution (mol/L)

5.24.3.14 std::string MasterSpeciesList::speciesName (int i)

Function to return the name of the ith species.

5.24.3.15 double MasterSpeciesList::Eval_ChargeResidual (const Matrix< double > & x)

Calculate charge balance residual for the electroneutrality constraint.

This function returns the value of the residual for the electroneutrality equation in the system. Electroneutrality is based on the concentrations and charges of each species in the system so the charges of each molecule must be appropriately set. Concentrations of those species are fed into this function via the argument x, but come in as the log(C) values (i.e., x = log(C)).

Parameters

x matrix of the log(C) concentration values at the current non-linear step

```
5.24.4 Member Data Documentation
5.24.4.1 int MasterSpeciesList::size [protected]
Size of the list.
5.24.4.2 std::vector<Molecule> MasterSpeciesList::species [protected]
List of Molecule Objects.
5.24.4.3 double MasterSpeciesList::residual_alkalinity [protected]
Conc of strong base - conc of strong acid in solution (mol/L)
The documentation for this class was generated from the following file:
    · shark.h
5.25 Matrix < T > Class Template Reference
Templated C++ Matrix Class Object (click Matrix to go to function definitions)
#include <macaw.h>
Public Member Functions
    • Matrix (int rows, int columns)
           Constructor for matrix with given number of rows and columns.

    T & operator() (int i, int j)

          Access operator for the matrix element at row i and column j (e.g., aij = A(i,j))
    • T operator() (int i, int j) const
           Constant access operator for the the matrix element at row i and column j.

    Matrix (const Matrix &M)

          Copy constructor for constructing a matrix as a copy of another matrix.

    Matrix & operator= (const Matrix &M)

          Equals operator for setting one matrix equal to another matrix.

    Matrix ()

          Default constructor for creating an empty matrix.
    • ~Matrix ()
          Default destructor for clearing out memory.
    void set_size (int i, int j)
          Function to set/change the size of a matrix to i rows and j columns.
    · void zeros ()
          Function to set/change all values in a matrix to zeros.

    void edit (int i, int j, T value)

          Function to set/change the element of a matrix at row i and column j to given value.
    • int rows ()
          Function to return the number of rows in a given matrix.
```

int columns ()

Function to return the number of columns in a matrix.

T determinate ()

Function to compute the determinate of a matrix and return that value.

• T norm ()

Function to compute the L2-norm of a matrix and return that value.

• T sum ()

Function to compute the sum of all elements in a matrix and return that value.

T inner product (const Matrix &x)

Function to compute the inner product between this matrix and matrix x.

Matrix & cofactor (const Matrix &M)

Function to convert this matrix to a cofactor matrix of the given matrix M.

Matrix operator+ (const Matrix &M)

Operator to add this matrix and matrix M and return the new matrix result.

Matrix operator- (const Matrix &M)

Operator to subtract this matrix and matrix M and return the new matrix result.

Matrix operator* (const T)

Operator to multiply this matrix by a scalar T return the new matrix result.

Matrix operator/ (const T)

Operator to divide this matrix by a scalar T and return the new matrix result.

Matrix operator* (const Matrix &M)

Operator to multiply this matrix and matrix M and return the new matrix result.

Matrix outer product (const Matrix &M)

Operator to perform an outer product between this and M and return result.

- Matrix & transpose (const Matrix &M)
- Matrix & transpose_multiply (const Matrix &MT, const Matrix &v)

Function to convert this matrix into the result of the given matrix M transposed and multiplied by the other given matrix v

Matrix & adjoint (const Matrix &M)

Function to convert this matrix to the adjoint of the given matrix.

Matrix & inverse (const Matrix &M)

Function to convert this matrix to the inverse of the given matrix.

void Display (const std::string Name)

Function to display the contents of this matrix given a Name for the matrix.

Matrix & tridiagonalSolve (const Matrix &A, const Matrix &b)

Function to solve Ax=b for x if A is symmetric, tridiagonal (this->x)

Matrix & ladshawSolve (const Matrix &A, const Matrix &d)

Function to solve Ax=d for x if A is non-symmetric, tridiagonal (this->x)

Matrix & tridiagonalFill (const T A, const T B, const T C, bool Spherical)

Function to fill in this matrix with coefficients A, B, and C to form a tridiagonal matrix.

Matrix & naturalLaplacian3D (int m)

Function to fill out this matrix with coefficients from a 3D Laplacian function.

• Matrix & sphericalBCFill (int node, const T coeff, T variable)

Function to fill out a column matrix with spherical specific boundary conditions.

Matrix & ConstantICFill (const T IC)

Function to set all values in a column matrix to a given constant.

Matrix & SolnTransform (const Matrix &A, bool Forward)

Function to transform the values in a column matrix from cartesian to spherical coordinates.

T sphericalAvg (double radius, double dr, double bound, bool Dirichlet)

Function to compute a spatial average of this column matrix in spherical coordinates.

• T IntegralAvg (double radius, double dr, double bound, bool Dirichlet)

Function to compute a spatial average of this column matrix in spherical coordinates.

T IntegralTotal (double dr, double bound, bool Dirichlet)

Function to compute a spatial total of this column matrix in spherical coordinates.

Matrix & tridiagonalVectorFill (const std::vector< T > &A, const std::vector< T > &B, const std::vector< T > &C)

Function to fill in this matrix, in tridiagonal fashion, using the vectors of coefficients.

Matrix & columnVectorFill (const std::vector < T > &A)

Function to fill in a column matrix with the values of the given vector object.

Matrix & columnProjection (const Matrix &b, const Matrix &b_old, const double dt, const double dt_old)

Function to project a column matrix solution in time based on older state vectors.

• Matrix & dirichletBCFill (int node, const T coeff, T variable)

Function to fill in a column matrix with all zeros except at the given node.

Matrix & diagonalSolve (const Matrix &D, const Matrix &v)

Function to solve the system Dx=v for x given that D is diagonal (this->x)

Matrix & upperTriangularSolve (const Matrix &U, const Matrix &v)

Function to solve the system Ux=v for x given that U is upper Triagular (this->x)

• Matrix & lowerTriangularSolve (const Matrix &L, const Matrix &v)

Function to solve the system Lx=v for x given that L is lower Triagular (this->x)

Matrix & upperHessenberg2Triangular (Matrix &b)

Function to convert this square matrix to upper Triangular (assuming this is upper Hessenberg)

• Matrix & lowerHessenberg2Triangular (Matrix &b)

Function to convert this square matrix to lower Triangular (assuming this is lower Hessenberg)

• Matrix & upperHessenbergSolve (const Matrix &H, const Matrix &v)

Function to solve the system Hx=v for x given that H is upper Hessenberg (this->x)

• Matrix & lowerHessenbergSolve (const Matrix &H, const Matrix &v)

Function to solve the system Hx=v for x given that H is lower Hessenberg (this->x)

Matrix & qrSolve (const Matrix &M, const Matrix &b)

Function to solve the system Mx=b using QR factorization for x given that M is invertable.

Matrix & columnExtract (int j, const Matrix &M)

Function to set this column matrix to the jth column of the given matrix M.

Matrix & rowExtract (int i, const Matrix &M)

Function to set this row matrix to the ith row of the given matrix M.

Matrix & columnReplace (int j, const Matrix &v)

Function to this matrices' jth column with the given column matrix v.

Matrix & rowReplace (int i, const Matrix &v)

Function to this matrices' ith row with the given row matrix v.

• void rowShrink ()

Function to delete the last row of this matrix.

void columnShrink ()

Function to delete the last column of this matrix.

void rowExtend (const Matrix &v)

Function to add the row matrix v to the end of this matrix.

void columnExtend (const Matrix &v)

Function to add the column matrix v to the end of this matrix.

Protected Attributes

• int num rows

Number of rows of the matrix.

· int num_cols

Number of columns of the matrix.

std::vector< T > Data

Storage vector for the elements of the matrix.

5.25.1 Detailed Description

```
template < class T > class Matrix < T >
```

Templated C++ Matrix Class Object (click Matrix to go to function definitions)

C++ templated class object containing many different functions, actions, and solver routines associated with Dense Matrices. Operator overloads are also provided to give the user a more natural way of operating matrices on other matrices or scalars. These operator overloads are especially useful for reducing the amount of code needed to be written when working with matrix-based problems.

```
5.25.2 Constructor & Destructor Documentation
```

```
5.25.2.1 template < class T > Matrix < T >::Matrix ( int rows, int columns )
```

Constructor for matrix with given number of rows and columns.

```
5.25.2.2 template < class T > Matrix < T >::Matrix ( const Matrix < T > & \it M )
```

Copy constructor for constructing a matrix as a copy of another matrix.

```
5.25.2.3 template < class T > Matrix < T >::Matrix ( )
```

Default constructor for creating an empty matrix.

```
5.25.2.4 template < class T > Matrix < T >::\sim Matrix ( )
```

Default destructor for clearing out memory.

5.25.3 Member Function Documentation

```
5.25.3.1 template < class T > T & Matrix < T >::operator() ( int i, int j )
```

Access operator for the matrix element at row i and column j (e.g., aij = A(i,j))

```
5.25.3.2 template < class T > T Matrix < T >::operator() ( int i, int j ) const
```

Constant access operator for the the matrix element at row i and column j.

```
5.25.3.3 template < class T > Matrix < T > & Matrix < T > ::operator= ( const Matrix < T > & M )
```

Equals operator for setting one matrix equal to another matrix.

```
5.25.3.4 template < class T > void Matrix < T >::set_size ( int i, int j )
```

Function to set/change the size of a matrix to i rows and j columns.

```
5.25.3.5 template < class T > void Matrix < T >::zeros ( )
```

Function to set/change all values in a matrix to zeros.

```
5.25.3.6 template < class T > void Matrix < T > ::edit ( int i, int j, T value )
```

Function to set/change the element of a matrix at row i and column j to given value.

```
5.25.3.7 template < class T > int Matrix < T >::rows ( )
```

Function to return the number of rows in a given matrix.

```
5.25.3.8 template < class T > int Matrix < T >::columns ( )
```

Function to return the number of columns in a matrix.

```
5.25.3.9 template < class T > T Matrix < T >::determinate ( )
```

Function to compute the determinate of a matrix and return that value.

```
5.25.3.10 template < class T > T Matrix < T >::norm ( )
```

Function to compute the L2-norm of a matrix and return that value.

```
5.25.3.11 template < class T > T Matrix < T >::sum ( )
```

Function to compute the sum of all elements in a matrix and return that value.

```
5.25.3.12 template < class T > T Matrix < T >::inner_product ( const Matrix < T > & x )
```

Function to compute the inner product between this matrix and matrix \boldsymbol{x} .

```
5.25.3.13 template < class T > Matrix < T > & Matrix < T > ::cofactor ( const Matrix < T > & M )
```

Function to convert this matrix to a cofactor matrix of the given matrix M.

```
5.25.3.14 template < class T > Matrix < T > Matrix < T > :: operator + ( const Matrix < T > & M )
```

Operator to add this matrix and matrix M and return the new matrix result.

```
5.25.3.15 template < class T > Matrix < T > Matrix < T > :: operator- ( const Matrix < T > & M )
```

Operator to subtract this matrix and matrix M and return the new matrix result.

```
5.25.3.16 template < class T > Matrix < T > Matrix < T > ::operator* ( const T a )
```

Operator to multiply this matrix by a scalar T return the new matrix result.

```
5.25.3.17 template < class T> Matrix < T> Matrix < T>::operator/ (const Ta)
```

Operator to divide this matrix by a scalar T and return the new matrix result.

```
5.25.3.18 template < class T> Matrix < T> Matrix < T>::operator * ( const Matrix < T> & M )
```

Operator to multiply this matrix and matrix M and return the new matrix result.

```
5.25.3.19 template < class T > Matrix < T > Matrix < T > ::outer_product ( const Matrix < T > & M )
```

Operator to perform an outer product between this and M and return result.

```
5.25.3.20 template < class T > Matrix < T > & Matrix < T >::transpose (const Matrix < T > & M
```

Function to convert this matrix to the transpose of the given matrix M

```
5.25.3.21 template < class T > Matrix < T > & Matrix < T > ::transpose_multiply ( const Matrix < T > & MT, const Matrix < T > & \nu)
```

Function to convert this matrix into the result of the given matrix M transposed and multiplied by the other given matrix v.

```
5.25.3.22 template < class T > Matrix < T > & Matrix < T > ::adjoint (const Matrix < T > & M)
```

Function to convert this matrix to the adjoint of the given matrix.

```
5.25.3.23 template < class T > Matrix < T > & Matrix < T > ::inverse (const Matrix < T > & M)
```

Function to convert this matrix to the inverse of the given matrix.

```
5.25.3.24 template < class T > void Matrix < T >::Display (const std::string Name)
```

Function to display the contents of this matrix given a Name for the matrix.

```
5.25.3.25 template < class T > Matrix < T > & Matrix < T >::tridiagonal Solve (const Matrix < T > & A, const Matrix < T > & b)
```

Function to solve Ax=b for x if A is symmetric, tridiagonal (this->x)

```
5.25.3.26 template < class T > Matrix < T > & Matrix < T > ::ladshawSolve ( const Matrix < T > & A, const Matrix < T > & d )
```

Function to solve Ax=d for x if A is non-symmetric, tridiagonal (this->x)

```
5.25.3.27 template < class T > Matrix < T > & Matrix < T > ::tridiagonalFill ( const T A, const T B, const T C, bool Spherical )
```

Function to fill in this matrix with coefficients A, B, and C to form a tridiagonal matrix.

This function fills in the diagonal elements of a square matrix with coefficient B, upper diagonal with C, and lower diagonal with A. The boolean will apply a transformation to those coefficients, if the problem happens to stem from 1-D diffusion in spherical coordinates.

```
5.25.3.28 template < class T > Matrix < T > & Matrix < T >::naturalLaplacian3D ( int m )
```

Function to fill out this matrix with coefficients from a 3D Laplacian function.

This function will fill out the coefficients of the matrix with the coefficients that stem from discretizing a 3D Laplacian on a natural grid with 2nd order finite differences.

```
5.25.3.29 template < class T > Matrix < T > & Matrix < T >::spherical BCFill (int node, const T coeff, T variable)
```

Function to fill out a column matrix with spherical specific boundary conditions.

This function will fille out a column matrix with zeros at all nodes expect for the node indicated. That node's value will be the product of the node id with the coeff and variable values given.

```
5.25.3.30 template < class T > Matrix < T > & Matrix < T >::ConstantICFill (const T /C)
```

Function to set all values in a column matrix to a given constant.

```
5.25.3.31 template < class T > Matrix < T > & Matrix < T > ::SolnTransform ( const Matrix < T > & A, bool Forward )
```

Function to transform the values in a column matrix from cartesian to spherical coordinates.

```
5.25.3.32 template < class T > T Matrix < T >::sphericalAvg ( double radius, double dr, double bound, bool Dirichlet )
```

Function to compute a spatial average of this column matrix in spherical coordinates.

This function is used to compute an average value of a variable, represented in this column matrix, by integrating over the domain of the sphere. (Assumes you have variable value at center node)

Parameters

radius	radius of the sphere
dr	space between each node
bound	value of the variable at the boundary
Dirichlet	True if problem has a Dirichlet BC, False if Neumann

 $5.25.3.33 \quad template < class \ T > T \ Matrix < T > ::Integral Avg \ (\ double \ \textit{radius}, \ double \ \textit{dr,} \ double \ \textit{bound}, \ bool \ \textit{Dirichlet} \)$

Function to compute a spatial average of this column matrix in spherical coordinates.

This function is used to compute an average value of a variable, represented in this column matrix, by integrating over the domain of the sphere. (Assumes you DO NOT have variable value at center node)

Parameters

radius	radius of the sphere
dr	space between each node
bound	value of the variable at the boundary
Dirichlet	True if problem has a Dirichlet BC, False if Neumann

5.25.3.34 template < class T > T Matrix < T >::IntegralTotal (double dr, double bound, bool Dirichlet)

Function to compute a spatial total of this column matrix in spherical coordinates.

This function is used to compute an average value of a variable, represented in this column matrix, by integrating over the domain of the sphere. (Assumes you DO NOT have variable value at center node)

Parameters

dr	space between each node
bound	value of the variable at the boundary
Dirichlet	True if problem has a Dirichlet BC, False if Neumann

5.25.3.35 template < class T > Matrix < T > & Matrix < T > ::tridiagonal Vector Fill (const std::vector < T > & A, const std::vector < T > & B, const std::vector < T > & C)

Function to fill in this matrix, in tridiagonal fashion, using the vectors of coefficients.

5.25.3.36 template < class T > Matrix < T > & Matrix < T >::columnVectorFill (const std::vector < T > & A)

Function to fill in a column matrix with the values of the given vector object.

5.25.3.37 template < class T > Matrix < T > & Matrix < T > :::columnProjection (const Matrix < T > & b, const Matrix < T > & const Matrix < T > &

Function to project a column matrix solution in time based on older state vectors.

This function is used in finch.h to form Matrix u_star. It uses the size of the current step and old step, dt and dt_old respectively, to form an approximation for the next state. The current state and olde state of the variables are passed as b and b_old respectively.

5.25.3.38 template < class T > Matrix < T > & Matrix < T > ::dirichletBCFill (int node, const T coeff, T variable)

Function to fill in a column matrix with all zeros except at the given node.

Similar to sphericalBCFill, this function will set the values of all elements in the column matrix to zero except at the given node, where the value is set to the product of coeff and variable. This is often used to set BCs in finch.h or other related files/simulations.

5.25.3.39 template < class T > Matrix < T > & Matrix < T > ::diagonal Solve (const Matrix < T > & D, const Matrix < T > & ν)

Function to solve the system Dx=v for x given that D is diagonal (this->x)

5.25.3.40 template < class T > Matrix < T > & Matrix < T > ::upperTriangularSolve (const Matrix < T > & U, const Matrix < T > & v)

Function to solve the system Ux=v for x given that U is upper Triagular (this->x)

5.25.3.41 template < class T > Matrix < T > & Matrix < T > ::lowerTriangularSolve (const Matrix < T > & L, const Matrix < T > & ν)

Function to solve the system Lx=v for x given that L is lower Triagular (this->x)

```
5.25.3.42 template < class T > Matrix < T > & Matrix < T > ::upperHessenberg2Triangular (Matrix < T > & b)
```

Function to convert this square matrix to upper Triangular (assuming this is upper Hessenberg)

During this transformation, a column vector (b) is also being transformed to represent the BCs in a linear system. This algorithm uses Givens Rotations to efficiently convert the upper Hessenberg matrix to an upper triangular matrix.

```
5.25.3.43 template < class T > Matrix < T > & Matrix < T >:: lower Hessenberg 2 Triangular (Matrix < T > & b)
```

Function to convert this square matrix to lower Triangular (assuming this is lower Hessenberg)

During this transformation, a column vector (b) is also being transformed to represent the BCs in a linear system. This algorithm uses Givens Rotations to efficiently convert the lower Hessenberg matrix to an lower triangular matrix.

```
5.25.3.44 template < class T > Matrix < T > & Matrix < T > ::upperHessenbergSolve ( const Matrix < T > & H, const Matrix < T > & \nu)
```

Function to solve the system Hx=v for x given that H is upper Hessenberg (this->x)

```
5.25.3.45 template < class T > Matrix < T > & Matrix < T > ::lowerHessenbergSolve ( const Matrix < T > & \it H, const Matrix < T > & \it v )
```

Function to solve the system Hx=v for x given that H is lower Hessenberg (this->x)

```
5.25.3.46 template < class T > Matrix < T > & Matrix < T > ::qrSolve ( const Matrix < T > & M, const Matrix < T > & b )
```

Function to solve the system Mx=b using QR factorization for x given that M is invertable.

```
5.25.3.47 template < class T > Matrix < T > & Matrix < T > :::columnExtract ( int j, const Matrix < T > & M)
```

Function to set this column matrix to the jth column of the given matrix M.

```
5.25.3.48 template < class T > Matrix < T > & Matrix < T > ::rowExtract ( int i, const Matrix < T > & M )
```

Function to set this row matrix to the ith row of the given matrix M.

```
5.25.3.49 template < class T > Matrix < T > & Matrix < T > ::columnReplace (int j, const Matrix < T > & \nu)
```

Function to this matrices' jth column with the given column matrix v.

```
5.25.3.50 template < class T > Matrix < T > & Matrix < T >::rowReplace (int i, const Matrix < T > & \nu)
```

Function to this matrices' ith row with the given row matrix v.

```
5.25.3.51 template < class T > void Matrix < T >::rowShrink ( )
```

Function to delete the last row of this matrix.

```
5.25.3.52 template < class T > void Matrix < T >::columnShrink ( )
```

Function to delete the last column of this matrix.

```
5.25.3.53 template < class T > void Matrix < T >::rowExtend ( const Matrix < T > & \nu )
```

Function to add the row matrix v to the end of this matrix.

```
5.25.3.54 template < class T > void Matrix < T >::columnExtend ( const Matrix < T > & \nu )
```

Function to add the column matrix v to the end of this matrix.

5.25.4 Member Data Documentation

```
5.25.4.1 template < class T > int Matrix < T >::num_rows [protected]
```

Number of rows of the matrix.

```
5.25.4.2 template < class T > int Matrix < T >::num_cols [protected]
```

Number of columns of the matrix.

```
5.25.4.3 template < class T > std::vector < T > Matrix < T >::Data [protected]
```

Storage vector for the elements of the matrix.

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

· macaw.h

5.26 MIXED_GAS Struct Reference

Data structure holding information necessary for computing mixed gas properties.

```
#include <egret.h>
```

Public Attributes

• int N

Given: Total number of gas species.

bool CheckMolefractions = true

Given: True = Check Molefractions for errors.

double total_pressure

Given: Total gas pressure (kPa)

· double gas_temperature

Given: Gas temperature (K)

· double velocity

Given: Gas phase velocity (cm/s)

· double char_length

Given: Characteristic Length (cm)

std::vector< double > molefraction

Given: Gas molefractions of each species (-)

double total_density

Calculated: Total gas density (g/cm[^]3) {use RE3}.

• double total_dyn_vis

Calculated: Total dynamic viscosity (g/cm/s)

· double kinematic_viscosity

Calculated: Kinematic viscosity (cm²/s)

• double total_molecular_weight

Calculated: Total molecular weight (g/mol)

· double total_specific_heat

Calculated: Total specific heat (J/g/K)

· double Reynolds

Calculated: Value of the Reynold's number (-)

• Matrix< double > binary_diffusion

Calculated: Tensor matrix of binary gas diffusivities (cm 2 /s)

std::vector< PURE_GAS > species_dat

Vector of the pure gas info of all species.

5.26.1 Detailed Description

Data structure holding information necessary for computing mixed gas properties.

C-style object holding the mixed gas information necessary for performing gas dynamic simulations. This object works in conjunction with the calculate_variables function and uses the kinetic theory of gases to estimate mixed gas properties.

5.26.2 Member Data Documentation

5.26.2.1 int MIXED_GAS::N

Given: Total number of gas species.

5.26.2.2 bool MIXED_GAS::CheckMolefractions = true

Given: True = Check Molefractions for errors.

5.26.2.3 double MIXED_GAS::total_pressure

Given: Total gas pressure (kPa)

5.26.2.4 double MIXED_GAS::gas_temperature

Given: Gas temperature (K)

5.26.2.5 double MIXED_GAS::velocity

Given: Gas phase velocity (cm/s)

5.26.2.6 double MIXED_GAS::char_length

Given: Characteristic Length (cm)

5.26.2.7 std::vector<double> MIXED_GAS::molefraction

Given: Gas molefractions of each species (-)

5.26.2.8 double MIXED_GAS::total_density

Calculated: Total gas density (g/cm³) {use RE3}.

5.26.2.9 double MIXED_GAS::total_dyn_vis

Calculated: Total dynamic viscosity (g/cm/s)

5.26.2.10 double MIXED_GAS::kinematic_viscosity

Calculated: Kinematic viscosity (cm^2/s)

5.26.2.11 double MIXED_GAS::total_molecular_weight

Calculated: Total molecular weight (g/mol)

5.26.2.12 double MIXED_GAS::total_specific_heat

Calculated: Total specific heat (J/g/K)

5.26.2.13 double MIXED_GAS::Reynolds

Calculated: Value of the Reynold's number (-)

 ${\bf 5.26.2.14} \quad \textbf{Matrix}{<} \textbf{double}{>} \, \textbf{MIXED_GAS::binary_diffusion}$

Calculated: Tensor matrix of binary gas diffusivities (cm²/s)

```
5.26.2.15 std::vector<PURE_GAS> MIXED_GAS::species_dat
```

Vector of the pure gas info of all species.

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

· egret.h

5.27 Molecule Class Reference

C++ Molecule Object built from Atom Objects (click Molecule to go to function definitions)

```
#include <mola.h>
```

Public Member Functions

• Molecule ()

Default Constructor (builds an empty molecule object)

∼Molecule ()

Default Destructor (clears out memory)

• Molecule (int charge, double enthalpy, double entropy, double energy, bool HS, bool G, std::string Phase, std::string Name, std::string Formula, std::string lin_formula)

Construct any molecule from the available information.

 void Register (int charge, double enthalpy, double entropy, double energy, bool HS, bool G, std::string Phase, std::string Name, std::string Formula, std::string lin formula)

Function to register this molecule from the available information.

void Register (std::string formula)

Function to register this molecule based on the given formula (if formula is in library)

void setFormula (std::string form)

Sets the formula for a molecule.

• void calculateMolarWeight ()

Forces molecule to calculate its molar weight.

void calculateMolarVolume ()

Force molecule to calculate van der Waals volume.

void calculateMolarArea ()

Force molecule to calculate van der Waals area.

void setMolarWeigth (double mw)

Set the molar weight of species to a constant.

• void setMolarVolume (double v)

Set the van der Waals volume of the species to a constant.

void setMolarArea (double a)

Set the van der Waals area of the species to a constant.

void editCharge (int c)

Change the ionic charge of a molecule.

void editOneOxidationState (int state, std::string Symbol)

Change oxidation state of one of the given atoms (always first match found)

void editAllOxidationStates (int state, std::string Symbol)

Change oxidation state of all of the given atoms.

void calculateAvgOxiState (std::string Symbol)

Function to calculate the average oxidation state of the atoms.

void editEnthalpy (double enthalpy)

Edit the molecules formation enthalpy (J/mol)

void editEntropy (double entropy)

Edit the molecules formation entropy (J/K/mol)

void editHS (double H, double S)

Edit both formation enthalpy and entropy.

void editEnergy (double energy)

Edit Gibb's formation energy.

void removeOneAtom (std::string Symbol)

Removes one atom of the symbol given (always the first atom found)

• void removeAllAtoms (std::string Symbol)

Removes all atoms of the symbol given.

• int Charge ()

Return the charge of the molecule.

double MolarWeight ()

Return the molar weight of the molecule.

• double MolarVolume ()

Return the van der Waals volume of the molecule.

• double MolarArea ()

Return the van der Waals area of the molecule.

· bool HaveHS ()

Returns true if enthalpy and entropy are known.

bool HaveEnergy ()

Returns true if the Gibb's energy is known.

bool isRegistered ()

Returns true if the molecule has been registered.

• double Enthalpy ()

Return the formation enthalpy of the molecule.

double Entropy ()

Return the formation entropy of the molecule.

• double Energy ()

Return the Gibb's formation energy of the molecule.

• std::string MoleculeName ()

Return the common name of the molecule.

• std::string MolecularFormula ()

Return the molecular formula of the molecule.

• std::string MoleculePhase ()

Return the phase of the molecule.

• int MoleculePhaseID ()

Return the enum phase ID of the molecule.

void DisplayInfo ()

Function to display molecule information.

Protected Attributes

· int charge

Ionic charge of the molecule - specified.

· double molar weight

Molar weight of the molecule (g/mol) - determined from atoms or specified.

double molar_volume

van der Waals Volume of the molecule (cubic angstroms) - determined from atoms or specified

· double molar area

van der Waals Area of the molecule (square angstroms) - determined from atoms or specified

· double formation enthalpy

Enthalpy of formation of the molecule (J/mol) - constant.

double formation_entropy

Entropy of formation of the molecule (J/K/mol) - constant.

· double formation_energy

Gibb's energy of formation (J/mol) - given.

std::string Phase

Phase of the molecule (i.e. Solid, Liquid, Aqueous, Gas...)

int PhaseID

Phase ID of the molecule (from the enum)

std::vector< Atom > atoms

Atoms which make up the molecule - based on Formula.

Private Attributes

std::string Name

Name of the Molecule - Common Name (i.e. H2O = Water)

std::string Formula

Formula for the molecule - specified (i.e. H2O)

bool haveG

True = given Gibb's energy of formation.

bool haveHS

True = give enthalpy and entropy of formation.

· bool registered

True = the object was registered.

5.27.1 Detailed Description

C++ Molecule Object built from Atom Objects (click Molecule to go to function definitions)

C++ Class Object that stores information and certain operations associated with molecules. Registered molecules are built up from their respective atoms so that the molecule can keep track of information such as molecular weight and oxidation states. Primarily, this object is used in conjunction with shark.h to formulate the system of equations necessary for solving speciation type problems in aqueous systems. However, this object is generalized enough to be of use in RedOx calculations, reaction formulation, and molecular transformations.

All information for a molecule should be initialized prior to performing operations with or on the object. There are several molecules already defined for construction by the formulas listed at the top of this section.

```
5.27.2 Constructor & Destructor Documentation
```

5.27.2.1 Molecule::Molecule ()

Default Constructor (builds an empty molecule object)

5.27.2.2 Molecule::∼Molecule ()

Default Destructor (clears out memory)

5.27.2.3 Molecule::Molecule (int *charge*, double *enthalpy*, double *entropy*, double *energy*, bool *HS*, bool *G*, std::string *Phase*, std::string *Name*, std::string *Formula*, std::string *lin_formula*)

Construct any molecule from the available information.

This constructor will build a user defined custom molecule.

Parameters

charge	the ionic charge of the molecule
enthalpy	the standard formation enthalpy of the molecule (J/mol)
entropy	the standard formation entropy of the molecule (J/K/mol)
energy	the standard Gibb's Free Energy of formation of the molecule (J/mol)
HS	boolean to be set to true if enthalpy and entropy were given
G	boolean to be set to true if the energy was given
Phase	string denoting molecule's phase (i.e., Liquid, Aqueous, Gas, Solid)
Name	string denoting the common name of the molecule (i.e., H2O -> Water)
Formula	string denoting the formula by which the molecule is referened (i.e., CI - (aq))
lin_formula	string denoting all the atoms in the molecule (i.e., UO2(OH)2 -> UO4H2)

5.27.3 Member Function Documentation

5.27.3.1 void Molecule::Register (int *charge*, double *enthalpy*, double *entropy*, double *energy*, bool *HS*, bool *G*, std::string *Phase*, std::string *Name*, std::string *Formula*, std::string *lin_formula*)

Function to register this molecule from the available information.

This function will build a user defined custom molecule.

Parameters

charge	the ionic charge of the molecule
enthalpy	the standard formation enthalpy of the molecule (J/mol)
entropy	the standard formation entropy of the molecule (J/K/mol)
energy	the standard Gibb's Free Energy of formation of the molecule (J/mol)
HS	boolean to be set to true if enthalpy and entropy were given
G	boolean to be set to true if the energy was given
Phase	string denoting molecule's phase (i.e., Liquid, Aqueous, Gas, Solid)
Name	string denoting the common name of the molecule (i.e., H2O -> Water)
Formula	string denoting the formula by which the molecule is referened (i.e., CI - (aq))
lin_formula	string denoting all the atoms in the molecule (i.e., UO2(OH)2 -> UO4H2)

5.27.3.2 void Molecule::Register (std::string formula)

Function to register this molecule based on the given formula (if formula is in library)

This function will create this molecule object from the given formula, but only if that formula is already registered in the library. See the top of this class section for a list of all currently registered formulas.

Note

The formula is checked against a known set of molecules inside of the registration function If the formula is unknown, an error will print to the screen. Unknown molecules should be registered using the full registration function from above. The library can only be added to by a going in and editing the source code of the mola.cpp file. However, this is a relatively simple task.

5.27.3.3 void Molecule::setFormula (std::string form)

Sets the formula for a molecule.

5.27.3.4 void Molecule::calculateMolarWeight ()

Forces molecule to calculate its molar weight.

5.27.3.5 void Molecule::calculateMolarVolume ()

Force molecule to calculate van der Waals volume.

5.27.3.6 void Molecule::calculateMolarArea ()

Force molecule to calculate van der Waals area.

5.27.3.7 void Molecule::setMolarWeigth (double mw)

Set the molar weight of species to a constant.

5.27.3.8 void Molecule::setMolarVolume (double v)

Set the van der Waals volume of the species to a constant.

5.27.3.9 void Molecule::setMolarArea (double a)

Set the van der Waals area of the species to a constant.

5.27.3.10 void Molecule::editCharge (int c)

Change the ionic charge of a molecule.

5.27.3.11 void Molecule::editOneOxidationState (int state, std::string Symbol)

Change oxidation state of one of the given atoms (always first match found)

This function will search the list of Atoms that make up the Molecule for the given atomic Symbol. It will change the oxidation state of the first found matching atom with the given state.

5.27.3.12 void Molecule::editAllOxidationStates (int state, std::string Symbol)

Change oxidation state of all of the given atoms.

This function will search the list of Atoms that make up the Molecule for the given atomic Symbol. It will change the oxidation state of all found matching atoms with the given state.

5.27.3.13 void Molecule::calculateAvgOxiState (std::string Symbol)

Function to calculate the average oxidation state of the atoms.

This function search the atoms in the molecule for the matching atomic Symbol. It then looks at all oxidation states of that atom in the molecule and then sets all the oxidation states of that atom to the average value calculated.

5.27.3.14 void Molecule::editEnthalpy (double enthalpy)

Edit the molecules formation enthalpy (J/mol)

5.27.3.15 void Molecule::editEntropy (double entropy)

Edit the molecules formation entropy (J/K/mol)

5.27.3.16 void Molecule::editHS (double H, double S)

Edit both formation enthalpy and entropy.

This function will change or set the values for formation enthalpy (J/mol) and formation entropy (J/K/mol) based on the given values.

Parameters

Н	formation enthalpy (J/mol)
S	formation entropy (J/K/mol)

5.27.3.17 void Molecule::editEnergy (double energy)

Edit Gibb's formation energy.

5.27.3.18 void Molecule::removeOneAtom (std::string Symbol)

Removes one atom of the symbol given (always the first atom found)

5.27.3.19 void Molecule::removeAllAtoms (std::string Symbol)

Removes all atoms of the symbol given.

5.27.3.20 int Molecule::Charge ()

Return the charge of the molecule.

5.27.3.21 double Molecule::MolarWeight ()

Return the molar weight of the molecule.

5.27.3.22 double Molecule::MolarVolume ()

Return the van der Waals volume of the molecule.

5.27.3.23 double Molecule::MolarArea ()

Return the van der Waals area of the molecule.

5.27.3.24 bool Molecule::HaveHS ()

Returns true if enthalpy and entropy are known.

5.27.3.25 bool Molecule::HaveEnergy ()

Returns true if the Gibb's energy is known.

5.27.3.26 bool Molecule::isRegistered ()

Returns true if the molecule has been registered.

5.27.3.27 double Molecule::Enthalpy ()

Return the formation enthalpy of the molecule.

```
5.27.3.28 double Molecule::Entropy ( )
Return the formation entropy of the molecule.
5.27.3.29 double Molecule::Energy ( )
Return the Gibb's formation energy of the molecule.
5.27.3.30 std::string Molecule::MoleculeName ( )
Return the common name of the molecule.
5.27.3.31 std::string Molecule::MolecularFormula ( )
Return the molecular formula of the molecule.
5.27.3.32 std::string Molecule::MoleculePhase ( )
Return the phase of the molecule.
5.27.3.33 int Molecule::MoleculePhaseID ( )
Return the enum phase ID of the molecule.
5.27.3.34 void Molecule::DisplayInfo ( )
Function to display molecule information.
5.27.4 Member Data Documentation
5.27.4.1 int Molecule::charge [protected]
lonic charge of the molecule - specified.
5.27.4.2 double Molecule::molar_weight [protected]
Molar weight of the molecule (g/mol) - determined from atoms or specified.
5.27.4.3 double Molecule::molar_volume [protected]
van der Waals Volume of the molecule (cubic angstroms) - determined from atoms or specified
5.27.4.4 double Molecule::molar_area [protected]
van der Waals Area of the molecule (square angstroms) - determined from atoms or specified
5.27.4.5 double Molecule::formation_enthalpy [protected]
Enthalpy of formation of the molecule (J/mol) - constant.
```

```
5.27.4.6 double Molecule::formation_entropy [protected]
Entropy of formation of the molecule (J/K/mol) - constant.
5.27.4.7 double Molecule::formation_energy [protected]
Gibb's energy of formation (J/mol) - given.
5.27.4.8 std::string Molecule::Phase [protected]
Phase of the molecule (i.e. Solid, Liquid, Aqueous, Gas...)
5.27.4.9 int Molecule::PhaseID [protected]
Phase ID of the molecule (from the enum)
5.27.4.10 std::vector < Atom > Molecule::atoms [protected]
Atoms which make up the molecule - based on Formula.
5.27.4.11 std::string Molecule::Name [private]
Name of the Molecule - Common Name (i.e. H2O = Water)
5.27.4.12 std::string Molecule::Formula [private]
Formula for the molecule - specified (i.e. H2O)
5.27.4.13 bool Molecule::haveG [private]
True = given Gibb's energy of formation.
5.27.4.14 bool Molecule::haveHS [private]
True = give enthalpy and entropy of formation.
5.27.4.15 bool Molecule::registered [private]
True = the object was registered.
The documentation for this class was generated from the following file:
    • mola.h
```

5.28 MONKFISH_DATA Struct Reference

Primary data structure for running MONKFISH.

```
#include <monkfish.h>
```

Public Attributes

• unsigned long int total_steps = 0 Total number of steps taken by the algorithm (iterations and time steps) • double time old = 0.0 Old value of time in the simulation (hrs) • double time = 0.0 Current value of time in the simulation (hrs) • bool Print2File = true True = results to .txt; False = no printing. • bool Print2Console = true True = results to console; False = no printing. • bool DirichletBC = true False = uses film mass transfer for BC, True = Dirichlet BC. • bool NonLinear = false False = Solve directly, True = Solve iteratively. • bool haveMinMax = false True = know min and max fiber density, False = only know avg density (Used in ICs) • bool MultiScale = true True = solve single fiber model at nodes, False = solve equilibrium at nodes. • int level = 2 Level of coupling between multiple scales (default = 2) • double t counter = 0.0 Counter for the time output. double t_print Print output at every t_print time (hrs) int NumComp Number of species to track. · double end time Units: hours. · double total sorption old Old total adsorption per mass of woven nest (mg/g) double total_sorption Current total adsorption per mass woven nest (mg/g) · double single_fiber_density Units: g/L. double avg_fiber_density Units: g/L (Used in ICs) · double max_fiber_density Units: g/L (Used in ICs) · double min_fiber_density Units: g/L (Used in ICs) · double max_porosity Units: -. · double min porosity Units: -. · double domain diameter Nominal diameter of the woven fiber ball - Units: cm. FILE * Output

Output file pointer for printing to text file.

• double(* eval_eps)(int i, int I, const void *user_data)

Function pointer to evaluate the porosity of the woven bundle of fibers.

double(* eval_rho)(int i, int I, const void *user_data)

Function pointer to evaluate the fiber density in the domain.

double(* eval_Dex)(int i, int I, const void *user_data)

Function pointer to evaluate the interparticle diffusivity.

double(* eval_ads)(int i, int I, const void *user_data)

Function pointer to evaluate the adsorption strength for the macro-scale.

double(* eval Ret)(int i, int I, const void *user data)

Function pointer to evaluate the retardation coefficient for the macro-scale.

double(* eval_Cex)(int i, const void *user_data)

Function pointer to evaluate the exterior concentration for the domain.

double(* eval_kf)(int i, const void *user_data)

Function pointer to evalutate the film mass transfer coefficient for the macro-scale.

const void * user_data

User supplied data function to evaluate the function pointers (Default = MONKFISH_DATA)

std::vector< FINCH DATA > finch dat

FINCH data structures to solve each species interparticle diffusion equation.

• std::vector< MONKFISH PARAM > param dat

MONKFISH parameter data structure for each species adsorbing.

std::vector< DOGFISH_DATA > dog_dat

DOGFISH data structures for each node in the macro-scale problem.

5.28.1 Detailed Description

Primary data structure for running MONKFISH.

C-style object holding simulation information for MONKFISH as well as common system parameters like fiber density, fiber diameter, fiber length, etc. This object also contains function pointers to different parameter evaluation functions that can be changed to suit a particular problem. Default functions will be given, so not every user needs to override these functions. This structure also contains vectors of other objects including FINCH and DOGFISH objects to resolve the diffusion physics at both the macro- and micro-scale.

```
5.28.2 Member Data Documentation
```

5.28.2.1 unsigned long int MONKFISH DATA::total steps = 0

Total number of steps taken by the algorithm (iterations and time steps)

5.28.2.2 double MONKFISH_DATA::time_old = 0.0

Old value of time in the simulation (hrs)

5.28.2.3 double MONKFISH_DATA::time = 0.0

Current value of time in the simulation (hrs)

5.28.2.4 bool MONKFISH_DATA::Print2File = true

True = results to .txt; False = no printing.

5.28.2.5 bool MONKFISH_DATA::Print2Console = true

True = results to console; False = no printing.

5.28.2.6 bool MONKFISH_DATA::DirichletBC = true

False = uses film mass transfer for BC, True = Dirichlet BC.

5.28.2.7 bool MONKFISH_DATA::NonLinear = false

False = Solve directly, True = Solve iteratively.

5.28.2.8 bool MONKFISH_DATA::haveMinMax = false

True = know min and max fiber density, False = only know avg density (Used in ICs)

5.28.2.9 bool MONKFISH_DATA::MultiScale = true

True = solve single fiber model at nodes, False = solve equilibrium at nodes.

5.28.2.10 int MONKFISH_DATA::level = 2

Level of coupling between multiple scales (default = 2)

5.28.2.11 double MONKFISH_DATA::t_counter = 0.0

Counter for the time output.

5.28.2.12 double MONKFISH_DATA::t_print

Print output at every t_print time (hrs)

5.28.2.13 int MONKFISH_DATA::NumComp

Number of species to track.

5.28.2.14 double MONKFISH_DATA::end_time

Units: hours.

5.28.2.15 double MONKFISH_DATA::total_sorption_old

Old total adsorption per mass of woven nest (mg/g)

5.28.2.16 double MONKFISH_DATA::total_sorption

Current total adsorption per mass woven nest (mg/g)

```
5.28.2.17 double MONKFISH_DATA::single_fiber_density
Units: g/L.
5.28.2.18 double MONKFISH_DATA::avg_fiber_density
Units: g/L (Used in ICs)
5.28.2.19 double MONKFISH_DATA::max_fiber_density
Units: g/L (Used in ICs)
5.28.2.20 double MONKFISH_DATA::min_fiber_density
Units: g/L (Used in ICs)
5.28.2.21 double MONKFISH_DATA::max_porosity
Units: -.
5.28.2.22 double MONKFISH_DATA::min_porosity
Units: -.
5.28.2.23 double MONKFISH_DATA::domain_diameter
Nominal diameter of the woven fiber ball - Units: cm.
5.28.2.24 FILE* MONKFISH_DATA::Output
Output file pointer for printing to text file.
5.28.2.25 double(* MONKFISH_DATA::eval_eps) (int i, int I, const void *user_data)
Function pointer to evaluate the porosity of the woven bundle of fibers.
5.28.2.26 double(* MONKFISH_DATA::eval_rho) (int i, int l, const void *user_data)
Function pointer to evaluate the fiber density in the domain.
5.28.2.27 double(* MONKFISH_DATA::eval_Dex) (int i, int I, const void *user_data)
Function pointer to evaluate the interparticle diffusivity.
5.28.2.28 double(* MONKFISH_DATA::eval_ads) (int i, int I, const void *user_data)
Function pointer to evaluate the adsorption strength for the macro-scale.
```

5.28.2.29 double(* MONKFISH_DATA::eval_Ret) (int i, int I, const void *user_data)

Function pointer to evaluate the retardation coefficient for the macro-scale.

5.28.2.30 double(* MONKFISH_DATA::eval_Cex) (int i, const void *user_data)

Function pointer to evaluate the exterior concentration for the domain.

5.28.2.31 double(* MONKFISH_DATA::eval_kf) (int i, const void *user_data)

Function pointer to evalutate the film mass transfer coefficient for the macro-scale.

5.28.2.32 const void* MONKFISH_DATA::user_data

User supplied data function to evaluate the function pointers (Default = MONKFISH_DATA)

5.28.2.33 std::vector<FINCH_DATA> MONKFISH_DATA::finch_dat

FINCH data structures to solve each species interparticle diffusion equation.

5.28.2.34 std::vector<MONKFISH_PARAM> MONKFISH_DATA::param_dat

MONKFISH parameter data structure for each species adsorbing.

5.28.2.35 std::vector<DOGFISH_DATA> MONKFISH_DATA::dog_dat

DOGFISH data structures for each node in the macro-scale problem.

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

· monkfish.h

5.29 MONKFISH_PARAM Struct Reference

Data structure for species specific information and parameters.

#include <monkfish.h>

Public Attributes

· double interparticle_diffusion

Units: cm[^]2/hr.

· double exterior_concentration

Units: mol/L.

double exterior_transfer_coeff

Units: cm/hr.

• double sorbed_molefraction

Units: -.

· double initial_sorption

Units: mg/g.

· double sorption_bc

Units: mg/g.

· double intraparticle_diffusion

Units: um^2/hr .

· double film_transfer_coeff

Units: um/hr.

Matrix< double > avg_sorption

Units: mg/g.

Matrix< double > avg_sorption_old

Units: mg/g.Molecule species

Species in the liquid phase.

5.29.1 Detailed Description

Data structure for species specific information and parameters.

C-style object to hold information associated with the different species present in the interparticle diffusion problem. Each species may have different diffusivities, mass transfer coefficients, etc. Average adsorption for each species will be held in matrix objects.

5.29.2 Member Data Documentation

 $5.29.2.1 \quad double \ MONKFISH_PARAM:: interparticle_diffusion$

Units: cm²/hr.

5.29.2.2 double MONKFISH_PARAM::exterior_concentration

Units: mol/L.

 $5.29.2.3 \quad double \ MONKFISH_PARAM:: exterior_transfer_coeff$

Units: cm/hr.

5.29.2.4 double MONKFISH_PARAM::sorbed_molefraction

Units: -.

5.29.2.5 double MONKFISH_PARAM::initial_sorption Units: mg/g. 5.29.2.6 double MONKFISH_PARAM::sorption_bc Units: mg/g. 5.29.2.7 double MONKFISH_PARAM::intraparticle_diffusion Units: um^2/hr. 5.29.2.8 double MONKFISH_PARAM::film_transfer_coeff Units: um/hr. 5.29.2.9 Matrix<double> MONKFISH_PARAM::avg_sorption Units: mg/g. 5.29.2.10 Matrix<double> MONKFISH_PARAM::avg_sorption_old Units: mg/g. 5.29.2.11 Molecule MONKFISH_PARAM::species Species in the liquid phase. The documentation for this struct was generated from the following file: · monkfish.h 5.30 mSPD_DATA Struct Reference MSPD Data Structure. #include <magpie.h> **Public Attributes** • double s

Area shape factor.

double v

van der Waals Volume (cm[^]3/mol)

• double eMax

Maximum lateral interaction energy (J/mol)

• std::vector < double > eta

Binary interaction parameter matrix (i,j)

· double gama

Activity coefficient calculated from mSPD.

5.30.1 Detailed Description MSPD Data Structure. C-Style object holding all parameter information associated with the Modified Spreading Pressure Dependent (SPD) activity model. Each species in the gas phase will have one of these objects. 5.30.2 Member Data Documentation 5.30.2.1 double mSPD_DATA::s Area shape factor. 5.30.2.2 double mSPD_DATA::v van der Waals Volume (cm³/mol) 5.30.2.3 double mSPD_DATA::eMax Maximum lateral interaction energy (J/mol) 5.30.2.4 std::vector<double> mSPD_DATA::eta

Binary interaction parameter matrix (i,j)

5.30.2.5 double mSPD_DATA::gama

Activity coefficient calculated from mSPD.

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

· magpie.h

5.31 MultiligandAdsorption Class Reference

Multi-ligand Adsorption Reaction Object.

#include <shark.h>

Public Member Functions

MultiligandAdsorption ()

Default Constructor.

→MultiligandAdsorption ()

Default Destructor.

- void Initialize_Object (MasterSpeciesList &List, int I, std::vector< int > n)

Function to call the initialization of objects sequentially.

void modifyDeltas (MassBalance &mbo)

Modify the Deltas in the MassBalance Object.

• int setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object in each ligand object.

int checkAqueousIndices ()

Function to check and report errors in the aqueous species indices.

 void setActivityModelInfo (int(*act)(const Matrix< double > &logq, Matrix< double > &activity, const void *data), const void *act data)

Function to set the surface activity model and data pointer.

int setAqueousIndexAuto ()

Automatically sets the primary aqueous species index based on reactions for each ligand.

void setActivityEnum (int act)

Set the activity enum to the value of act.

void setMolarFactor (int ligand, int rxn, double m)

Set the molar factor for the rxn reaction of the ligand ligand to a value of m.

void setVolumeFactor (int i, double v)

Set all ith volume factors for the species list (cm[^]3/mol)

• void setAreaFactor (int i, double a)

Set all ith area factors for the species list (m²/mol)

• void setSpecificMolality (int ligand, double a)

Set the specific molality for the ligand (mol/kg)

- void setSurfaceCharge (int ligand, double c)
- void setAdsorbentName (std::string name)

Set the name of the adsorbent material or particle.

• void setLigandName (int i, std::string name)

Set the name of the ith ligand.

void setSpecificArea (double area)

Set the specific area of the adsorbent.

void setTotalMass (double mass)

Set the mass of the adsorbent.

void setTotalVolume (double volume)

Set the total volume of the system.

void setSurfaceChargeBool (bool opt)

Set the surface charge boolean.

void setElectricPotential (double a)

Set the surface electric potential.

void calculateAreaFactors ()

Calculates the area factors used from the van der Waals volumes.

void calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

void setChargeDensity (const Matrix< double > &x)

Calculates and sets the current value of charge density.

void setlonicStrength (const Matrix< double > &x)

Calculates and sets the current value of ionic strength.

int callSurfaceActivity (const Matrix< double > &x)

Calls the activity model and returns an int flag for success or failure.

• void calculateElecticPotential (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

• double calculate Equilibrium Correction (double sigma, double T, double I, double rel_epsilon, int rxn, int ligand)

Function to calculate the correction term for the equilibrium parameter.

double Eval_Residual (const Matrix < double > &x, const Matrix < double > &gama, double T, double rel_←
perm, int rxn, int ligand)

Calculates the residual for the ith reaction and Ith ligand in the system.

AdsorptionReaction & getAdsorptionObject (int i)

Return reference to the adsortpion object corresponding to ligand i.

int getNumberLigands ()

Get the number of ligands involved with the surface.

• int getActivityEnum ()

Get the value of the activity enum set by user.

double getActivity (int i)

Get the ith activity coefficient from the matrix object.

double getSpecificArea ()

Get the specific area of the adsorbent (m^2/kg) or (mol/kg)

• double getBulkDensity ()

Calculate and return bulk density of adsorbent in system (kg/L)

double getTotalMass ()

Get the total mass of adsorbent in the system (kg)

double getTotalVolume ()

Get the total volume of the system (L)

double getChargeDensity ()

Get the value of the surface charge density (C/m^2)

• double getlonicStrength ()

Get the value of the ionic strength of solution (mol/L)

double getElectricPotential ()

Get the value of the electric surface potential (V)

• bool includeSurfaceCharge ()

Returns true if we are considering surface charging during adsorption.

std::string getLigandName (int i)

Get the name of the ligand object indexed by i.

• std::string getAdsorbentName ()

Get the name of the adsorbent.

Protected Attributes

MasterSpeciesList * List

Pointer to the MasterSpeciesList object.

· int num ligands

Number of different ligands to consider.

std::string adsorbent_name

Name of the adsorbent.

int(* surface activity)(const Matrix< double > &logg, Matrix< double > &activity, const void *data)

Pointer to a surface activity model.

• const void * activity_data

Pointer to the data structure needed for surface activities.

int act_fun

Enumeration to represent the choosen surface activity function.

Matrix< double > activities

List of the activities calculated by the activity model.

double specific_area

Specific surface area of the adsorbent (m^2/kg)

· double total mass

Total mass of the adsorbent in the system (kg)

· double total volume

Total volume of the system (L)

· double ionic_strength

Ionic Strength of the system used to adjust equilibria constants (mol/L)

· double charge_density

Surface charge density of the adsorbent used to adjust equilbria (C/m²)

• double electric_potential

Electric surface potential of the adsorbent used to adjust equilibria (V)

· bool IncludeSurfCharge

True = Includes surface charging corrections, False = Does not consider surface charge.

Private Attributes

std::vector< AdsorptionReaction > ligand obj

List of the ligands and reactions they have on the surface.

5.31.1 Detailed Description

Multi-ligand Adsorption Reaction Object.

C++ Object to handle data and functions associated with forumlating multi-ligand adsorption reactions in a aqueous mixture. Each unique surface in a system will require an instance of this structure. This object is made from a vector of AdsorptionReaction objects, but differentiate between different ligands that exist on the surface.

5.31.2 Constructor & Destructor Documentation

5.31.2.1 MultiligandAdsorption::MultiligandAdsorption ()

Default Constructor.

5.31.2.2 MultiligandAdsorption:: ~MultiligandAdsorption ()

Default Destructor.

5.31.3 Member Function Documentation

5.31.3.1 void MultiligandAdsorption::Initialize_Object (MasterSpeciesList & List, int I, std::vector< int > n)

Function to call the initialization of objects sequentially.

Function will initialize each ligand adsorption object.

Parameters

List	reference to MasterSpeciesList object
1	number of ligands on the surface
n	number of reactions for each ligand (ligands must be correctly indexed)

5.31.3.2 void MultiligandAdsorption::modifyDeltas (MassBalance & mbo)

Modify the Deltas in the MassBalance Object.

This function will take a mass balance object as an argument and modify the deltas in that object to correct for how adsorption affects that particular mass balance. Since adsorption can effect multiple mass balances, this function must be called for each mass balance in the system.

Parameters

mbo	reference to the MassBalance Object the adsorption is acting on
-----	---

5.31.3.3 int MultiligandAdsorption::setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object in each ligand object.

This function searches through the Reaction objects in AdsorptionReaction to find the solid species and their indices to set that information in the adsorb_index structure. That information will be used later to approximate maximum capacities and equilibrium parameters for use in a modified extended Langmuir type expression. Function will return 0 if successful and -1 on a failure.

5.31.3.4 int MultiligandAdsorption::checkAqueousIndices ()

Function to check and report errors in the aqueous species indices.

5.31.3.5 void MultiligandAdsorption::setActivityModelInfo (int(*)(const Matrix< double > &logq, Matrix< double > &activity, const void *data) act, const void * act_data)

Function to set the surface activity model and data pointer.

This function will setup the surface activity model based on the given pointer arguments. If no arguments are given, or are given as NULL, then the activity model will default to ideal solution assumption.

5.31.3.6 int MultiligandAdsorption::setAqueousIndexAuto ()

Automatically sets the primary aqueous species index based on reactions for each ligand.

This function will go through all species and all reactions in each adsorption object and automatically set the primary aqueous species index based on the stoicheometry of the reaction. It will also check and make sure that the primary aqueous index species appears opposite of the adsorbed species in the reactions. Note: This function assumes that the adsorbed indices have already been set.

5.31.3.7 void MultiligandAdsorption::setActivityEnum (int act)

Set the activity enum to the value of act.

```
5.31.3.8 void MultiligandAdsorption::setMolarFactor (int ligand, int rxn, double m)
Set the molar factor for the rxn reaction of the ligand ligand to a value of m.
5.31.3.9 void MultiligandAdsorption::setVolumeFactor (int i, double v)
Set all ith volume factors for the species list (cm<sup>^3</sup>/mol)
5.31.3.10 void MultiligandAdsorption::setAreaFactor (int i, double a)
Set all ith area factors for the species list (m<sup>2</sup>/mol)
5.31.3.11 void MultiligandAdsorption::setSpecificMolality (int ligand, double a)
Set the specific molality for the ligand (mol/kg)
5.31.3.12 void MultiligandAdsorption::setSurfaceCharge (int ligand, double c)
5.31.3.13 void MultiligandAdsorption::setAdsorbentName ( std::string name )
Set the name of the adsorbent material or particle.
5.31.3.14 void MultiligandAdsorption::setLigandName ( int i, std::string name )
Set the name of the ith ligand.
5.31.3.15 void MultiligandAdsorption::setSpecificArea ( double area )
Set the specific area of the adsorbent.
5.31.3.16 void MultiligandAdsorption::setTotalMass ( double mass )
Set the mass of the adsorbent.
5.31.3.17 void MultiligandAdsorption::setTotalVolume ( double volume )
Set the total volume of the system.
5.31.3.18 void MultiligandAdsorption::setSurfaceChargeBool (bool opt)
Set the surface charge boolean.
5.31.3.19 void MultiligandAdsorption::setElectricPotential ( double a )
Set the surface electric potential.
5.31.3.20 void MultiligandAdsorption::calculateAreaFactors ( )
Calculates the area factors used from the van der Waals volumes.
```

5.31.3.21 void MultiligandAdsorption::calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

5.31.3.22 void MultiligandAdsorption::setChargeDensity (const Matrix< double > & x)

Calculates and sets the current value of charge density.

5.31.3.23 void MultiligandAdsorption::setlonicStrength (const Matrix< double > & x)

Calculates and sets the current value of ionic strength.

5.31.3.24 int MultiligandAdsorption::callSurfaceActivity (const Matrix< double > & x)

Calls the activity model and returns an int flag for success or failure.

5.31.3.25 void MultiligandAdsorption::calculateElecticPotential (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

This function will calculate the electric surface potential of the adsorbent under the current conditions of charge density, temperature, ionic strength, and relative permittivity.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)

5.31.3.26 double MultiligandAdsorption::calculateEquilibriumCorrection (double *sigma*, double *T*, double *I*, double *rel_epsilon*, int *rxn*, int *ligand*)

Function to calculate the correction term for the equilibrium parameter.

This function calculates the correction term that gets applied to the equilibrium parameter to correct for surface charge and charge accumulation/depletion effects. It will call the psi approximation and charge exchange functions, therefore it needs to have those functions arguments passed to it as well.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)
rxn	index of the reaction of interest for the adsorption object
ligand	index of the ligand of interest for the adsorption object

5.31.3.27 double MultiligandAdsorption::Eval_Residual (const Matrix < double > & x, const Matrix < double > & gama, double T, double rel_perm, int rxn, int ligand)

Calculates the residual for the ith reaction and lth ligand in the system.

This function will provide a system residual for the ith reaction object involved in the lth ligand's Adsorption Reaction object. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously. This function will also adjust the equilibrium parameter for the reaction

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
rxn	index of the reaction of interest for the adsorption object
ligand	index of the ligand of interest for the adsorption object

5.31.3.28 AdsorptionReaction& MultiligandAdsorption::getAdsorptionObject (int i)

Return reference to the adsortpion object corresponding to ligand i.

5.31.3.29 int MultiligandAdsorption::getNumberLigands ()

Get the number of ligands involved with the surface.

5.31.3.30 int MultiligandAdsorption::getActivityEnum ()

Get the value of the activity enum set by user.

5.31.3.31 double MultiligandAdsorption::getActivity (int i)

Get the ith activity coefficient from the matrix object.

5.31.3.32 double MultiligandAdsorption::getSpecificArea ()

Get the specific area of the adsorbent (m^2/kg) or (mol/kg)

5.31.3.33 double MultiligandAdsorption::getBulkDensity ()

Calculate and return bulk density of adsorbent in system (kg/L)

5.31.3.34 double MultiligandAdsorption::getTotalMass ()

Get the total mass of adsorbent in the system (kg)

5.31.3.35 double MultiligandAdsorption::getTotalVolume ()

Get the total volume of the system (L)

```
5.31.3.36 double MultiligandAdsorption::getChargeDensity ( )
Get the value of the surface charge density (C/m^{\wedge}2)
5.31.3.37 double MultiligandAdsorption::getlonicStrength ( )
Get the value of the ionic strength of solution (mol/L)
5.31.3.38 double MultiligandAdsorption::getElectricPotential ( )
Get the value of the electric surface potential (V)
5.31.3.39 bool MultiligandAdsorption::includeSurfaceCharge ( )
Returns true if we are considering surface charging during adsorption.
5.31.3.40 std::string MultiligandAdsorption::getLigandName (int i)
Get the name of the ligand object indexed by i.
5.31.3.41 std::string MultiligandAdsorption::getAdsorbentName ( )
Get the name of the adsorbent.
5.31.4 Member Data Documentation
5.31.4.1 MasterSpeciesList* MultiligandAdsorption::List [protected]
Pointer to the MasterSpeciesList object.
5.31.4.2 int MultiligandAdsorption::num_ligands [protected]
Number of different ligands to consider.
5.31.4.3 std::string MultiligandAdsorption::adsorbent_name [protected]
Name of the adsorbent.
5.31.4.4 int(* MultiligandAdsorption::surface_activity) (const Matrix< double > &logq, Matrix< double > &activity, const
         void *data) [protected]
```

This is a function pointer for a surface activity model. The function must accept the log of the surface concentrations as an argument (logq) and provide the activities for each species (activity). The pointer data is used to pass any additional arguments needed.

Pointer to a surface activity model.

Parameters

logq	matrix of the log (base 10) of surface concentrations of all species
activity	matrix of activity coefficients for all surface species (must be overriden)
data	pointer to a data structure needed to calculate activities

5.31.4.5 const void* MultiligandAdsorption::activity_data [protected]

Pointer to the data structure needed for surface activities.

5.31.4.6 int MultiligandAdsorption::act_fun [protected]

Enumeration to represent the choosen surface activity function.

5.31.4.7 Matrix<double> MultiligandAdsorption::activities [protected]

List of the activities calculated by the activity model.

5.31.4.8 double MultiligandAdsorption::specific_area [protected]

Specific surface area of the adsorbent (m²/kg)

5.31.4.9 double MultiligandAdsorption::total_mass [protected]

Total mass of the adsorbent in the system (kg)

5.31.4.10 double MultiligandAdsorption::total_volume [protected]

Total volume of the system (L)

5.31.4.11 double MultiligandAdsorption::ionic_strength [protected]

Ionic Strength of the system used to adjust equilibria constants (mol/L)

5.31.4.12 double MultiligandAdsorption::charge_density [protected]

Surface charge density of the adsorbent used to adjust equilbria (C/m²)

 $\textbf{5.31.4.13} \quad \textbf{double MultiligandAdsorption::electric_potential} \quad \texttt{[protected]}$

Electric surface potential of the adsorbent used to adjust equilibria (V)

5.31.4.14 bool MultiligandAdsorption::IncludeSurfCharge [protected]

True = Includes surface charging corrections, False = Does not consider surface charge.

5.31.4.15 std::vector<AdsorptionReaction> MultiligandAdsorption::ligand_obj [private]

List of the ligands and reactions they have on the surface.

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

· shark.h

5.32 MultiligandChemisorption Class Reference

Multi-ligand Chemisorption Reaction Object.

```
#include <shark.h>
```

Public Member Functions

• MultiligandChemisorption ()

Default Constructor.

∼MultiligandChemisorption ()

Default Destructor.

void Initialize Object (MasterSpeciesList &List, int I, std::vector< int > n)

Function to call the initialization of objects sequentially.

void Display_Info ()

Display the adsorption reaction information.

void modifyMBEdeltas (MassBalance &mbo)

Modify the Deltas in the MassBalance Object.

int setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object in each ligand object.

int setLigandIndices ()

Find and set the ligand species index.

• int setDeltas ()

Find and set all the delta values for the site balance.

 void setActivityModelInfo (int(*act)(const Matrix< double > &logq, Matrix< double > &activity, const void *data), const void *act_data)

Function to set the surface activity model and data pointer.

void setActivityEnum (int act)

Set the activity enum to the value of act.

void setVolumeFactor (int i, double v)

Set all ith volume factors for the species list (cm[^]3/mol)

• void setAreaFactor (int i, double a)

Set all ith area factors for the species list (m^2/mol)

void setSpecificMolality (int ligand, double a)

Set the specific molality for the ligand (mol/kg)

void setAdsorbentName (std::string name)

Set the name of the adsorbent material or particle.

void setLigandName (int ligand, std::string name)

Set the name of the ith ligand.

void setSpecificArea (double area)

Set the specific area of the adsorbent.

void setTotalMass (double mass)

Set the mass of the adsorbent.

• void setTotalVolume (double volume)

Set the total volume of the system.

void setSurfaceChargeBool (bool opt)

Set the surface charge boolean.

void setElectricPotential (double a)

Set the surface electric potential.

void calculateAreaFactors ()

Calculates the area factors used from the van der Waals volumes.

void calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

void setChargeDensity (const Matrix< double > &x)

Calculates and sets the current value of charge density.

void setlonicStrength (const Matrix< double > &x)

Calculates and sets the current value of ionic strength.

int callSurfaceActivity (const Matrix< double > &x)

Calls the activity model and returns an int flag for success or failure.

void calculateElecticPotential (double sigma, double T, double I, double rel epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

· double calculateEquilibriumCorrection (double sigma, double T, double I, double rel epsilon, int rxn, int ligand)

Function to calculate the correction term for the equilibrium parameter.

double Eval_RxnResidual (const Matrix< double > &x, const Matrix< double > &gama, double T, double rel perm, int rxn, int ligand)

Calculates the residual for the ith reaction and Ith ligand in the system.

double Eval_SiteBalanceResidual (const Matrix< double > &x, int ligand)

Calculates the residual for the overall site balance for a given ligand.

ChemisorptionReaction & getChemisorptionObject (int ligand)

Return reference to the adsortpion object corresponding to the ligand.

int getNumberLigands ()

Get the number of ligands involved with the surface.

int getActivityEnum ()

Get the value of the activity enum set by user.

• double getActivity (int i)

Get the ith activity coefficient from the matrix object.

double getSpecificArea ()

Get the specific area of the adsorbent (m^2/kg) or (mol/kg)

double getBulkDensity ()

Calculate and return bulk density of adsorbent in system (kg/L)

double getTotalMass ()

Get the total mass of adsorbent in the system (kg)

double getTotalVolume ()

Get the total volume of the system (L)

double getChargeDensity ()

Get the value of the surface charge density (C/m^2)

• double getlonicStrength ()

Get the value of the ionic strength of solution (mol/L)

double getElectricPotential ()

Get the value of the electric surface potential (V)

bool includeSurfaceCharge ()

Returns true if we are considering surface charging during adsorption.

std::string getLigandName (int ligand)

Get the name of the ligand object indexed by ligand.

std::string getAdsorbentName ()

Get the name of the adsorbent.

Protected Attributes

MasterSpeciesList * List

Pointer to the MasterSpeciesList object.

int num_ligands

Number of different ligands to consider.

· std::string adsorbent_name

Name of the adsorbent.

int(* surface_activity)(const Matrix < double > &logq, Matrix < double > &activity, const void *data)

Pointer to a surface activity model.

const void * activity data

Pointer to the data structure needed for surface activities.

int act_fun

Enumeration to represent the choosen surface activity function.

• Matrix< double > activities

List of the activities calculated by the activity model.

double specific_area

Specific surface area of the adsorbent (m²/kg)

· double total mass

Total mass of the adsorbent in the system (kg)

· double total volume

Total volume of the system (L)

· double ionic_strength

Ionic Strength of the system used to adjust equilibria constants (mol/L)

double charge_density

Surface charge density of the adsorbent used to adjust equilbria (C/m^2)

· double electric potential

Electric surface potential of the adsorbent used to adjust equilibria (V)

bool IncludeSurfCharge

True = Includes surface charging corrections, False = Does not consider surface charge.

Private Attributes

std::vector < ChemisorptionReaction > ligand_obj

List of the ligands and reactions they have on the surface.

5.32.1 Detailed Description

Multi-ligand Chemisorption Reaction Object.

C++ Object to handle data and functions associated with forumlating multi-ligand chemisorption reactions in a aqueous mixture. Each unique surface in a system will require an instance of this structure. This object is made from a vector of ChemisorptionReaction objects, but differentiate between different ligands that exist on the surface. It is based largely off of the original Multiligand Adsorption object, but will include an explict way to handle the site balances associated with each ligand.

5.32.2 Constructor & Destructor Documentation

5.32.2.1 MultiligandChemisorption::MultiligandChemisorption ()

Default Constructor.

5.32.2.2 MultiligandChemisorption:: ~MultiligandChemisorption ()

Default Destructor.

5.32.3 Member Function Documentation

5.32.3.1 void MultiligandChemisorption::Initialize_Object (MasterSpeciesList & List, int I, std::vector< int > n)

Function to call the initialization of objects sequentially.

Function will initialize each ligand adsorption object.

Parameters

List	reference to MasterSpeciesList object
1	number of ligands on the surface
n	number of reactions for each ligand (ligands must be correctly indexed)

5.32.3.2 void MultiligandChemisorption::Display_Info ()

Display the adsorption reaction information.

5.32.3.3 void MultiligandChemisorption::modifyMBEdeltas (MassBalance & mbo)

Modify the Deltas in the MassBalance Object.

This function will take a mass balance object as an argument and modify the deltas in that object to correct for how adsorption affects that particular mass balance. Since adsorption can effect multiple mass balances, this function must be called for each mass balance in the system.

Parameters

mbo reference to the MassBalance Object the adsorption is acting of	mbo	the MassBalance Object the adsorption	is acting on
---	-----	---------------------------------------	--------------

5.32.3.4 int MultiligandChemisorption::setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object in each ligand object.

This function searches through the Reaction objects in ChemisorptionReaction to find the solid species and their indices to set that information in the adsorb_index structure. That information will be used later to approximate maximum capacities and equilibrium parameters for use in a modified extended Langmuir type expression. Function will return 0 if successful and -1 on a failure.

5.32.3.5 int MultiligandChemisorption::setLigandIndices ()

Find and set the ligand species index.

This function searches through the Reaction objects in ChemisorptionReaction to find the ligand species and its index to set that information in the ligand_index structure. Function will return 0 if successful and -1 on a failure.

5.32.3.6 int MultiligandChemisorption::setDeltas ()

Find and set all the delta values for the site balance.

This function searches through all reaction object instances for the stoicheometry of the ligand in each adsorption reaction. That stoicheometry serves as the basis for determining the site balance. NOTE: the delta for the ligand is set automatically in the setLigandIndex() function, so we can ignore that species. In addition, this function must be called after setLigandIndex() and setAdsorbIndices() are called and after the stoicheometry of each reaction has been determined.

5.32.3.7 void MultiligandChemisorption::setActivityModelInfo (int(*)(const Matrix< double > &logq, Matrix< double > &activity, const void *data) act, const void * act_data)

Function to set the surface activity model and data pointer.

This function will setup the surface activity model based on the given pointer arguments. If no arguments are given, or are given as NULL, then the activity model will default to ideal solution assumption.

5.32.3.8 void MultiligandChemisorption::setActivityEnum (int act)

Set the activity enum to the value of act.

5.32.3.9 void MultiligandChemisorption::setVolumeFactor (int i, double v)

Set all ith volume factors for the species list (cm³/mol)

5.32.3.10 void MultiligandChemisorption::setAreaFactor (int i, double a)

Set all ith area factors for the species list (m²/mol)

 $5.32.3.11 \quad \text{void MultiligandChemisorption::setSpecificMolality (} \ \text{int } \textit{ligand, } \ \text{double } \textit{a} \ \text{)}$

Set the specific molality for the ligand (mol/kg)

5.32.3.12 void MultiligandChemisorption::setAdsorbentName (std::string name)

Set the name of the adsorbent material or particle.

5.32.3.13 void MultiligandChemisorption::setLigandName (int ligand, std::string name)

Set the name of the ith ligand.

5.32.3.14 void MultiligandChemisorption::setSpecificArea (double area)

Set the specific area of the adsorbent.

5.32.3.15 void MultiligandChemisorption::setTotalMass (double mass)

Set the mass of the adsorbent.

5.32.3.16 void MultiligandChemisorption::setTotalVolume (double volume)

Set the total volume of the system.

5.32.3.17 void MultiligandChemisorption::setSurfaceChargeBool (bool opt)

Set the surface charge boolean.

5.32.3.18 void MultiligandChemisorption::setElectricPotential (double a)

Set the surface electric potential.

5.32.3.19 void MultiligandChemisorption::calculateAreaFactors ()

Calculates the area factors used from the van der Waals volumes.

5.32.3.20 void MultiligandChemisorption::calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

5.32.3.21 void MultiligandChemisorption::setChargeDensity (const Matrix < double > & x)

Calculates and sets the current value of charge density.

5.32.3.22 void MultiligandChemisorption::setlonicStrength (const Matrix < double > & x)

Calculates and sets the current value of ionic strength.

5.32.3.23 int MultiligandChemisorption::callSurfaceActivity (const Matrix< double > & x)

Calls the activity model and returns an int flag for success or failure.

5.32.3.24 void MultiligandChemisorption::calculateElecticPotential (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

This function will calculate the electric surface potential of the adsorbent under the current conditions of charge density, temperature, ionic strength, and relative permittivity.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)

5.32.3.25 double MultiligandChemisorption::calculateEquilibriumCorrection (double *sigma*, double *T*, double *l*, double *rel_epsilon*, int *rxn*, int *ligand*)

Function to calculate the correction term for the equilibrium parameter.

This function calculates the correction term that gets applied to the equilibrium parameter to correct for surface charge and charge accumulation/depletion effects. It will call the psi approximation and charge exchange functions, therefore it needs to have those functions arguments passed to it as well.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)
rxn	index of the reaction of interest for the adsorption object
ligand	index of the ligand of interest for the adsorption object

5.32.3.26 double MultiligandChemisorption::Eval_RxnResidual (const Matrix < double > & x, const Matrix < double > & gama, double T, double rel_perm, int rxn, int ligand)

Calculates the residual for the ith reaction and Ith ligand in the system.

This function will provide a system residual for the ith reaction object involved in the lth ligand's Adsorption Reaction object. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously. This function will also adjust the equilibrium parameter for the reaction

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
rxn	index of the reaction of interest for the adsorption object
ligand	index of the ligand of interest for the adsorption object

5.32.3.27 double MultiligandChemisorption::Eval_SiteBalanceResidual (const Matrix < double > & x, int ligand)

Calculates the residual for the overall site balance for a given ligand.

This function will provide a system residual for the site/ligand balance for the Chemisorption Reaction object. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
ligand	index of the ligand of interest of the chemisorption object

```
5.32.3.28 ChemisorptionReaction& MultiligandChemisorption::getChemisorptionObject (int ligand)
Return reference to the adsortpion object corresponding to the ligand.
5.32.3.29 int MultiligandChemisorption::getNumberLigands ( )
Get the number of ligands involved with the surface.
5.32.3.30 int MultiligandChemisorption::getActivityEnum ( )
Get the value of the activity enum set by user.
5.32.3.31 double MultiligandChemisorption::getActivity (int i)
Get the ith activity coefficient from the matrix object.
5.32.3.32 double MultiligandChemisorption::getSpecificArea ( )
Get the specific area of the adsorbent (m<sup>2</sup>/kg) or (mol/kg)
5.32.3.33 double MultiligandChemisorption::getBulkDensity ( )
Calculate and return bulk density of adsorbent in system (kg/L)
5.32.3.34 double MultiligandChemisorption::getTotalMass ( )
Get the total mass of adsorbent in the system (kg)
5.32.3.35 double MultiligandChemisorption::getTotalVolume ( )
Get the total volume of the system (L)
5.32.3.36 double MultiligandChemisorption::getChargeDensity ( )
Get the value of the surface charge density (C/m<sup>2</sup>)
5.32.3.37 double MultiligandChemisorption::getlonicStrength ( )
Get the value of the ionic strength of solution (mol/L)
5.32.3.38 double MultiligandChemisorption::getElectricPotential ( )
Get the value of the electric surface potential (V)
5.32.3.39 bool MultiligandChemisorption::includeSurfaceCharge ( )
```

Returns true if we are considering surface charging during adsorption.

5.32.3.40 std::string MultiligandChemisorption::getLigandName (int ligand)

Get the name of the ligand object indexed by ligand.

5.32.3.41 std::string MultiligandChemisorption::getAdsorbentName ()

Get the name of the adsorbent.

5.32.4 Member Data Documentation

5.32.4.1 MasterSpeciesList* MultiligandChemisorption::List [protected]

Pointer to the MasterSpeciesList object.

5.32.4.2 int MultiligandChemisorption::num_ligands [protected]

Number of different ligands to consider.

5.32.4.3 std::string MultiligandChemisorption::adsorbent_name [protected]

Name of the adsorbent.

5.32.4.4 int(* MultiligandChemisorption::surface_activity) (const Matrix< double > &logq, Matrix< double > &activity, const void *data) [protected]

Pointer to a surface activity model.

This is a function pointer for a surface activity model. The function must accept the log of the surface concentrations as an argument (logq) and provide the activities for each species (activity). The pointer data is used to pass any additional arguments needed.

Parameters

logq	matrix of the log (base 10) of surface concentrations of all species
activity	matrix of activity coefficients for all surface species (must be overriden)
data	pointer to a data structure needed to calculate activities

5.32.4.5 const void* MultiligandChemisorption::activity_data [protected]

Pointer to the data structure needed for surface activities.

5.32.4.6 int MultiligandChemisorption::act_fun [protected]

Enumeration to represent the choosen surface activity function.

5.32.4.7 Matrix<double> MultiligandChemisorption::activities [protected]

List of the activities calculated by the activity model.

```
5.32.4.8 double MultiligandChemisorption::specific_area [protected]
Specific surface area of the adsorbent (m<sup>2</sup>/kg)
5.32.4.9 double MultiligandChemisorption::total_mass [protected]
Total mass of the adsorbent in the system (kg)
5.32.4.10 double MultiligandChemisorption::total_volume [protected]
Total volume of the system (L)
5.32.4.11 double MultiligandChemisorption::ionic_strength [protected]
Ionic Strength of the system used to adjust equilibria constants (mol/L)
5.32.4.12 double MultiligandChemisorption::charge_density [protected]
Surface charge density of the adsorbent used to adjust equilbria (C/m<sup>2</sup>)
5.32.4.13 double MultiligandChemisorption::electric_potential [protected]
Electric surface potential of the adsorbent used to adjust equilibria (V)
5.32.4.14 bool MultiligandChemisorption::IncludeSurfCharge [protected]
True = Includes surface charging corrections, False = Does not consider surface charge.
5.32.4.15 std::vector<ChemisorptionReaction>MultiligandChemisorption::ligand_obj [private]
List of the ligands and reactions they have on the surface.
The documentation for this class was generated from the following file:
```

shark.h

5.33 NUM_JAC_DATA Struct Reference

Data structure to form a numerical jacobian matrix with finite differences.

```
#include <lark.h>
```

Public Attributes

```
• double eps = sqrt(DBL_EPSILON)
```

Perturbation value.

Matrix< double > Fx

Vector of function evaluations at x.

Matrix< double > Fxp

Vector of function evaluations at x+eps.

Matrix< double > dxj

Vector of perturbed x values.

5.33.1 Detailed Description

Data structure to form a numerical jacobian matrix with finite differences.

C-style object to be used in conjunction with the Numerical Jacobian algorithm. This algorithm will used double-precision finite-differences to formulate an approximate Jacobian matrix at the given variable state for the given residual/non-linear function.

5.33.2 Member Data Documentation

```
5.33.2.1 double NUM_JAC_DATA::eps = sqrt(DBL_EPSILON)
```

Perturbation value.

```
5.33.2.2 Matrix<double> NUM_JAC_DATA::Fx
```

Vector of function evaluations at x.

```
5.33.2.3 Matrix < double > NUM_JAC_DATA::Fxp
```

Vector of function evaluations at x+eps.

```
5.33.2.4 Matrix<double> NUM_JAC_DATA::dxj
```

Vector of perturbed x values.

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

· lark.h

5.34 OPTRANS_DATA Struct Reference

Data structure for implementation of linear operator transposition.

```
#include <lark.h>
```

Public Attributes

• Matrix< double > li

The ith column vector of the identity operator.

Matrix< double > Ai

The ith column vector of the user's linear operator.

5.34.1 Detailed Description

Data structure for implementation of linear operator transposition.

C-style object used in conjunction with the Operator Transpose algorithm to form an action of $A^{\wedge}T*r$ when A is only available as a linear operator and not a matrix. This is a sub-routine required by GCR and GMRESR to stabilize the outer iterations.

```
5.34.2 Member Data Documentation
```

```
\textbf{5.34.2.1} \quad \textbf{Matrix} \small{<} \textbf{double} \small{>} \textbf{OPTRANS\_DATA::} \textbf{li}
```

The ith column vector of the identity operator.

```
5.34.2.2 Matrix<double> OPTRANS_DATA::Ai
```

The ith column vector of the user's linear operator.

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

· lark.h

5.35 PCG_DATA Struct Reference

Data structure for implementation of the PCG algorithms for symmetric linear systems.

```
#include <lark.h>
```

Public Attributes

• int maxit = 0

Maximum allowable iterations - default = min(vector_size,1000)

• int iter = 0

Actual number of iterations taken.

· double alpha

Step size for new solution.

· double beta

Step size for new search direction.

• double tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

• double tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

• double res

Absolute residual norm.

· double relres

Relative residual norm.

• double relres_base

Initial residual norm.

· double bestres

Best found residual norm.

• bool Output = true

True = print messages to console.

Matrix< double > x

Current solution to the linear system.

Matrix< double > bestx

Best found solution to the linear system.

• Matrix< double > r

Residual vector for the linear system.

• Matrix< double > r_old

Previous residual vector.

Matrix< double > z

Preconditioned residual vector (result of precon function)

Matrix< double > z_old

Previous preconditioned residual vector.

Matrix< double > p

Search direction.

Matrix< double > Ap

Result of matrix-vector multiplication.

5.35.1 Detailed Description

Data structure for implementation of the PCG algorithms for symmetric linear systems.

C-style object used in conjunction with the Preconditioned Conjugate Gradient (PCG) algorithm to iteratively solve a symmetric linear system of equations. This algorithm is optimal if your linear system is symmetric, but will not work at all if your system is asymmetric. For asymmetric systems, use one of the other linear methods.

5.35.2 Member Data Documentation

5.35.2.1 int PCG_DATA::maxit = 0

Maximum allowable iterations - default = min(vector_size,1000)

5.35.2.2 int PCG_DATA::iter = 0

Actual number of iterations taken.

5.35.2.3 double PCG_DATA::alpha

Step size for new solution.

5.35.2.4 double PCG_DATA::beta

Step size for new search direction.

5.35.2.5 double PCG_DATA::tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

5.35.2.6 double PCG_DATA::tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

5.35.2.7 double PCG_DATA::res

Absolute residual norm.

5.35.2.8 double PCG_DATA::relres

Relative residual norm.

5.35.2.9 double PCG_DATA::relres_base

Initial residual norm.

5.35.2.10 double PCG_DATA::bestres

Best found residual norm.

5.35.2.11 bool PCG_DATA::Output = true

True = print messages to console.

5.35.2.12 Matrix<double> PCG_DATA::x

Current solution to the linear system.

5.35.2.13 Matrix < double > PCG_DATA::bestx

Best found solution to the linear system.

5.35.2.14 Matrix<double> PCG_DATA::r

Residual vector for the linear system.

5.35.2.15 Matrix < double > PCG_DATA::r_old

Previous residual vector.

5.35.2.16 Matrix<double> PCG_DATA::z

Preconditioned residual vector (result of precon function)

5.35.2.17 Matrix<double> PCG_DATA::z_old

Previous preconditioned residual vector.

5.35.2.18 Matrix<double> PCG_DATA::p

Search direction.

5.35.2.19 Matrix < double > PCG_DATA::Ap

Result of matrix-vector multiplication.

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

· lark.h

5.36 PeriodicTable Class Reference

Class object that store a digitial copy of all Atom objects.

```
#include <eel.h>
```

Public Member Functions

PeriodicTable ()

Default Constructor - Build Perodic Table.

∼PeriodicTable ()

Default Destructor - Destroy the table.

PeriodicTable (int *n, int N)

Construct a partial table from a list of atomic numbers.

PeriodicTable (std::vector < std::string > &Symbol)

Construct a partial table from a vector of atom symbols.

PeriodicTable (std::vector< int > &n)

Construct a partial table from a vector of atomic numbers.

void DisplayTable ()

Displays the periodic table via symbols.

Protected Attributes

• std::vector < Atom > Table

Storage vector for all atoms in the table.

Private Attributes

• int number_elements

Number of atom objects being stored.

5.36.1 Detailed Description

Class object that store a digitial copy of all Atom objects.

C++ class object to hold digitally registered Atom objects. All registered atoms (Hydrogen to Ununoctium) are stored as in a vector. Currently, this object is unused, but could be modified to be explorable and used as a constant referece for all atoms in the table.

5.36.2 Constructor & Destructor Documentation

5.36.2.1 PeriodicTable::PeriodicTable ()

Default Constructor - Build Perodic Table.

5.36.2.2 PeriodicTable:: ∼PeriodicTable ()

Default Destructor - Destroy the table.

```
5.36.2.3 PeriodicTable::PeriodicTable ( int * n, int N )
Construct a partial table from a list of atomic numbers.
5.36.2.4 PeriodicTable::PeriodicTable ( std::vector < std::string > & Symbol )
Construct a partial table from a vector of atom symbols.
5.36.2.5 PeriodicTable::PeriodicTable ( std::vector < int > & n )
Construct a partial table from a vector of atomic numbers.
5.36.3 Member Function Documentation
5.36.3.1 void PeriodicTable::DisplayTable ( )
Displays the periodic table via symbols.
5.36.4 Member Data Documentation
5.36.4.1 std::vector<Atom> PeriodicTable::Table [protected]
Storage vector for all atoms in the table.
5.36.4.2 int PeriodicTable::number_elements [private]
Number of atom objects being stored.
The documentation for this class was generated from the following file:
    · eel.h
5.37 PICARD_DATA Struct Reference
Data structure for the implementation of a Picard or Fixed-Point iteration for non-linear systems.
```

#include <lark.h>

Public Attributes

• int maxit = 0

Maximum allowable iterations - default = min(3*vec_size,1000)

• int iter = 0

Actual number of iterations.

• double tol rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

• double tol abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

· double res

Residual norm of the iterate.

double relres

Relative residual norm of the iterate.

· double relres_base

Initial residual norm.

· double bestres

Best found residual norm.

• bool Output = true

True = print messages to console.

Matrix< double > x0

Previous iterate solution vector.

Matrix< double > bestx

Best found solution vector.

Matrix< double > r

Residual of the non-linear system.

5.37.1 Detailed Description

Data structure for the implementation of a Picard or Fixed-Point iteration for non-linear systems.

C-style object used in conjunction with the Picard algorithm for solving a non-linear system of equations. This is an extradorinarily simple iterative method by which a weak or loose form of the non-linear system is solved based on an initial guess. User must supplied a residual function for the non-linear system and a function representing the weak solution. Generally, this method is less efficient than Newton methods, but is significantly cheaper.

```
5.37.2 Member Data Documentation
```

```
5.37.2.1 int PICARD DATA::maxit = 0
```

Maximum allowable iterations - default = min(3*vec_size,1000)

5.37.2.2 int PICARD_DATA::iter = 0

Actual number of iterations.

5.37.2.3 double PICARD_DATA::tol_rel = 1e-6

Relative tolerance for convergence - default = 1e-6.

5.37.2.4 double PICARD_DATA::tol_abs = 1e-6

Absolution tolerance for convergence - default = 1e-6.

5.37.2.5 double PICARD_DATA::res

Residual norm of the iterate.

5.37.2.6 double PICARD_DATA::relres

Relative residual norm of the iterate.

5.37.2.7 double PICARD_DATA::relres_base

Initial residual norm.

5.37.2.8 double PICARD_DATA::bestres

Best found residual norm.

5.37.2.9 bool PICARD_DATA::Output = true

True = print messages to console.

5.37.2.10 Matrix < double > PICARD_DATA::x0

Previous iterate solution vector.

5.37.2.11 Matrix < double > PICARD_DATA::bestx

Best found solution vector.

5.37.2.12 Matrix<double> PICARD_DATA::r

Residual of the non-linear system.

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

· lark.h

5.38 PJFNK_DATA Struct Reference

Data structure for the implementation of the PJFNK algorithm for non-linear systems.

#include <lark.h>

Public Attributes

• int nl_iter = 0

Number of non-linear iterations.

• int | iter = 0

Number of linear iterations.

• int fun call = 0

Actual number of function calls made.

int nl maxit = 0

Maximum allowable non-linear steps.

• int linear_solver = -1

Flag to denote which linear solver to use - default = PJFNK Chooses.

double nl_tol_abs = 1e-6

Absolute Convergence tolerance for non-linear system - default = 1e-6.

• double nl_tol_rel = 1e-6

Relative Convergence tol for the non-linear system - default = 1e-6.

double lin_tol_rel = 1e-6

Relative tolerance of the linear solver - default = 1e-6.

• double lin tol abs = 1e-6

Absolute tolerance of the linear solver - default = 1e-6.

· double nl res

Absolute redidual norm for the non-linear system.

· double nl relres

Relative residual for the non-linear system.

• double nl_res_base

Initial residual norm for the non-linear system.

· double nl bestres

Best found residual norm.

double eps =sqrt(DBL_EPSILON)

Value of epsilon used jacvec - default = sqrt(DBL_EPSILON)

• bool NL Output = true

True = print PJFNK messages to console.

bool L_Output = false

True = print Linear messages to console.

• bool LineSearch = false

True = use Backtracking Linesearch for global convergence.

• bool Bounce = false

True = allow Linesearch to go outside local well, False = Strict local convergence.

Matrix< double > F

Stored fuction evaluation at x (also the residual)

Matrix< double > Fv

Stored function evaluation at x+eps*v.

Matrix< double > v

Stored vector of x+eps*v.

Matrix< double > x

Current solution vector for the non-linear system.

Matrix< double > bestx

Best found solution vector to the non-linear system.

· GMRESLP DATA gmreslp dat

Data structure for the GMRESLP method.

• PCG_DATA pcg_dat

Data structure for the PCG method.

BiCGSTAB_DATA bicgstab_dat

Data structure for the BiCGSTAB method.

· CGS DATA cgs dat

Data structure for the CGS method.

GMRESRP_DATA gmresrp_dat

Data structure for the GMRESRP method.

· GCR DATA gcr dat

Data structure for the GCR method.

GMRESR_DATA gmresr_dat

Data structure for the GMRESR method.

· KMS DATA kms dat

Data structure for the KMS method.

• QR_DATA qr_dat

Data structure for the QR solve method.

BACKTRACK_DATA backtrack_dat

Data structure for the Backtracking Linesearch algorithm.

const void * res data

Data structure pointer for user's residual data.

const void * precon_data

Data structure pointer for user's preconditioning data.

int(* funeval)(const Matrix< double > &x, Matrix< double > &F, const void *res data)

Function pointer for the user's function F(x) using there data.

int(* precon)(const Matrix< double > &r, Matrix< double > &p, const void *precon_data)

Function pointer for the user's preconditioning function for the linear system.

5.38.1 Detailed Description

Data structure for the implementation of the PJFNK algorithm for non-linear systems.

C-style object to be used in conjunction with the Preconditioned Jacobian-Free Newton-Krylov (PJFNK) method for solving a non-linear system of equations. You can use any of the Krylov methods listed in the krylov_method enum to solve the linear sub-problem. When FOM is specified as the Krylov method, this algorithm becomes equivalent to an exact Newton method. If no Krylov method is specified, then the algorithm will try to pick a method based on the problem size and availability of preconditioning.

5.38.2 Member Data Documentation

5.38.2.1 int PJFNK_DATA::nl_iter = 0

Number of non-linear iterations.

5.38.2.2 int PJFNK_DATA::I_iter = 0

Number of linear iterations.

5.38.2.3 int PJFNK_DATA::fun_call = 0

Actual number of function calls made.

5.38.2.4 int PJFNK_DATA::nl_maxit = 0

Maximum allowable non-linear steps.

5.38.2.5 int PJFNK_DATA::linear_solver = -1

Flag to denote which linear solver to use - default = PJFNK Chooses.

5.38.2.6 double PJFNK_DATA::nl_tol_abs = 1e-6

Absolute Convergence tolerance for non-linear system - default = 1e-6.

5.38.2.7 double PJFNK_DATA::nl_tol_rel = 1e-6

Relative Convergence tol for the non-linear system - default = 1e-6.

5.38.2.8 double PJFNK_DATA::lin_tol_rel = 1e-6

Relative tolerance of the linear solver - default = 1e-6.

5.38.2.9 double PJFNK_DATA::lin_tol_abs = 1e-6

Absolute tolerance of the linear solver - default = 1e-6.

5.38.2.10 double PJFNK_DATA::nl_res

Absolute redidual norm for the non-linear system.

5.38.2.11 double PJFNK_DATA::nl_relres

Relative residual for the non-linear system.

5.38.2.12 double PJFNK_DATA::nl_res_base

Initial residual norm for the non-linear system.

5.38.2.13 double PJFNK_DATA::nl_bestres

Best found residual norm.

5.38.2.14 double PJFNK_DATA::eps =sqrt(DBL_EPSILON)

Value of epsilon used jacvec - default = sqrt(DBL_EPSILON)

5.38.2.15 bool PJFNK_DATA::NL_Output = true

True = print PJFNK messages to console.

5.38.2.16 bool PJFNK_DATA::L_Output = false

True = print Linear messages to console.

5.38.2.17 bool PJFNK_DATA::LineSearch = false

True = use Backtracking Linesearch for global convergence.

5.38.2.18 bool PJFNK_DATA::Bounce = false

True = allow Linesearch to go outside local well, False = Strict local convergence.

5.38.2.19 Matrix<double> PJFNK_DATA::F

Stored fuction evaluation at x (also the residual)

5.38.2.20 Matrix<double> PJFNK_DATA::Fv

Stored function evaluation at x+eps*v.

5.38.2.21 Matrix<double> PJFNK_DATA::v

Stored vector of x+eps*v.

5.38.2.22 Matrix < double > PJFNK_DATA::x

Current solution vector for the non-linear system.

5.38.2.23 Matrix<double> PJFNK_DATA::bestx

Best found solution vector to the non-linear system.

5.38.2.24 GMRESLP_DATA PJFNK_DATA::gmreslp_dat

Data structure for the GMRESLP method.

5.38.2.25 PCG_DATA PJFNK_DATA::pcg_dat

Data structure for the PCG method.

5.38.2.26 BICGSTAB_DATA PJFNK_DATA::bicgstab_dat

Data structure for the BiCGSTAB method.

5.38.2.27 CGS_DATA PJFNK_DATA::cgs_dat

Data structure for the CGS method.

5.38.2.28 GMRESRP_DATA PJFNK_DATA::gmresrp_dat

Data structure for the GMRESRP method.

5.38.2.29 GCR DATA PJFNK_DATA::gcr_dat

Data structure for the GCR method.

5.38.2.30 GMRESR_DATA PJFNK_DATA::gmresr_dat

Data structure for the GMRESR method.

5.38.2.31 KMS DATA PJFNK_DATA::kms_dat

Data structure for the KMS method.

5.38.2.32 QR_DATA PJFNK_DATA::qr_dat

Data structure for the QR solve method.

5.38.2.33 BACKTRACK_DATA PJFNK_DATA::backtrack_dat

Data structure for the Backtracking Linesearch algorithm.

5.38.2.34 const void* PJFNK_DATA::res_data

Data structure pointer for user's residual data.

5.38.2.35 const void* PJFNK_DATA::precon_data

Data structure pointer for user's preconditioning data.

5.38.2.36 int(* PJFNK_DATA::funeval) (const Matrix < double > &x, Matrix < double > &F, const void *res_data)

Function pointer for the user's function F(x) using there data.

5.38.2.37 int(* PJFNK_DATA::precon) (const Matrix < double > &r, Matrix < double > &p, const void *precon_data)

Function pointer for the user's preconditioning function for the linear system.

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

lark.h

5.39 PURE_GAS Struct Reference

Data structure holding all the parameters for each pure gas spieces.

#include <egret.h>

Public Attributes

· double molecular_weight

Given: molecular weights (g/mol)

double Sutherland_Temp

Given: Sutherland's Reference Temperature (K)

· double Sutherland Const

Given: Sutherland's Constant (K)

· double Sutherland_Viscosity

Given: Sutherland's Reference Viscosity (g/cm/s)

· double specific_heat

Given: Specific heat of the gas (J/g/K)

• double molecular_diffusion

Calculated: molecular diffusivities (cm²/s)

double dynamic_viscosity

Calculated: dynamic viscosities (g/cm/s)

· double density

Calculated: gas densities (g/cm^{\(\)}3) {use RE3}.

double Schmidt

Calculated: Value of the Schmidt number (-)

5.39.1 Detailed Description

Data structure holding all the parameters for each pure gas spieces.

C-style object that holds the constants and parameters associated with each pure gas species in the overall mixture. This information is used in conjunction with the kinetic theory of gases to produce approximations to many different gas properties needed in simulating gas dynamics, mobility of a gas through porous media, as well as some kinetic adsorption parameters such as diffusivities.

5.39.2 Member Data Documentation

5.39.2.1 double PURE_GAS::molecular_weight

Given: molecular weights (g/mol)

5.39.2.2 double PURE_GAS::Sutherland_Temp

Given: Sutherland's Reference Temperature (K)

5.39.2.3 double PURE_GAS::Sutherland_Const

Given: Sutherland's Constant (K)

5.39.2.4 double PURE_GAS::Sutherland_Viscosity

Given: Sutherland's Reference Viscosity (g/cm/s)

5.39.2.5 double PURE_GAS::specific_heat

Given: Specific heat of the gas (J/g/K)

5.39.2.6 double PURE_GAS::molecular_diffusion

Calculated: molecular diffusivities (cm²/s)

5.39.2.7 double PURE_GAS::dynamic_viscosity

Calculated: dynamic viscosities (g/cm/s)

5.39.2.8 double PURE_GAS::density

Calculated: gas densities (g/cm³) {use RE3}.

5.39.2.9 double PURE_GAS::Schmidt

Calculated: Value of the Schmidt number (-)

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

· egret.h

5.40 QR_DATA Struct Reference

Data structure for the implementation of a QR solver given some invertable linear operator.

```
#include <lark.h>
```

Public Attributes

Matrix< double > ek

Unit vector used to extract columns from the linear operator.

Matrix< double > Ro

Upper triangular matrix formed from factoring the linear operator.

Matrix< double > x

Solution to the linear system.

5.40.1 Detailed Description

Data structure for the implementation of a QR solver given some invertable linear operator.

C-style object to be used in conjuction with a QR solver for invertable linear operators. This method will extract columns from the linear operator and use Householder Reflections to factor the operator into an upper triangular matrix and a unitary reflection matrix. It is generally less efficient to use this method for sparse systems, but is more stable and occassionally more efficient for dense systems.

5.40.2 Member Data Documentation

5.40.2.1 Matrix<double> QR_DATA::ek

Unit vector used to extract columns from the linear operator.

5.40.2.2 Matrix<double> QR_DATA::Ro

Upper triangular matrix formed from factoring the linear operator.

5.40.2.3 Matrix<double> QR_DATA::x

Solution to the linear system.

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

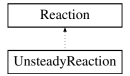
· lark.h

5.41 Reaction Class Reference

Reaction Object.

#include <shark.h>

Inheritance diagram for Reaction:



Public Member Functions

• Reaction ()

Default constructor.

∼Reaction ()

Default destructor.

• void Initialize_Object (MasterSpeciesList &List)

Function to initialize the Reaction object from the MasterSpeciesList.

• void Display_Info ()

Display the reaction information.

• void Set Stoichiometric (int i, double v)

Set the ith stoichiometric value.

• void Set_Equilibrium (double logK)

Set the equilibrium constant in log(K) units.

void Set Enthalpy (double H)

Set the enthalpy of the reaction (J/mol)

void Set_Entropy (double S)

Set the entropy of the reaction (J/K/mol)

void Set_EnthalpyANDEntropy (double H, double S)

Set both the enthalpy and entropy (J/mol) & (J/K/mol)

void Set Energy (double G)

Set the Gibb's free energy of reaction (J/mol)

void checkSpeciesEnergies ()

Function to check MasterList Reference for species energy info.

- void calculateEnergies ()
- void calculateEquilibrium (double T)

Function to calculate the equilibrium constant based on temperature in K.

bool haveEquilibrium ()

Function to return true if equilibrium constant is given or can be calculated.

• double Get Stoichiometric (int i)

Fetch the ith stoichiometric value.

• double Get_Equilibrium ()

Fetch the equilibrium constant (logK)

double Get_Enthalpy ()

Fetch the enthalpy of the reaction (J/mol)

double Get_Entropy ()

Fetch the entropy of the reaction (J/K/mol)

• double Get_Energy ()

Fetch the energy of the reaction (J/mol)

double Eval_Residual (const Matrix< double > &x, const Matrix< double > &gama)

Protected Attributes

MasterSpeciesList * List

Pointer to a master species object.

• std::vector< double > Stoichiometric

Vector of stoichiometric constants corresponding to species list.

double Equilibrium

Equilibrium constant for the reaction (logK)

· double enthalpy

Reaction enthalpy (J/mol)

· double entropy

Reaction entropy (J/K/mol)

· double energy

Gibb's Free energy of reaction (J/mol)

bool CanCalcHS

True if all molecular info is avaiable to calculate dH and dS.

bool CanCalcG

True if all molecular info is available to calculate dG.

bool HaveHS

True if dH and dS is given, or can be calculated.

bool HaveG

True if dG is given, or can be calculated.

bool HaveEquil

True as long as Equilibrium is given, or can be calculated.

5.41.1 Detailed Description

Reaction Object.

C++ style object that holds data and functions associated with standard chemical reactions...

```
i.e., aA + bB \le cC + dD
```

These reactions are assumed steady state and are characterized by stoichiometry coefficients and equilibrium/stability constants. Types of reactions that these are valid for would be acid/base reactions, metal-ligand complexation reactions, oxidation-reduction reactions, Henry's Law phase changes, and more. Reactions that this may not be suitable for include mechanisms, adsorption, and precipitation. Those types of reactions would be better handled by more specific objects that inherit from this object.

If all species in the reaction are registered and known species in mola.h AND have known formation energies, then the equilibrium constants for that particular reaction will be calculated based on the species involved in the reaction. However, if using some custom molecule objects, then the reaction equilibrium may not be able to be automatically formed by the routine. In this case, you would need to also supply the equilibrium constant for the particular reaction.

```
5.41.2 Constructor & Destructor Documentation
```

```
5.41.2.1 Reaction::Reaction ( )
```

Default constructor.

```
5.41.2.2 Reaction::~Reaction ( )
```

Default destructor.

5.41.3 Member Function Documentation

5.41.3.1 void Reaction::Initialize_Object (MasterSpeciesList & List)

Function to initialize the Reaction object from the MasterSpeciesList.

```
5.41.3.2 void Reaction::Display_Info()
```

Display the reaction information.

```
5.41.3.3 void Reaction::Set_Stoichiometric (int i, double v)
```

Set the ith stoichiometric value.

This function will set the stoichiometric constant of the ith species in the master list to the given value of v. All values of v are set to zero unless overriden by this function.

Parameters

i	index of the species in the MasterSpeciesList
V	value of the stoichiometric constant for that species in the reaction

```
5.41.3.4 void Reaction::Set_Equilibrium ( double logK )
Set the equilibrium constant in log(K) units.
5.41.3.5 void Reaction::Set_Enthalpy ( double H )
Set the enthalpy of the reaction (J/mol)
5.41.3.6 void Reaction::Set_Entropy ( double S )
Set the entropy of the reaction (J/K/mol)
5.41.3.7 void Reaction::Set_EnthalpyANDEntropy ( double H, double S )
Set both the enthalpy and entropy (J/mol) & (J/K/mol)
5.41.3.8 void Reaction::Set_Energy ( double G )
Set the Gibb's free energy of reaction (J/mol)
5.41.3.9 void Reaction::checkSpeciesEnergies ( )
Function to check MasterList Reference for species energy info.
This function will go through the stoichiometry of this reaction and check the molecules in the MasterSpeciesList
that correspond to the species present in this reaction for the existance of their formation energies. Based on the
states of those energies, it will note internally whether or not it can determine the equilibrium constants based soley
on individual species information. If it cannot, then the user must provide either the reaction energies to form the
equilibrium constant or the equilibrium constant itself. Function to calculate and set the energy of the reaction
5.41.3.10 void Reaction::calculateEnergies ( )
If the energies of the reaction can be determined from the individual species in the reaction, then this function uses
that information. Otherwise, it sets the energies equal to the constants given to the object by the user.
5.41.3.11 void Reaction::calculateEquilibrium ( double T )
Function to calculate the equilibrium constant based on temperature in K.
5.41.3.12 bool Reaction::haveEquilibrium ( )
Function to return true if equilibrium constant is given or can be calculated.
5.41.3.13 double Reaction::Get_Stoichiometric ( int i )
Fetch the ith stoichiometric value.
5.41.3.14 double Reaction::Get_Equilibrium ( )
Fetch the equilibrium constant (logK)
```

```
5.41.3.15 double Reaction::Get_Enthalpy ( )

Fetch the enthalpy of the reaction (J/mol)

5.41.3.16 double Reaction::Get_Entropy ( )

Fetch the entropy of the reaction (J/K/mol)

5.41.3.17 double Reaction::Get_Energy ( )

Fetch the energy of the reaction (J/mol)
```

Evaluate a residual for the reaction given variable x=log(C) and activity coefficients gama

```
5.41.3.18 double Reaction::Eval_Residual ( const Matrix< double > & x, const Matrix< double > & gama)
```

This function will calculate the reaction residual from this object's stoichiometry, equilibrium constant, log(C) concentrations, and activity coefficients.

Parameters

	Х	matrix of the log(C) concentration values at the current non-linear step
ſ	gama	matrix of activity coefficients for each species at the current non-linear step

5.41.4 Member Data Documentation

```
5.41.4.1 MasterSpeciesList* Reaction::List [protected]
```

Pointer to a master species object.

```
5.41.4.2 std::vector<double> Reaction::Stoichiometric [protected]
```

Vector of stoichiometric constants corresponding to species list.

```
5.41.4.3 double Reaction::Equilibrium [protected]
```

Equilibrium constant for the reaction (logK)

```
5.41.4.4 double Reaction::enthalpy [protected]
```

Reaction enthalpy (J/mol)

5.41.4.5 double Reaction::entropy [protected]

Reaction entropy (J/K/mol)

5.41.4.6 double Reaction::energy [protected]

Gibb's Free energy of reaction (J/mol)

```
5.41.4.7 bool Reaction::CanCalcHS [protected]
```

True if all molecular info is avaiable to calculate dH and dS.

```
5.41.4.8 bool Reaction::CanCalcG [protected]
```

True if all molecular info is available to calculate dG.

```
5.41.4.9 bool Reaction::HaveHS [protected]
```

True if dH and dS is given, or can be calculated.

```
5.41.4.10 bool Reaction::HaveG [protected]
```

True if dG is given, or can be calculated.

```
5.41.4.11 bool Reaction::HaveEquil [protected]
```

True as long as Equilibrium is given, or can be calculated.

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

• shark.h

5.42 SCOPSOWL_DATA Struct Reference

Primary data structure for SCOPSOWL simulations.

```
#include <scopsowl.h>
```

Public Attributes

· unsigned long int total steps

Running total of all calculation steps.

int coord_macro

Coordinate system for large pellet.

int coord_micro

Coordinate system for small crystal (if any)

• int level = 2

Level of coupling between the different scales (default = 2)

double sim_time

Stopping time for the simulation (hrs)

· double t old

Old time of the simulations (hrs)

double t

Current time of the simulations (hrs)

• double t counter = 0.0

Counter for the time output.

double t_print

Print output at every t_print time (hrs) • bool Print2File = true True = results to .txt; False = no printing. • bool Print2Console = true True = results to console; False = no printing. bool SurfDiff = true True = includes SKUA simulation if Heterogeneous; False = only uses MAGPIE. • bool Heterogeneous = true True = pellet is made of binder and crystals, False = all one phase. · double gas_velocity Superficial Gas Velocity arount pellet (cm/s) • double total_pressure Gas phase total pressure (kPa) • double gas_temperature Gas phase temperature (K) double pellet radius Nominal radius of the pellet - macroscale domain (cm) · double crystal radius Nominal radius of the crystal - microscale domain (um) · double char macro Characteristic size for macro scale (cm or cm²) - only if pellet is not spherical. · double char_micro Characteristic size for micro scale (um or um $^{\wedge}$ 2) - only if crystal is not spherical. double binder_fraction Volume of binder per total volume of pellet (-) · double binder_porosity Volume of pores per volume of binder (-) · double binder poresize Nominal radius of the binder pores (cm) · double pellet density Mass of the pellet per volume of pellet (kg/L) bool DirichletBC = false True = Dirichlet BC; False = Neumann BC. • bool NonLinear = true True = Non-linear solver; False = Linear solver. std::vector< double > y Outside mole fractions of each component (-) std::vector< double > tempy Temporary place holder for gas mole fractions in other locations (-) FILE * OutputFile Output file pointer to the output file for postprocesses. double(* eval ads)(int i, int I, const void *user data) Function pointer for evaluating adsorption (mol/kg) double(* eval_retard)(int i, int I, const void *user_data) Function pointer for evaluating retardation (-) double(* eval_diff)(int i, int I, const void *user_data) Function pointer for evaluating pore diffusion (cm²/hr) double(* eval_surfDiff)(int i, int I, const void *user_data) Function pointer for evaluating surface diffusion (um²/hr) double(* eval_kf)(int i, const void *user_data)

Function pointer for evaluating film mass transfer (cm/hr)

const void * user_data

Data structure for users info to calculate parameters.

MIXED GAS * gas dat

Pointer to the MIXED_GAS data structure (may or may not be used)

MAGPIE_DATA magpie_dat

Data structure for a magpie problem (to be used if not using skua)

std::vector< FINCH DATA > finch dat

Data structure for pore adsorption kinetics for all species (u in mol/L)

std::vector < SCOPSOWL_PARAM_DATA > param_dat

Data structure for parameter info for all species.

std::vector < SKUA DATA > skua dat

Data structure holding a skua object for all nodes (each skua has an object for each species)

5.42.1 Detailed Description

Primary data structure for SCOPSOWL simulations.

C-style object holding necessary information to run a SCOPSOWL simulation. SCOPSOWL is a multi-scale problem involving PDE solution for the macro-scale adsorbent pellet and the micro-scale adsorbent crystals. As such, each SCOPSOWL simulation involves multiple SKUA simulations at the nodes in the macro-scale domain. Alternatively, if the user wishes to specify that the adsorbent is homogeneous, then you can run SCOPSOWL as a single-scale problem. Additionally, you can simplfy the model by assuming that the micro-scale diffusion is very fast, and therefore replace each SKUA simulation with a simpler MAGPIE evaluation. Details on running SCOPSOWL with the various options will be discussed in the SCOPSOWL_SCENARIOS function.

5.42.2 Member Data Documentation

5.42.2.1 unsigned long int SCOPSOWL_DATA::total_steps

Running total of all calculation steps.

5.42.2.2 int SCOPSOWL_DATA::coord_macro

Coordinate system for large pellet.

5.42.2.3 int SCOPSOWL_DATA::coord_micro

Coordinate system for small crystal (if any)

5.42.2.4 int SCOPSOWL_DATA::level = 2

Level of coupling between the different scales (default = 2)

5.42.2.5 double SCOPSOWL_DATA::sim_time

Stopping time for the simulation (hrs)

5.42.2.6 double SCOPSOWL_DATA::t_old

Old time of the simulations (hrs)

5.42.2.7 double SCOPSOWL_DATA::t

Current time of the simulations (hrs)

5.42.2.8 double SCOPSOWL_DATA::t_counter = 0.0

Counter for the time output.

5.42.2.9 double SCOPSOWL_DATA::t_print

Print output at every t_print time (hrs)

5.42.2.10 bool SCOPSOWL_DATA::Print2File = true

True = results to .txt; False = no printing.

5.42.2.11 bool SCOPSOWL_DATA::Print2Console = true

True = results to console; False = no printing.

5.42.2.12 bool SCOPSOWL_DATA::SurfDiff = true

True = includes SKUA simulation if Heterogeneous; False = only uses MAGPIE.

5.42.2.13 bool SCOPSOWL_DATA::Heterogeneous = true

True = pellet is made of binder and crystals, False = all one phase.

5.42.2.14 double SCOPSOWL_DATA::gas_velocity

Superficial Gas Velocity arount pellet (cm/s)

5.42.2.15 double SCOPSOWL_DATA::total_pressure

Gas phase total pressure (kPa)

5.42.2.16 double SCOPSOWL_DATA::gas_temperature

Gas phase temperature (K)

5.42.2.17 double SCOPSOWL_DATA::pellet_radius

Nominal radius of the pellet - macroscale domain (cm)

5.42.2.18 double SCOPSOWL_DATA::crystal_radius

Nominal radius of the crystal - microscale domain (um)

5.42.2.19 double SCOPSOWL_DATA::char_macro

Characteristic size for macro scale (cm or cm²) - only if pellet is not spherical.

5.42.2.20 double SCOPSOWL_DATA::char_micro

Characteristic size for micro scale (um or um²) - only if crystal is not spherical.

5.42.2.21 double SCOPSOWL_DATA::binder_fraction

Volume of binder per total volume of pellet (-)

5.42.2.22 double SCOPSOWL_DATA::binder_porosity

Volume of pores per volume of binder (-)

5.42.2.23 double SCOPSOWL_DATA::binder_poresize

Nominal radius of the binder pores (cm)

5.42.2.24 double SCOPSOWL_DATA::pellet_density

Mass of the pellet per volume of pellet (kg/L)

5.42.2.25 bool SCOPSOWL_DATA::DirichletBC = false

True = Dirichlet BC; False = Neumann BC.

5.42.2.26 bool SCOPSOWL_DATA::NonLinear = true

True = Non-linear solver; False = Linear solver.

5.42.2.27 std::vector<double> SCOPSOWL_DATA::y

Outside mole fractions of each component (-)

5.42.2.28 std::vector<double> SCOPSOWL_DATA::tempy

Temporary place holder for gas mole fractions in other locations (-)

5.42.2.29 FILE* SCOPSOWL_DATA::OutputFile

Output file pointer to the output file for postprocesses.

 $5.42.2.30 \quad double(*\ SCOPSOWL_DATA::eval_ads)\ (int\ i,\ int\ I,\ const\ void\ *user_data)$

Function pointer for evaluating adsorption (mol/kg)

5.42.2.31 double(* SCOPSOWL_DATA::eval_retard) (int i, int I, const void *user_data)

Function pointer for evaluating retardation (-)

5.42.2.32 double(* SCOPSOWL_DATA::eval_diff) (int i, int I, const void *user_data)

Function pointer for evaluating pore diffusion (cm²/hr)

5.42.2.33 double(* SCOPSOWL_DATA::eval_surfDiff) (int i, int I, const void *user_data)

Function pointer for evaluating surface diffusion (um²/hr)

5.42.2.34 double(* SCOPSOWL_DATA::eval_kf) (int i, const void *user_data)

Function pointer for evaluating film mass transfer (cm/hr)

5.42.2.35 const void* SCOPSOWL_DATA::user_data

Data structure for users info to calculate parameters.

5.42.2.36 MIXED_GAS* SCOPSOWL_DATA::gas_dat

Pointer to the MIXED_GAS data structure (may or may not be used)

5.42.2.37 MAGPIE_DATA SCOPSOWL_DATA::magpie_dat

Data structure for a magpie problem (to be used if not using skua)

 $5.42.2.38 \quad std:: vector < \textbf{FINCH_DATA} > \textbf{SCOPSOWL_DATA}:: finch_dat$

Data structure for pore adsorption kinetics for all species (u in mol/L)

5.42.2.39 std::vector<SCOPSOWL_PARAM_DATA> SCOPSOWL_DATA::param_dat

Data structure for parameter info for all species.

5.42.2.40 std::vector<SKUA DATA> SCOPSOWL_DATA::skua_dat

Data structure holding a skua object for all nodes (each skua has an object for each species)

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

scopsowl.h

5.43 SCOPSOWL_OPT_DATA Struct Reference

Data structure for the SCOPSOWL optmization routine.

#include <scopsowl_opt.h>

Public Attributes

int num_curves

Number of adsorption curves to analyze.

· int evaluation

Number of times the eval function has been called for a single curve.

· unsigned long int total_eval

Total number of evaluations needed for completion.

· int current_points

Number of points in the current curve.

• int num_params = 1

Number of adjustable parameters for the current curve (currently only supports 1)

· int diffusion_type

Flag to identify type of diffusion function to use.

· int adsorb_index

Component index for adsorbable species.

int max_guess_iter = 20

Maximum allowed guess iterations (default = 20)

· bool Optimize

True = run optimization, False = run a comparison.

bool Rough

True = use only a rough estimate, False = run full optimization.

· double current temp

Temperature for current curve.

· double current_press

Partial pressure for current curve.

double current_equil

Equilibrium data point for the current curve.

double simulation_equil

Equilibrium simulation point for the current curve.

· double max bias

Positive maximum bias plausible for fitting.

double min_bias

Negative minimum bias plausible for fitting.

• double e norm

Euclidean norm of current fit.

• double f_bias

Function bias of current fit.

double e_norm_old

Euclidean norm of the previous fit.

· double f_bias_old

Function bias of the previous fit.

• double param_guess

Parameter guess for the surface/crystal diffusivity.

· double param guess old

Parameter guess for the previous curve.

double rel_tol_norm = 0.01

Tolerance for convergence of the guess norm.

• double abs tol bias = 1.0

Tolerance for convergence of the guess bias.

std::vector< double > y_base

Gas phase mole fractions in absense of adsorbing species.

std::vector< double > q_data

Amount adsorbed at a particular point in current curve.

std::vector< double > q_sim

Amount adsorbed based on the simulation.

std::vector< double > t

Time points in the current curve.

FILE * ParamFile

Output file for parameter results.

FILE * CompareFile

Output file for comparison of results.

SCOPSOWL_DATA owl_dat

Data structure for the SCOPSOWL simulation.

5.43.1 Detailed Description

Data structure for the SCOPSOWL optmization routine.

C-style object holding information about the optimization routine as well as the standard SCOPSOWI_DATA structure for SCOPSOWL simulations.

5.43.2 Member Data Documentation

5.43.2.1 int SCOPSOWL_OPT_DATA::num_curves

Number of adsorption curves to analyze.

5.43.2.2 int SCOPSOWL_OPT_DATA::evaluation

Number of times the eval function has been called for a single curve.

5.43.2.3 unsigned long int SCOPSOWL_OPT_DATA::total_eval

Total number of evaluations needed for completion.

5.43.2.4 int SCOPSOWL_OPT_DATA::current_points

Number of points in the current curve.

5.43.2.5 int SCOPSOWL_OPT_DATA::num_params = 1

Number of adjustable parameters for the current curve (currently only supports 1)

5.43.2.6 int SCOPSOWL_OPT_DATA::diffusion_type

Flag to identify type of diffusion function to use.

5.43.2.7 int SCOPSOWL_OPT_DATA::adsorb_index

Component index for adsorbable species.

5.43.2.8 int SCOPSOWL_OPT_DATA::max_guess_iter = 20

Maximum allowed guess iterations (default = 20)

5.43.2.9 bool SCOPSOWL_OPT_DATA::Optimize

True = run optimization, False = run a comparison.

5.43.2.10 bool SCOPSOWL_OPT_DATA::Rough

True = use only a rough estimate, False = run full optimization.

5.43.2.11 double SCOPSOWL_OPT_DATA::current_temp

Temperature for current curve.

5.43.2.12 double SCOPSOWL_OPT_DATA::current_press

Partial pressure for current curve.

5.43.2.13 double SCOPSOWL_OPT_DATA::current_equil

Equilibrium data point for the current curve.

5.43.2.14 double SCOPSOWL_OPT_DATA::simulation_equil

Equilibrium simulation point for the current curve.

5.43.2.15 double SCOPSOWL_OPT_DATA::max_bias

Positive maximum bias plausible for fitting.

5.43.2.16 double SCOPSOWL_OPT_DATA::min_bias

Negative minimum bias plausible for fitting.

5.43.2.17 double SCOPSOWL_OPT_DATA::e_norm

Euclidean norm of current fit.

5.43.2.18 double SCOPSOWL_OPT_DATA::f_bias

Function bias of current fit.

5.43.2.19 double SCOPSOWL_OPT_DATA::e_norm_old

Euclidean norm of the previous fit.

5.43.2.20 double SCOPSOWL_OPT_DATA::f_bias_old

Function bias of the previous fit.

5.43.2.21 double SCOPSOWL_OPT_DATA::param_guess

Parameter guess for the surface/crystal diffusivity.

5.43.2.22 double SCOPSOWL_OPT_DATA::param_guess_old

Parameter guess for the previous curve.

5.43.2.23 double SCOPSOWL_OPT_DATA::rel_tol_norm = 0.01

Tolerance for convergence of the guess norm.

5.43.2.24 double SCOPSOWL_OPT_DATA::abs_tol_bias = 1.0

Tolerance for convergence of the guess bias.

5.43.2.25 std::vector<double> SCOPSOWL_OPT_DATA::y_base

Gas phase mole fractions in absense of adsorbing species.

5.43.2.26 std::vector<double> SCOPSOWL_OPT_DATA::q_data

Amount adsorbed at a particular point in current curve.

 $5.43.2.27 \quad std::vector < double > SCOPSOWL_OPT_DATA::q_sim$

Amount adsorbed based on the simulation.

 $5.43.2.28 \quad std::vector < double > SCOPSOWL_OPT_DATA::t$

Time points in the current curve.

5.43.2.29 FILE* SCOPSOWL_OPT_DATA::ParamFile

Output file for parameter results.

5.43.2.30 FILE* SCOPSOWL_OPT_DATA::CompareFile

Output file for comparison of results.

5.43.2.31 SCOPSOWL_DATA SCOPSOWL_OPT_DATA::owl_dat

Data structure for the SCOPSOWL simulation.

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

· scopsowl opt.h

5.44 SCOPSOWL_PARAM_DATA Struct Reference

Data structure for the species' parameters in SCOPSOWL.

```
#include <scopsowl.h>
```

Public Attributes

Matrix< double > qAvg

Average adsorbed amount for a species at each node (mol/kg)

Matrix< double > qAvg_old

Old Average adsorbed amount for a species at each node (mol/kg)

Matrix< double > Qst

Heat of adsorption for all nodes (J/mol)

Matrix< double > Qst_old

Old Heat of adsorption for all nodes (J/mol)

Matrix< double > dq dc

Storage vector for current adsorption slope/strength (dq/dc) (L/kg)

double xIC

Initial conditions for adsorbed molefractions.

double qIntegralAvg

Integral average of adsorption over the entire pellet (mol/kg)

double qIntegralAvg_old

Old Integral average of adsorption over the entire pellet (mol/kg)

double QstAvg

Integral average heat of adsorption (J/mol)

double QstAvg_old

Old integral average heat of adsorption (J/mol)

• double qo

Boundary value of adsorption if using Dirichlet BCs (mol/kg)

· double Qsto

Boundary value of adsorption heat if using Dirichlet BCs (J/mol)

double dq_dco

Boundary value of adsorption slope for Dirichelt BCs (L/kg)

• double pore diffusion

Value for constant pore diffusion (cm²/hr)

double film_transfer

Value for constant film mass transfer (cm/hr)

· double activation energy

Activation energy for surface diffusion (J/mol)

• double ref_diffusion

Reference state surface diffusivity (um²/hr)

• double ref_temperature

Reference temperature for empirical adjustments (K)

· double affinity

Affinity parameter used in empirical adjustments (-)

- double ref_pressure
- · bool Adsorbable

True = species can adsorb; False = species cannot adsorb.

std::string speciesName

String to hold the name of each species.

5.44.1 Detailed Description

Data structure for the species' parameters in SCOPSOWL.

C-style object that holds information on all species for a particular SCOPSOWL simulation. Initial conditions, kinetic parameters, and interim matrix objects are stored here for use in various SCOSPSOWL functions.

5.44.2 Member Data Documentation

5.44.2.1 Matrix<double> SCOPSOWL_PARAM_DATA::qAvg

Average adsorbed amount for a species at each node (mol/kg)

 $5.44.2.2 \quad Matrix{<} double{>} SCOPSOWL_PARAM_DATA::qAvg_old$

Old Average adsorbed amount for a species at each node (mol/kg)

5.44.2.3 Matrix<double> SCOPSOWL_PARAM_DATA::Qst

Heat of adsorption for all nodes (J/mol)

5.44.2.4 Matrix<double> SCOPSOWL_PARAM_DATA::Qst_old

Old Heat of adsorption for all nodes (J/mol)

 $5.44.2.5 \quad \textbf{Matrix} {<} \textbf{double} {>} \ \textbf{SCOPSOWL_PARAM_DATA} {::} \textbf{dq_dc}$

Storage vector for current adsorption slope/strength (dq/dc) (L/kg)

5.44.2.6 double SCOPSOWL_PARAM_DATA::xIC

Initial conditions for adsorbed molefractions.

 $5.44.2.7 \quad double \ SCOPSOWL_PARAM_DATA:: qIntegral Avg$

Integral average of adsorption over the entire pellet (mol/kg)

5.44.2.8 double SCOPSOWL_PARAM_DATA::qintegralAvg_old

Old Integral average of adsorption over the entire pellet (mol/kg)

5.44.2.9 double SCOPSOWL_PARAM_DATA::QstAvg

Integral average heat of adsorption (J/mol)

5.44.2.10 double SCOPSOWL_PARAM_DATA::QstAvg_old

Old integral average heat of adsorption (J/mol)

5.44.2.11 double SCOPSOWL_PARAM_DATA::qo

Boundary value of adsorption if using Dirichlet BCs (mol/kg)

5.44.2.12 double SCOPSOWL_PARAM_DATA::Qsto

Boundary value of adsorption heat if using Dirichlet BCs (J/mol)

5.44.2.13 double SCOPSOWL_PARAM_DATA::dq_dco

Boundary value of adsorption slope for Dirichelt BCs (L/kg)

5.44.2.14 double SCOPSOWL_PARAM_DATA::pore_diffusion

Value for constant pore diffusion (cm²/hr)

5.44.2.15 double SCOPSOWL_PARAM_DATA::film_transfer

Value for constant film mass transfer (cm/hr)

5.44.2.16 double SCOPSOWL_PARAM_DATA::activation_energy

Activation energy for surface diffusion (J/mol)

5.44.2.17 double SCOPSOWL_PARAM_DATA::ref_diffusion

Reference state surface diffusivity (um²/hr)

5.44.2.18 double SCOPSOWL_PARAM_DATA::ref_temperature

Reference temperature for empirical adjustments (K)

5.44.2.19 double SCOPSOWL_PARAM_DATA::affinity

Affinity parameter used in empirical adjustments (-)

5.44.2.20 double SCOPSOWL_PARAM_DATA::ref_pressure

5.44.2.21 bool SCOPSOWL_PARAM_DATA::Adsorbable

True = species can adsorb; False = species cannot adsorb.

5.44.2.22 std::string SCOPSOWL_PARAM_DATA::speciesName

String to hold the name of each species.

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

· scopsowl.h

5.45 SHARK_DATA Struct Reference

Data structure for SHARK simulations.

```
#include <shark.h>
```

Public Attributes

MasterSpeciesList MasterList

Master List of species object.

• std::vector< Reaction > ReactionList

Equilibrium reaction objects.

std::vector< MassBalance > MassBalanceList

Mass balance objects.

• std::vector< UnsteadyReaction > UnsteadyList

Unsteady Reaction objects.

std::vector< AdsorptionReaction > AdsorptionList

Equilibrium Adsorption Reaction Objects.

std::vector< UnsteadyAdsorption > UnsteadyAdsList

Unsteady Adsorption Reaction Objects.

 $\bullet \ \, std:: vector < Multiligand Adsorption > Multi Ads List$

Multiligand Adsorptioin Objects.

• std::vector< ChemisorptionReaction > ChemisorptionList

Chemisorption Reaction objects.

• std::vector< MultiligandChemisorption > MultiChemList

Multiligand Chemisorption Reaction Objects.

std::vector< double(*)(const Matrix< double > &x, SHARK_DATA *shark_dat, const void *data) > OtherList

Array of Other Residual functions to be defined by user.

· int numvar

Total number of functions and species.

• int num_ssr

Number of steady-state reactions.

· int num mbe

Number of mass balance equations.

• int num_usr = 0

Number of unsteady-state reactions.

• int num_ssao = 0

Number of steady-state adsorption objects.

• int num usao = 0

Number of unsteady adsorption objects.

• int num multi ssao = 0

Number of multiligand steady-state adsorption objects.

• int num sschem = 0

Number of steady-state chemisorption objects.

int num_multi_sschem = 0

Number of multiligand steady-state chemisorption objects.

std::vector< int > num_ssar

List of the numbers of reactions in each adsorption object.

std::vector< int > num_usar

List of the numbers of reactions in each unsteady adsorption object.

std::vector< int > num_sschem_rxns

List of the numbers of reactions in each steady-state chemisorption object.

std::vector< std::vector< int > > num_multi_ssar

List of all multiligand objects -> List of ligands and rxns of that ligand.

• std::vector< std::vector< int > > num multichem rxns

List of all multiligand chemisorption objects -> List of num rxns for that ligand.

std::vector< std::string > ss_ads_names

List of the steady-state adsorbent object names.

std::vector< std::string > us_ads_names

List of the unsteady adsorption object names.

 $\bullet \ \, \mathsf{std} :: \mathsf{vector} < \mathsf{std} :: \mathsf{string} > \mathsf{ss_chem_names} \\$

List of the steady-state chemisorption object names.

std::vector< std::vector< std::string >> ssmulti_names

List of the names of the ligands in each multiligand object.

• std::vector< std::vector< std::string >> ssmultichem names

List of the names of the ligands in each multiligand chemisorption object.

• int num other = 0

Number of other functions to be used (default is always 0)

int act_fun = IDEAL

Flag denoting the activity function to use (default is IDEAL)

int reactor_type = BATCH

Flag denoting the type of reactor considered for the system (default is BATCH)

• int totalsteps = 0

Total number of iterations.

• int totalcalls = 0

Total number of residual function calls.

• int timesteps = 0

Number of time steps taken to complete simulation.

• int pH_index = -1

Contains the index of the pH variable (set internally)

• int pOH index = -1

Contains the index of the pOH variable (set internally)

• double simulationtime = 0.0

Time to simulate unsteady reactions for (default = 0.0 hrs)

double dt = 0.1

Time step size (hrs)

```
    double dt_min = sqrt(DBL_EPSILON)

      Minimum allowable step size.

    double dt max = 744.0

      Maximum allowable step size (\sim1 month in time)
• double t out = 0.0
      Time increment by which file output is made (default = print all time steps)
• double t count = 0.0
      Running count of time increments.
• double time = 0.0
      Current value of time (starts from t = 0.0 \text{ hrs})
• double time old = 0.0
      Previous value of time (start from t = 0.0 \text{ hrs})

    double pH = 7.0

      Value of pH if needed (default = 7)
• double pH step = 0.5
      Value by which to increment pH when doing a speciation curve (default = 0.5)
double start_temp = 277.15
      Value of the starting temperature used for Temperature Curves (default = 277.15 K)
• double end_temp = 323.15
      Value of the ending temperature used for Temperature Curves (default = 323.15 K)
• double temp_step = 10.0
      Size of the step changes to use for Temperature Curves (default = 10.0 K);.

 double volume = 1.0

      Volume of the domain in liters (default = 1 L)
double flow_rate = 1.0
      Flow rate in the reactor in L/hr (default = 1 L/hr)
• double xsec area = 1.0
      Cross sectional area of the reactor in m^2 (default = 1 m^22)
• double Norm = 0.0
      Current value of euclidean norm in solution.
• double dielectric const = 78.325
      Dielectric constant used in many activity models (default: water = 78.325 (1/K))
• double relative_permittivity = 80.1
      Relative permittivity of the medium (default: water = 80.1 (-))
• double temperature = 298.15
      Solution temperature (default = 25 oC or 298.15 K)
• double ionic strength = 0.0
      Solution ionic strength in Molar (calculated internally)

    bool steadystate = true

      True = solve steady problem; False = solve transient problem.
• bool ZeroInitialSolids = false
      True = no solids or adsorption initially in the reactor.
• bool TimeAdaptivity = false
      True = solve using variable time step.
• bool const pH = false
      True = set pH to a constant; False = solve for pH.

    bool SpeciationCurve = false

      True = runs a series of constant pH steady-state problems to produce curves.

    bool TemperatureCurve = false

      True = runs a series of constant temperature steady-state problmes to produce curves.
```

bool Console_Output = true

True = display output to console.

• bool File Output = false

True = write output to a file.

• bool Contains_pH = false

True = system contains pH as a variable (set internally)

bool Contains pOH = false

True = system contains pOH as a variable (set internally)

• bool Converged = false

True = system converged within tolerance.

bool LocalMin = true

True = allow the system to settle for a local minimum if tolerance not reached.

Matrix< double > X old

Solution vector for old time step - log(C)

Matrix< double > X_new

Solution vector for current time step - log(C)

Matrix< double > Conc_old

Concentration vector for old time step - 10° x.

Matrix< double > Conc new

Concentration vector for current time step - 10° x.

Matrix< double > activity new

Activity matrix for current time step.

Matrix< double > activity_old

Activity matrix from prior time step.

int(* EvalActivity)(const Matrix< double > &x, Matrix< double > &F, const void *data)

Function pointer to evaluate activity coefficients.

int(* Residual)(const Matrix < double > &x, Matrix < double > &F, const void *data)

Function pointer to evaluate all residuals in the system.

• int(* lin_precon)(const Matrix< double > &r, Matrix< double > &p, const void *data)

Function pointer to form a linear preconditioning operation for the Jacobian.

PJFNK_DATA Newton_data

Data structure for the Newton-Krylov solver (see lark.h)

const void * activity data

User defined data structure for an activity model.

const void * residual_data

User defined data structure for the residual function.

const void * precon_data

User defined data structure for preconditioning.

const void * other_data

User define data structure used for user defined residuals.

FILE * OutputFile

Output File pointer.

yaml_cpp_class yaml_object

yaml object to read and access digitized yaml documents (see yaml_wrapper.h)

5.45.1 Detailed Description

Data structure for SHARK simulations.

C-style object holding data and function pointers associated with solving aqueous speciation and reaction kinetics. This object couples all other objects available in shark.h in order to provide residual calculations for each individual function that makes up the overall system model. Those residuals are brought together inside the residual function and fed into the lark.h PJFNK solver routine. That solver then attempts to find a solution to all non-linear variables simultaneously. Any function or data pointers in this structure can be overriden to change how you interface with and solve the problem. Users may also provide a set of custom residual functions through the "OtherList" vector object. Those residual function must all have the same format.

5.45.2 Member Data Documentation

5.45.2.1 MasterSpeciesList SHARK_DATA::MasterList

Master List of species object.

5.45.2.2 std::vector < Reaction > SHARK_DATA::ReactionList

Equilibrium reaction objects.

5.45.2.3 std::vector < MassBalance > SHARK_DATA::MassBalanceList

Mass balance objects.

5.45.2.4 std::vector < UnsteadyReaction > SHARK_DATA::UnsteadyList

Unsteady Reaction objects.

5.45.2.5 std::vector < AdsorptionReaction > SHARK_DATA::AdsorptionList

Equilibrium Adsorption Reaction Objects.

5.45.2.6 std::vector<UnsteadyAdsorption> SHARK_DATA::UnsteadyAdsList

Unsteady Adsorption Reaction Objects.

 $5.45.2.7 \quad std:: vector < \textbf{MultiligandAdsorption} > SHARK_DATA:: MultiAdsList$

Multiligand Adsorption Objects.

 $5.45.2.8 \quad std:: vector < \textbf{ChemisorptionReaction} > \textbf{SHARK_DATA}:: ChemisorptionList$

Chemisorption Reaction objects.

5.45.2.9 std::vector<MultiligandChemisorption> SHARK_DATA::MultiChemList

Multiligand Chemisorption Reaction Objects.

 $\begin{array}{ll} \textbf{5.45.2.10} & \textbf{std::vector} < \textbf{double (*) (const Matrix} < \textbf{double} > \textbf{\&x, SHARK_DATA} * \textbf{shark_dat, const void} * \textbf{data)} > \\ & \textbf{SHARK_DATA::OtherList} \\ \end{array}$

Array of Other Residual functions to be defined by user.

This list of function pointers can be declared and set up by the user in order to add to or change the behavior of the SHARK system. Each one must be declared setup individually by the user. They will be called by the shark __residual function when needed. Alternatively, the user is free to provide their own shark_residual function for the overall system.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
shark_dat	pointer to the SHARK_DATA data structure
data	pointer to a user defined data structure that is used to evaluate this residual

5.45.2.11 int SHARK_DATA::numvar

Total number of functions and species.

5.45.2.12 int SHARK_DATA::num_ssr

Number of steady-state reactions.

5.45.2.13 int SHARK_DATA::num_mbe

Number of mass balance equations.

5.45.2.14 int SHARK_DATA::num_usr = 0

Number of unsteady-state reactions.

5.45.2.15 int SHARK_DATA::num_ssao = 0

Number of steady-state adsorption objects.

5.45.2.16 int SHARK_DATA::num_usao = 0

Number of unsteady adsorption objects.

5.45.2.17 int SHARK_DATA::num_multi_ssao = 0

Number of multiligand steady-state adsorption objects.

5.45.2.18 int SHARK_DATA::num_sschem = 0

Number of steady-state chemisorption objects.

5.45.2.19 int SHARK_DATA::num_multi_sschem = 0

Number of multiligand steady-state chemisorption objects.

5.45.2.20 std::vector<int> SHARK_DATA::num_ssar

List of the numbers of reactions in each adsorption object.

5.45.2.21 std::vector<int> SHARK_DATA::num_usar

List of the numbers of reactions in each unsteady adsorption object.

5.45.2.22 std::vector<int> SHARK_DATA::num_sschem_rxns

List of the numbers of reactions in each steady-state chemisorption object.

5.45.2.23 std::vector < std::vector < int > > SHARK_DATA::num_multi_ssar

List of all multiligand objects -> List of ligands and rxns of that ligand.

 $5.45.2.24 \quad std::vector < std::vector < int > > SHARK_DATA::num_multichem_rxns$

List of all multiligand chemisorption objects -> List of num rxns for that ligand.

5.45.2.25 std::vector<std::string> SHARK_DATA::ss_ads_names

List of the steady-state adsorbent object names.

5.45.2.26 std::vector<std::string> SHARK_DATA::us_ads_names

List of the unsteady adsorption object names.

5.45.2.27 std::vector<std::string> SHARK_DATA::ss_chem_names

List of the steady-state chemisorption object names.

5.45.2.28 std::vector< std::vector<std::string> > SHARK_DATA::ssmulti_names

List of the names of the ligands in each multiligand object.

 $5.45.2.29 \quad std::vector < std::string > > SHARK_DATA::ssmultichem_names$

List of the names of the ligands in each multiligand chemisorption object.

5.45.2.30 int SHARK_DATA::num_other = 0

Number of other functions to be used (default is always 0)

5.45.2.31 int SHARK_DATA::act_fun = IDEAL

Flag denoting the activity function to use (default is IDEAL)

5.45.2.32 int SHARK_DATA::reactor_type = BATCH

Flag denoting the type of reactor considered for the system (default is BATCH)

5.45.2.33 int SHARK_DATA::totalsteps = 0

Total number of iterations.

5.45.2.34 int SHARK_DATA::totalcalls = 0

Total number of residual function calls.

5.45.2.35 int SHARK_DATA::timesteps = 0

Number of time steps taken to complete simulation.

5.45.2.36 int SHARK_DATA::pH_index = -1

Contains the index of the pH variable (set internally)

5.45.2.37 int SHARK_DATA::pOH_index = -1

Contains the index of the pOH variable (set internally)

5.45.2.38 double SHARK_DATA::simulationtime = 0.0

Time to simulate unsteady reactions for (default = 0.0 hrs)

5.45.2.39 double SHARK_DATA::dt = 0.1

Time step size (hrs)

5.45.2.40 double SHARK_DATA::dt_min = sqrt(DBL_EPSILON)

Minimum allowable step size.

5.45.2.41 double SHARK_DATA::dt_max = 744.0

Maximum allowable step size (∼1 month in time)

5.45.2.42 double SHARK_DATA::t_out = 0.0

Time increment by which file output is made (default = print all time steps)

5.45.2.43 double SHARK_DATA::t_count = 0.0

Running count of time increments.

5.45.2.44 double SHARK_DATA::time = 0.0

Current value of time (starts from t = 0.0 hrs)

5.45.2.45 double SHARK_DATA::time_old = 0.0

Previous value of time (start from t = 0.0 hrs)

```
5.45.2.46 double SHARK_DATA::pH = 7.0
Value of pH if needed (default = 7)
5.45.2.47 double SHARK_DATA::pH_step = 0.5
Value by which to increment pH when doing a speciation curve (default = 0.5)
5.45.2.48 double SHARK_DATA::start_temp = 277.15
Value of the starting temperature used for Temperature Curves (default = 277.15 K)
5.45.2.49 double SHARK_DATA::end_temp = 323.15
Value of the ending temperature used for Temperature Curves (default = 323.15 K)
5.45.2.50 double SHARK_DATA::temp_step = 10.0
Size of the step changes to use for Temperature Curves (default = 10.0 K);.
5.45.2.51 double SHARK_DATA::volume = 1.0
Volume of the domain in liters (default = 1 L)
5.45.2.52 double SHARK_DATA::flow_rate = 1.0
Flow rate in the reactor in L/hr (default = 1 L/hr)
5.45.2.53 double SHARK_DATA::xsec_area = 1.0
Cross sectional area of the reactor in m^2 (default = 1 m^2)
5.45.2.54 double SHARK_DATA::Norm = 0.0
Current value of euclidean norm in solution.
5.45.2.55 double SHARK_DATA::dielectric_const = 78.325
Dielectric constant used in many activity models (default: water = 78.325 (1/K))
5.45.2.56 double SHARK_DATA::relative_permittivity = 80.1
Relative permittivity of the medium (default: water = 80.1 (-))
5.45.2.57 double SHARK_DATA::temperature = 298.15
Solution temperature (default = 25 oC or 298.15 K)
```

5.45.2.58 double SHARK_DATA::ionic_strength = 0.0

Solution ionic strength in Molar (calculated internally)

5.45.2.59 bool SHARK_DATA::steadystate = true

True = solve steady problem; False = solve transient problem.

5.45.2.60 bool SHARK_DATA::ZeroInitialSolids = false

True = no solids or adsorption initially in the reactor.

5.45.2.61 bool SHARK_DATA::TimeAdaptivity = false

True = solve using variable time step.

5.45.2.62 bool SHARK_DATA::const_pH = false

True = set pH to a constant; False = solve for pH.

5.45.2.63 bool SHARK_DATA::SpeciationCurve = false

True = runs a series of constant pH steady-state problems to produce curves.

5.45.2.64 bool SHARK_DATA::TemperatureCurve = false

True = runs a series of constant temperature steady-state problmes to produce curves.

5.45.2.65 bool SHARK_DATA::Console_Output = true

True = display output to console.

5.45.2.66 bool SHARK_DATA::File_Output = false

True = write output to a file.

5.45.2.67 bool SHARK_DATA::Contains_pH = false

True = system contains pH as a variable (set internally)

5.45.2.68 bool SHARK_DATA::Contains_pOH = false

True = system contains pOH as a variable (set internally)

5.45.2.69 bool SHARK_DATA::Converged = false

True = system converged within tolerance.

5.45.2.70 bool SHARK_DATA::LocalMin = true

True = allow the system to settle for a local minimum if tolerance not reached.

5.45.2.71 Matrix<double> SHARK_DATA::X_old

Solution vector for old time step - log(C)

5.45.2.72 Matrix < double > SHARK_DATA::X_new

Solution vector for current time step - log(C)

5.45.2.73 Matrix < double > SHARK_DATA::Conc_old

Concentration vector for old time step - 10° x.

5.45.2.74 Matrix < double > SHARK_DATA::Conc_new

Concentration vector for current time step - $10^{\land}x$.

5.45.2.75 Matrix < double > SHARK_DATA::activity_new

Activity matrix for current time step.

5.45.2.76 Matrix<double> SHARK_DATA::activity_old

Activity matrix from prior time step.

5.45.2.77 int(* SHARK_DATA::EvalActivity) (const Matrix < double > &x, Matrix < double > &F, const void *data)

Function pointer to evaluate activity coefficients.

This function pointer is called within the shark_residual function to calculate and modify the activity_new matrix entries. When using the SHARK default options, this function pointer will be automatically set to a cooresponding activity function for the list of valid functions from the valid_act enum. User may override this function pointer if they desire. Must be overriden after calling the setup function.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to a data structure needed to evaluate the activity model

5.45.2.78 int(* SHARK_DATA::Residual) (const Matrix< double > &x, Matrix< double > &F, const void *data)

Function pointer to evaluate all residuals in the system.

This function will be fed into the PJFNK solver (see lark.h) to solve the non-linear system of equations. By default, this pointer will be the shark_residual function (see below). However, the user may override the function and provide their own residuals for the PJFNK solver to operate on.

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of residuals that are to be altered from the functions in the system
data	pointer to a data structure needed to evaluate the activity model

5.45.2.79 int(* SHARK_DATA::lin_precon) (const Matrix < double > &r, Matrix < double > &p, const void *data)

Function pointer to form a linear preconditioning operation for the Jacobian.

This function will be fed into the linear solver used for each non-linear step in PJFNK (see lark.h). By default, we cannot provide any linear preconditioner, because we do not know the form or sparcity of the Jacobian before hand. It will be the user's responsibility to form their own preconditioner until we can figure out a generic way to precondition the system.

5.45.2.80 PJFNK DATA SHARK_DATA::Newton_data

Data structure for the Newton-Krylov solver (see lark.h)

5.45.2.81 const void* SHARK_DATA::activity_data

User defined data structure for an activity model.

5.45.2.82 const void* SHARK_DATA::residual_data

User defined data structure for the residual function.

5.45.2.83 const void* SHARK_DATA::precon_data

User defined data structure for preconditioning.

5.45.2.84 const void* SHARK_DATA::other_data

User define data structure used for user defined residuals.

5.45.2.85 FILE* SHARK_DATA::OutputFile

Output File pointer.

5.45.2.86 yaml_cpp_class SHARK_DATA::yaml_object

yaml object to read and access digitized yaml documents (see yaml_wrapper.h)

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

· shark.h

5.46 SKUA_DATA Struct Reference

Data structure for all simulation information in SKUA.

```
#include <skua.h>
```

Public Attributes

· unsigned long int total steps

Running total of all calculation steps.

int coord

Used to determine the coordinates of the problem.

· double sim_time

Stopping time for the simulation (hrs)

double t old

Old time of the simulations (hrs)

· double t

Current time of the simulations (hrs)

• double t counter = 0.0

Counts for print times for output (hrs)

double t_print

Prints out every t_print time (hrs)

double qTn

Old total amounts adsorbed (mol/kg)

double qTnp1

New total amounts adsorbed (mol/kg)

• bool Print2File = true

True = results to .txt; False = no printing.

• bool Print2Console = true

True = results to console; False = no printing.

double gas_velocity

Superficial Gas Velocity arount pellet (cm/s)

double pellet_radius

Nominal radius of the pellet/crystal (um)

· double char measure

Length or Area if in Cylindrical or Cartesian coordinates (um or um^{^2})

• bool DirichletBC = true

True = Dirichlet BC; False = Neumann BC.

• bool NonLinear = true

True = Non-linear solver; False = Linear solver.

std::vector< double > y

Outside mole fractions of each component (-)

FILE * OutputFile

Output file pointer to the output file.

double(* eval diff)(int i, int I, const void *user data)

Function pointer for evaluating surface diffusivity.

double(* eval_kf)(int i, const void *user_data)

Function pointer for evaluating film mass transfer.

• const void * user data

Data structure for user's information needed in parameter functions.

MAGPIE_DATA magpie_dat

Data structure for adsorption equilibria (see magpie.h)

MIXED_GAS * gas_dat

Pointer to the MIXED_GAS data structure (see egret.h)

std::vector< FINCH_DATA > finch_dat

Data structure for adsorption kinetics (see finch.h)

std::vector< SKUA_PARAM > param_dat

Data structure for SKUA specific parameters.

5.46.1 Detailed Description

Data structure for all simulation information in SKUA.

C-style object holding all data, functions, and other objects needed to successfully run a SKUA simulation. This object holds system information, such as boundary condition type, adsorbent size, and total adsorption, and also contains structure for EGRET (egret.h), FINCH (finch.h), and MAGPIE (magpie.h) calculations. Function pointers for evaluation of the surface diffusivity and film mass transfer coefficients can be overriden by the user to change the behavior of the SKUA simulation. However, defaults are also provided for these functions.

5.46.2 Member Data Documentation

5.46.2.1 unsigned long int SKUA_DATA::total_steps

Running total of all calculation steps.

5.46.2.2 int SKUA_DATA::coord

Used to determine the coordinates of the problem.

5.46.2.3 double SKUA_DATA::sim_time

Stopping time for the simulation (hrs)

5.46.2.4 double SKUA_DATA::t_old

Old time of the simulations (hrs)

5.46.2.5 double SKUA_DATA::t

Current time of the simulations (hrs)

5.46.2.6 double SKUA_DATA::t_counter = 0.0

Counts for print times for output (hrs)

5.46.2.7 double SKUA_DATA::t_print

Prints out every t_print time (hrs)

```
5.46.2.8 double SKUA_DATA::qTn
```

Old total amounts adsorbed (mol/kg)

5.46.2.9 double SKUA_DATA::qTnp1

New total amounts adsorbed (mol/kg)

5.46.2.10 bool SKUA_DATA::Print2File = true

True = results to .txt; False = no printing.

5.46.2.11 bool SKUA_DATA::Print2Console = true

True = results to console; False = no printing.

5.46.2.12 double SKUA_DATA::gas_velocity

Superficial Gas Velocity arount pellet (cm/s)

5.46.2.13 double SKUA_DATA::pellet_radius

Nominal radius of the pellet/crystal (um)

5.46.2.14 double SKUA_DATA::char_measure

Length or Area if in Cylindrical or Cartesian coordinates (um or um²)

5.46.2.15 bool SKUA_DATA::DirichletBC = true

True = Dirichlet BC; False = Neumann BC.

5.46.2.16 bool SKUA_DATA::NonLinear = true

True = Non-linear solver; False = Linear solver.

5.46.2.17 std::vector<double> SKUA_DATA::y

Outside mole fractions of each component (-)

5.46.2.18 FILE* SKUA_DATA::OutputFile

Output file pointer to the output file.

5.46.2.19 double(* SKUA_DATA::eval_diff) (int i, int I, const void *user_data)

Function pointer for evaluating surface diffusivity.

5.46.2.20 double(* SKUA_DATA::eval_kf) (int i, const void *user_data)

Function pointer for evaluating film mass transfer.

5.46.2.21 const void* SKUA_DATA::user_data

Data structure for user's information needed in parameter functions.

5.46.2.22 MAGPIE_DATA SKUA_DATA::magpie_dat

Data structure for adsorption equilibria (see magpie.h)

5.46.2.23 MIXED_GAS* SKUA_DATA::gas_dat

Pointer to the MIXED_GAS data structure (see egret.h)

5.46.2.24 std::vector<FINCH_DATA> SKUA_DATA::finch_dat

Data structure for adsorption kinetics (see finch.h)

5.46.2.25 std::vector < SKUA_PARAM > SKUA_DATA::param_dat

Data structure for SKUA specific parameters.

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

• skua.h

5.47 SKUA_OPT_DATA Struct Reference

Data structure for the SKUA Optimization Routine.

```
#include <skua_opt.h>
```

Public Attributes

• int num_curves

Number of adsorption curves to analyze.

· int evaluation

Number of times the eval function has been called for a single curve.

· unsigned long int total_eval

Total number of evaluations needed for completion.

· int current_points

Number of points in the current curve.

int num_params = 1

Number of adjustable parameters for the current curve.

· int diffusion type

Flag to identify type of diffusion function to use.

• int adsorb_index

Component index for adsorbable species.

• int max_guess_iter = 20

Maximum allowed guess iterations (default = 20)

· bool Optimize

True = run optimization, False = run a comparison.

· bool Rough

True = use only a rough estimate, False = run full optimization.

· double current_temp

Temperature for current curve.

· double current_press

Partial pressure for current curve.

• double current_equil

Equilibrium data point for the current curve.

· double simulation_equil

Equilibrium simulation point for the current curve.

· double max_bias

Positive maximum bias plausible for fitting.

• double min_bias

Negative minimum bias plausible for fitting.

double e_norm

Euclidean norm of current fit.

· double f bias

Function bias of current fit.

• double e_norm_old

Euclidean norm of the previous fit.

• double f_bias_old

Function bias of the previous fit.

• double param_guess

Parameter guess for the surface/crystal diffusivity.

· double param_guess_old

Parameter guess for the previous curve.

• double rel_tol_norm = 0.1

Tolerance for convergence of the guess norm.

• double abs_tol_bias = 0.1

Tolerance for convergence of the guess bias.

std::vector< double > y_base

Gas phase mole fractions in absense of adsorbing species.

std::vector< double > q_data

Amount adsorbed at a particular point in current curve.

std::vector< double > q_sim

Amount adsorbed based on the simulation.

• std::vector < double > t

Time points in the current curve.

FILE * ParamFile

Output file for parameter results.

FILE * CompareFile

Output file for comparison of results.

SKUA_DATA skua_dat

Data structure for the SKUA simulation.

5.47.1 Detailed Description

Data structure for the SKUA Optimization Routine.

C-style object holding data and pointers necessary for running a SKUA optimization. It contains information about the type of optimization requested, the current status of the optimization, the data being compared against, and the SKUA_DATA object for the evaluation of a SKUA simulation. The pointers in the structure are for the two output files produced by the routine: (i) parameter results and (ii) model comparison results.

5.47.2 Member Data Documentation

5.47.2.1 int SKUA_OPT_DATA::num_curves

Number of adsorption curves to analyze.

5.47.2.2 int SKUA_OPT_DATA::evaluation

Number of times the eval function has been called for a single curve.

5.47.2.3 unsigned long int SKUA_OPT_DATA::total_eval

Total number of evaluations needed for completion.

5.47.2.4 int SKUA_OPT_DATA::current_points

Number of points in the current curve.

5.47.2.5 int SKUA_OPT_DATA::num_params = 1

Number of adjustable parameters for the current curve.

5.47.2.6 int SKUA_OPT_DATA::diffusion_type

Flag to identify type of diffusion function to use.

5.47.2.7 int SKUA_OPT_DATA::adsorb_index

Component index for adsorbable species.

5.47.2.8 int SKUA_OPT_DATA::max_guess_iter = 20

Maximum allowed guess iterations (default = 20)

5.47.2.9 bool SKUA_OPT_DATA::Optimize

True = run optimization, False = run a comparison.

5.47.2.10 bool SKUA_OPT_DATA::Rough

True = use only a rough estimate, False = run full optimization.

5.47.2.11 double SKUA_OPT_DATA::current_temp

Temperature for current curve.

5.47.2.12 double SKUA_OPT_DATA::current_press

Partial pressure for current curve.

5.47.2.13 double SKUA_OPT_DATA::current_equil

Equilibrium data point for the current curve.

5.47.2.14 double SKUA_OPT_DATA::simulation_equil

Equilibrium simulation point for the current curve.

5.47.2.15 double SKUA_OPT_DATA::max_bias

Positive maximum bias plausible for fitting.

5.47.2.16 double SKUA_OPT_DATA::min_bias

Negative minimum bias plausible for fitting.

5.47.2.17 double SKUA_OPT_DATA::e_norm

Euclidean norm of current fit.

5.47.2.18 double SKUA_OPT_DATA::f_bias

Function bias of current fit.

5.47.2.19 double SKUA_OPT_DATA::e_norm_old

Euclidean norm of the previous fit.

5.47.2.20 double SKUA_OPT_DATA::f_bias_old

Function bias of the previous fit.

5.47.2.21 double SKUA_OPT_DATA::param_guess

Parameter guess for the surface/crystal diffusivity.

5.47.2.22 double SKUA_OPT_DATA::param_guess_old

Parameter guess for the previous curve.

5.47.2.23 double SKUA_OPT_DATA::rel_tol_norm = 0.1

Tolerance for convergence of the guess norm.

5.47.2.24 double SKUA_OPT_DATA::abs_tol_bias = 0.1

Tolerance for convergence of the guess bias.

5.47.2.25 std::vector<double> SKUA_OPT_DATA::y_base

Gas phase mole fractions in absense of adsorbing species.

5.47.2.26 std::vector<double> SKUA_OPT_DATA::q_data

Amount adsorbed at a particular point in current curve.

5.47.2.27 std::vector<double> SKUA_OPT_DATA::q_sim

Amount adsorbed based on the simulation.

5.47.2.28 std::vector<double> SKUA_OPT_DATA::t

Time points in the current curve.

5.47.2.29 FILE* SKUA_OPT_DATA::ParamFile

Output file for parameter results.

5.47.2.30 FILE* SKUA_OPT_DATA::CompareFile

Output file for comparison of results.

5.47.2.31 SKUA DATA SKUA_OPT_DATA::skua_dat

Data structure for the SKUA simulation.

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

skua_opt.h

5.48 SKUA_PARAM Struct Reference

Data structure for species' parameters in SKUA.

#include <skua.h>

Public Attributes

- double activation_energy
- double ref_diffusion
- double ref temperature
- double affinity
- double ref_pressure
- double film_transfer
- double xIC
- double y_eff
- · double Qstn
- double Qstnp1
- double xn
- double xnp1
- · bool Adsorbable
- std::string speciesName

5.48.1 Detailed Description

Data structure for species' parameters in SKUA.

C-style object holding data and parameters associated with the gas/solid species in the overall SKUA system. These parameters are used in to modify surface diffusivity with temperature, establish film mass transfer coefficients, formulate the initial conditions, and store solution results for heat of adsorption and adsorbed mole fractions. One of these objects will be created for each species in the gas system.

5.48.2	Member Data Documentation
5.48.2.1	double SKUA_PARAM::activation_energy
5.48.2.2	double SKUA_PARAM::ref_diffusion
5.48.2.3	double SKUA_PARAM::ref_temperature
5.48.2.4	double SKUA_PARAM::affinity
5.48.2.5	double SKUA_PARAM::ref_pressure
5.48.2.6	double SKUA_PARAM::film_transfer
5.48.2.7	double SKUA_PARAM::xIC
5.48.2.8	double SKUA_PARAM::y_eff
5.48.2.9	double SKUA_PARAM::Qstn
5.48.2.10	double SKUA_PARAM::Qstnp1
5.48.2.11	double SKUA_PARAM::xn
5.48.2.12	double SKUA_PARAM::xnp1
5.48.2.13	B bool SKUA_PARAM::Adsorbable
5.48.2.14	std::string SKUA_PARAM::speciesName

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

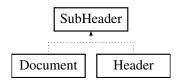
• skua.h

5.49 SubHeader Class Reference

Object for the Lowest level of Header for the yaml_wrapper.

```
#include <yaml_wrapper.h>
```

Inheritance diagram for SubHeader:



Public Member Functions

· SubHeader ()

Default Constructor.

∼SubHeader ()

Default Destructor.

• SubHeader (const SubHeader &subheader)

Copy constructor.

SubHeader (const KeyValueMap &map)

Construction by existing map.

SubHeader (std::string name)

Construction by name only.

SubHeader (std::string name, const KeyValueMap &map)

Construction by name and map.

• SubHeader & operator= (const SubHeader &sub)

Equals overload.

ValueTypePair & operator[] (const std::string key)

Return the ValueType reference at the given key.

ValueTypePair operator[] (const std::string key) const

Return the ValueType at the give key.

KeyValueMap & getMap ()

Returns reference to the KeyValueMap object.

• void clear ()

Empty out data contents.

void addPair (std::string key, std::string val)

Adds a pair object to the map (with only strings)

• void addPair (std::string key, std::string val, int type)

Adds a pair object and asserts a type.

void setName (std::string name)

Sets the name of the subheader.

· void setAlias (std::string alias)

Set the alias without type specification.

void setAlias (std::string alias, int state)

Sets the alias and state of the subheader.

void setNameAliasPair (std::string name, std::string alias, int state)

Sets the name and alias of the subheader.

void setState (int state)

Sets the state of the subheader.

void DisplayContents ()

Display the contents of the subheader.

• std::string getName ()

Return the name of the subheader.

• std::string getAlias ()

Return the alias of the subheader, if one exists.

• bool isAlias ()

Returns true if subheader is an alias.

· bool isAnchor ()

Returns true if subheader is an anchor.

• int getState ()

Returns the state of the subheader.

Protected Attributes

KeyValueMap Data_Map

A Map of Keys and Values.

• std::string name

Name of the subheader.

· std::string alias

Name of the alias for the subheader.

int state

State of the header.

5.49.1 Detailed Description

Object for the Lowest level of Header for the yaml_wrapper.

C++ Object for sub-headers in a yaml document. This object contains a KeyValueMap that holds a set of key-value pairs for data listed under a sub-header in yaml files. It is the lowest allowable recursion of headers in a yaml document and so is the base class for Header and Document, which themselves can contain KeyValueMaps as well as maps for other header-like objects.

SubHeaders are recognized by a unique name and/or alias while being put together in other higher document structures. Additionally, each header or sub-header will have a state to denote whether the object is a yaml alias, anchor, or niether. This is used in the yaml documents to ensure that aliases for anchors have the correct data moved over into the new structures.

5.49.2 Constructor & Destructor Documentation
5.49.2.1 SubHeader::SubHeader ()
Default Constructor.
5.49.2.2 SubHeader::~SubHeader ()

Default Destructor.

```
5.49.2.3 SubHeader::SubHeader ( const SubHeader & subheader )
Copy constructor.
5.49.2.4 SubHeader::SubHeader ( const KeyValueMap & map )
Construction by existing map.
5.49.2.5 SubHeader::SubHeader ( std::string name )
Construction by name only.
5.49.2.6 SubHeader::SubHeader ( std::string name, const KeyValueMap & map )
Construction by name and map.
5.49.3 Member Function Documentation
5.49.3.1 SubHeader & SubHeader::operator= ( const SubHeader & sub )
Equals overload.
5.49.3.2 ValueTypePair& SubHeader::operator[] ( const std::string key )
Return the ValueType reference at the given key.
5.49.3.3 ValueTypePair SubHeader::operator[] ( const std::string key ) const
Return the ValueType at the give key.
5.49.3.4 KeyValueMap& SubHeader::getMap ( )
Returns reference to the KeyValueMap object.
5.49.3.5 void SubHeader::clear ( )
Empty out data contents.
5.49.3.6 void SubHeader::addPair ( std::string key, std::string val )
Adds a pair object to the map (with only strings)
5.49.3.7 void SubHeader::addPair ( std::string key, std::string val, int type )
Adds a pair object and asserts a type.
5.49.3.8 void SubHeader::setName ( std::string name )
Sets the name of the subheader.
```

```
5.49.3.9 void SubHeader::setAlias ( std::string alias )
Set the alias without type specification.
5.49.3.10 void SubHeader::setAlias ( std::string alias, int state )
Sets the alias and state of the subheader.
5.49.3.11 void SubHeader::setNameAliasPair ( std::string name, std::string alias, int state )
Sets the name and alias of the subheader.
5.49.3.12 void SubHeader::setState (int state)
Sets the state of the subheader.
5.49.3.13 void SubHeader::DisplayContents ( )
Display the contents of the subheader.
5.49.3.14 std::string SubHeader::getName ( )
Return the name of the subheader.
5.49.3.15 std::string SubHeader::getAlias ( )
Return the alias of the subheader, if one exists.
5.49.3.16 bool SubHeader::isAlias ( )
Returns true if subheader is an alias.
5.49.3.17 bool SubHeader::isAnchor ( )
Returns true if subheader is an anchor.
5.49.3.18 int SubHeader::getState ( )
Returns the state of the subheader.
5.49.4 Member Data Documentation
5.49.4.1 KeyValueMap SubHeader::Data_Map [protected]
A Map of Keys and Values.
5.49.4.2 std::string SubHeader::name [protected]
Name of the subheader.
```

```
5.49.4.3 std::string SubHeader::alias [protected]
Name of the alias for the subheader.
5.49.4.4 int SubHeader::state [protected]
State of the header.
The documentation for this class was generated from the following file:
    yaml_wrapper.h
5.50 SYSTEM_DATA Struct Reference
System Data Structure.
#include <magpie.h>
Public Attributes

    double T

          System Temperature (K)
    · double PT
           Total Pressure (kPa)

    double qT

           Total Amount adsorbed (mol/kg)

    double PI

           Total Lumped Spreading Pressure (mol/kg)
          Actual Spreading pressure (J/m^2)

    double As

          Specific surface area of adsorbent (m<sup>^</sup>2/kg)

 int N

           Total Number of Components.
    int I

 int J

    • int K
          Special indices used to keep track of sub-systems.
    · unsigned long int total eval
          Counter to keep track of total number of non-linear steps.
    · double avg_norm
           Used to store all norms from evaluations then average at end of run.

    double max_norm

          Used to store the maximum e.norm calculated from non-linear iterations.
    • int Sys
          Number of sub-systems to solve.
    • int Par
          Number of binary parameters to solve for.

    bool Recover

          If Recover == false, standard GPAST using y's as knowns.

    bool Carrier

          If there is an inert carrier gas, Carrier == true.

    bool Ideal

          If the behavior of the system is determined to be ideal, then Ideal == true.
```

Boolean to suppress output if desired (true = display, false = no display.

bool Output

5.50.1 Detailed Description System Data Structure. C-style object holding all the data associated with the overall system to be modeled. 5.50.2 Member Data Documentation 5.50.2.1 double SYSTEM_DATA::T System Temperature (K) 5.50.2.2 double SYSTEM_DATA::PT Total Pressure (kPa) 5.50.2.3 double SYSTEM_DATA::qT Total Amount adsorbed (mol/kg) 5.50.2.4 double SYSTEM_DATA::PI Total Lumped Spreading Pressure (mol/kg) 5.50.2.5 double SYSTEM_DATA::pi Actual Spreading pressure (J/m²) 5.50.2.6 double SYSTEM_DATA::As

Specific surface area of adsorbent (m^2/kg)

5.50.2.7 int SYSTEM_DATA::N

Total Number of Components.

5.50.2.8 int SYSTEM_DATA::I

5.50.2.9 int SYSTEM_DATA::J

5.50.2.10 int SYSTEM_DATA::K

Special indices used to keep track of sub-systems.

5.50.2.11 unsigned long int SYSTEM_DATA::total_eval

Counter to keep track of total number of non-linear steps.

5.50.2.12 double SYSTEM_DATA::avg_norm

Used to store all norms from evaluations then average at end of run.

5.50.2.13 double SYSTEM_DATA::max_norm

Used to store the maximum e.norm calculated from non-linear iterations.

5.50.2.14 int SYSTEM_DATA::Sys

Number of sub-systems to solve.

5.50.2.15 int SYSTEM_DATA::Par

Number of binary parameters to solve for.

5.50.2.16 bool SYSTEM_DATA::Recover

If Recover == false, standard GPAST using y's as knowns.

5.50.2.17 bool SYSTEM_DATA::Carrier

If there is an inert carrier gas, Carrier == true.

5.50.2.18 bool SYSTEM_DATA::Ideal

If the behavior of the system is determined to be ideal, then Ideal == true.

5.50.2.19 bool SYSTEM_DATA::Output

Boolean to suppress output if desired (true = display, false = no display.

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

· magpie.h

5.51 TRAJECTORY_DATA Struct Reference

#include <Trajectory.h>

Public Attributes

• double $mu_0 = 12.57e-7$

permeability of free space, H/m

double rho_f = 1000.0

Fluid density, Kg/m3.

- double eta = 0.001
- double Hamaker = 1.3e-21
- double Temp = 298
- double k = 1.38e-23
- double Rs
- double L
- · double porosity
- double V_separator
- double a
- double V_wire
- double L_wire
- double A_separator
- double A_wire
- double B0
- double H0
- double Ms = 0.6
- double b
- double chi_p
- double rho_p = 8000.0
- double Q_in
- double V0
- double Y_initial = 20.0
- double dt
- double M
- double mp
- double beta
- double q bar
- double sigma_v
- double sigma_vz
- double sigma_z
- double sigma_n
- double sigma_m
- double n_rand
- double m_rand
- double s rand
- double t_rand
- Matrix< double > POL
- Matrix< double > H
- Matrix< double > dX
- Matrix< double > dY
- Matrix< double > Vr
- Matrix< double > Vt
- Matrix< double > X
- Matrix< double > Y
- Matrix< int > Cap

- 5.51.1 Member Data Documentation
- 5.51.1.1 double TRAJECTORY_DATA::mu_0 = 12.57e-7

permeability of free space, H/m

5.51.1.2 double TRAJECTORY_DATA::rho_f = 1000.0

Fluid density, Kg/m3.

- 5.51.1.3 double TRAJECTORY_DATA::eta = 0.001
- 5.51.1.4 double TRAJECTORY_DATA::Hamaker = 1.3e-21
- 5.51.1.5 double TRAJECTORY_DATA::Temp = 298
- 5.51.1.6 double TRAJECTORY_DATA::k = 1.38e-23
- 5.51.1.7 double TRAJECTORY_DATA::Rs
- 5.51.1.8 double TRAJECTORY_DATA::L
- 5.51.1.9 double TRAJECTORY_DATA::porosity
- 5.51.1.10 double TRAJECTORY_DATA::V_separator
- 5.51.1.11 double TRAJECTORY_DATA::a
- 5.51.1.12 double TRAJECTORY_DATA::V_wire
- 5.51.1.13 double TRAJECTORY_DATA::L_wire
- 5.51.1.14 double TRAJECTORY_DATA::A_separator
- 5.51.1.15 double TRAJECTORY_DATA::A_wire
- 5.51.1.16 double TRAJECTORY_DATA::B0
- 5.51.1.17 double TRAJECTORY_DATA::H0
- 5.51.1.18 double TRAJECTORY_DATA::Ms = 0.6
- 5.51.1.19 double TRAJECTORY_DATA::b
- 5.51.1.20 double TRAJECTORY_DATA::chi_p
- 5.51.1.21 double TRAJECTORY_DATA::rho_p = 8000.0
- 5.51.1.22 double TRAJECTORY_DATA::Q_in

5.51.1.23	double TRAJECTORY_DATA::V0
5.51.1.24	double TRAJECTORY_DATA::Y_initial = 20.0
5.51.1.25	double TRAJECTORY_DATA::dt
5.51.1.26	double TRAJECTORY_DATA::M
5.51.1.27	double TRAJECTORY_DATA::mp
5.51.1.28	double TRAJECTORY_DATA::beta
5.51.1.29	double TRAJECTORY_DATA::q_bar
5.51.1.30	double TRAJECTORY_DATA::sigma_v
5.51.1.31	double TRAJECTORY_DATA::sigma_vz
5.51.1.32	double TRAJECTORY_DATA::sigma_z
5.51.1.33	double TRAJECTORY_DATA::sigma_n
5.51.1.34	double TRAJECTORY_DATA::sigma_m
5.51.1.35	double TRAJECTORY_DATA::n_rand
5.51.1.36	double TRAJECTORY_DATA::m_rand
5.51.1.37	double TRAJECTORY_DATA::s_rand
5.51.1.38	double TRAJECTORY_DATA::t_rand
5.51.1.39	Matrix <double> TRAJECTORY_DATA::POL</double>
5.51.1.40	Matrix <double> TRAJECTORY_DATA::H</double>
5.51.1.41	Matrix <double> TRAJECTORY_DATA::dX</double>
5.51.1.42	Matrix <double> TRAJECTORY_DATA::dY</double>
5.51.1.43	Matrix <double> TRAJECTORY_DATA::Vr</double>
5.51.1.44	Matrix <double> TRAJECTORY_DATA::Vt</double>
5.51.1.45	Matrix <double> TRAJECTORY_DATA::X</double>
5.51.1.46	Matrix <double> TRAJECTORY_DATA::Y</double>
5.51.1.47	Matrix <int> TRAJECTORY_DATA::Cap</int>

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

• Trajectory.h

5.52 UI_DATA Struct Reference

Data structure holding the UI arguments.

```
#include <ui.h>
```

Public Attributes

ValueTypePair value_type

Data pair for input, tells what the input is and it's type.

std::vector< std::string > user_input

What is read in from the console at any point.

std::vector< std::string > input_files

A vector of input file names and directories given by user.

· std::string path

Path to where input files are located.

• int count = 0

Number of times a questing has been asked.

• int max = 3

Maximum allowable recursions of a question.

· int option

Current option choosen by the user.

• bool Path = false

True if user gives path as an option.

bool Files = false

True if user gives input files as an option.

• bool MissingArg = true

True if an input argument is missing; False if everything is ok.

bool BasicUI = true

True if using Basic UI; False if using Advanced UI.

· int argc

Number of console arguments given on input.

const char * argv []

Actual console arguments given at execution.

5.52.1 Detailed Description

Data structure holding the UI arguments.

C-Style object for interfacing with users request upon execution of the program. User input is stored in objects below and a series of booleans is used to determine how and what to execute.

5.52.2 Member Data Documentation

5.52.2.1 ValueTypePair UI_DATA::value_type

Data pair for input, tells what the input is and it's type.

5.52.2.2 std::vector<std::string> UI_DATA::user_input

What is read in from the console at any point.

5.52.2.3 std::vector<std::string> UI_DATA::input_files

A vector of input file names and directories given by user.

5.52.2.4 std::string UI_DATA::path

Path to where input files are located.

5.52.2.5 int UI_DATA::count = 0

Number of times a questing has been asked.

5.52.2.6 int UI_DATA::max = 3

Maximum allowable recursions of a question.

5.52.2.7 int UI_DATA::option

Current option choosen by the user.

5.52.2.8 bool UI_DATA::Path = false

True if user gives path as an option.

5.52.2.9 bool UI_DATA::Files = false

True if user gives input files as an option.

 $5.52.2.10 \quad bool \ UI_DATA::MissingArg = true$

True if an input argument is missing; False if everything is ok.

5.52.2.11 bool UI_DATA::BasicUI = true

True if using Basic UI; False if using Advanced UI.

5.52.2.12 int UI_DATA::argc

Number of console arguments given on input.

5.52.2.13 const char* UI_DATA::argv[]

Actual console arguments given at execution.

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

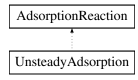
• ui.h

5.53 UnsteadyAdsorption Class Reference

Unsteady Adsorption Reaction Object.

#include <shark.h>

Inheritance diagram for UnsteadyAdsorption:



Public Member Functions

• UnsteadyAdsorption ()

Default Constructor.

∼UnsteadyAdsorption ()

Default Destructor.

void Initialize_Object (MasterSpeciesList &List, int n)

Function to call the initialization of objects sequentially.

void Display Info ()

Display the adsorption reaction information (PLACE HOLDER)

void modifyDeltas (MassBalance &mbo)

Modify the Deltas in the MassBalance Object.

• int setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object.

• int checkAqueousIndices ()

Function to check and report errors in the aqueous species indices.

 void setActivityModelInfo (int(*act)(const Matrix< double > &logq, Matrix< double > &activity, const void *data), const void *act_data)

Function to set the surface activity model and data pointer.

• void setAqueousIndex (int rxn_i, int species_i)

Set the primary aqueous species index for the ith reaction.

int setAqueousIndexAuto ()

Automatically sets the primary aqueous species index based on reactions.

void setActivityEnum (int act)

Set the surface activity enum value.

void setMolarFactor (int rxn_i, double m)

Set the molar factor for the ith reaction (mol/mol)

void setVolumeFactor (int i, double v)

Set the ith volume factor for the species list (cm[^]3/mol)

• void setAreaFactor (int i, double a)

Set the ith area factor for the species list (m^2/mol)

void setSpecificArea (double a)

Set the specific area for the adsorbent (m\^2/kg)

void setSpecificMolality (double a)

Set the specific molality for the adsorbent (mol/kg)

• void setSurfaceCharge (double c)

Set the surface charge of the uncomplexed ligands.

void setTotalMass (double m)

Set the total mass of the adsorbent (kg)

void setTotalVolume (double v)

Set the total volume of the system (L)

void setAreaBasisBool (bool opt)

Set the basis boolean directly.

void setSurfaceChargeBool (bool opt)

Set the boolean for inclusion of surface charging.

• void setBasis (std::string option)

Set the basis of the adsorption problem from the given string arg.

void setAdsorbentName (std::string name)

Set the name of the adsorbent to the given string.

void updateActivities ()

Set the old activities as the new activities before doing next time step.

void calculateAreaFactors ()

Calculates the area factors used from the van der Waals volumes.

void calculateEquilibria (double T)

Calculates all equilibrium parameters as a function of temperature.

• void calculateRates (double T)

Calculates all reaction rate parameters as a function of temperature.

void setChargeDensity (const Matrix< double > &x)

Calculates and sets the current value of charge density.

void setlonicStrength (const Matrix< double > &x)

Calculates and sets the current value of ionic strength.

int callSurfaceActivity (const Matrix< double > &x)

Calls the activity model and returns an int flag for success or failure.

double calculateActiveFraction (const Matrix< double > &x)

Calculates the fraction of the surface that is active and available.

double calculateSurfaceChargeDensity (const Matrix< double > &x)

Function to calculate the surface charge density based on concentrations.

• double calculatePsi (double sigma, double T, double I, double rel epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

double calculateAqueousChargeExchange (int i)

Function to calculate the net exchange of charges of the ageous species involved in a given reaction.

double calculateEquilibriumCorrection (double sigma, double T, double I, double rel_epsilon, int i)

Function to calculate the correction term for the equilibrium parameter.

double Eval_Residual (const Matrix< double > &x, const Matrix< double > &gama, double T, double rel_
 perm, int i)

Calculates the residual for the ith reaction in the system.

double Eval_Residual (const Matrix< double > &x_new, const Matrix< double > &x_old, const Matrix
 double > &gama new, const Matrix< double > &gama old, double T, double rel perm, int i)

Calculates the unsteady residual for the ith reaction in the system.

double Eval_ReactionRate (const Matrix< double > &x, const Matrix< double > &gama, double T, double rel_perm, int i)

Function to calculate the explicit or implicit rate of reaction.

double Eval_IC_Residual (const Matrix< double > &x, int i)

Calculate the unsteady residual for initial conditions.

double Explicit_Eval (const Matrix < double > &x, const Matrix < double > &gama, double T, double rel_perm, int i)

Return an approximate explicit solution to our unsteady adsorption variable (mol/kg)

• UnsteadyReaction & getReaction (int i)

Return reference to the ith reaction object in the adsorption object.

double getMolarFactor (int i)

Get the ith reaction's molar factor for adsorption (mol/mol)

double getVolumeFactor (int i)

Get the ith volume factor (species not involved return zeros) (cm[^]3/mol)

double getAreaFactor (int i)

Get the ith area factor (species not involved return zeros) (m²/mol)

• double getActivity (int i)

Get the ith activity factor for the surface species.

double getOldActivity (int i)

Get the ith old activity factor for the surface species.

double getSpecificArea ()

Get the specific area of the adsorbent (m^2/kg) or (mol/kg)

double getSpecificMolality ()

Get the specific molality of the adsorbent (mol/kg)

• double getSurfaceCharge ()

Get the surface charge of the adsorbent.

double getBulkDensity ()

Calculate and return bulk density of adsorbent in system (kg/L)

double getTotalMass ()

Get the total mass of adsorbent in the system (kg)

double getTotalVolume ()

Get the total volume of the system (L)

double getChargeDensity ()

Get the value of the surface charge density (C/m^2)

• double getlonicStrength ()

Get the value of the ionic strength of solution (mol/L)

• int getNumberRxns ()

Get the number of reactions involved in the adsorption object.

• int getAdsorbIndex (int i)

Get the index of the adsorbed species in the ith reaction.

• int getAqueousIndex (int i)

Get the index of the primary aqueous species in the ith reaction.

int getActivityEnum ()

Return the enum representing the choosen activity function.

• bool isAreaBasis ()

Returns true if we are in the Area Basis, False if in Molar Basis.

• bool includeSurfaceCharge ()

Returns true if we are considering surface charging during adsorption.

• std::string getAdsorbentName ()

Returns the name of the adsorbent as a string.

Protected Attributes

Matrix< double > activities_old

List of the old activities calculated by the activity model.

Private Attributes

std::vector < UnsteadyReaction > ads_rxn

List of reactions involved with adsorption.

Additional Inherited Members

5.53.1 Detailed Description

Unsteady Adsorption Reaction Object.

C++ Object to handle data and functions associated with forumlating unsteady adsorption reactions in a aqueous mixture. Each unique surface in a system will require an instance of this structure.

5.53.2 Constructor & Destructor Documentation

5.53.2.1 UnsteadyAdsorption::UnsteadyAdsorption()

Default Constructor.

5.53.2.2 UnsteadyAdsorption::~UnsteadyAdsorption()

Default Destructor.

5.53.3 Member Function Documentation

5.53.3.1 void UnsteadyAdsorption::Initialize_Object (MasterSpeciesList & List, int n)

Function to call the initialization of objects sequentially.

5.53.3.2 void UnsteadyAdsorption::Display_Info()

Display the adsorption reaction information (PLACE HOLDER)

5.53.3.3 void UnsteadyAdsorption::modifyDeltas (MassBalance & mbo)

Modify the Deltas in the MassBalance Object.

This function will take a mass balance object as an argument and modify the deltas in that object to correct for how adsorption affects that particular mass balance. Since adsorption can effect multiple mass balances, this function must be called for each mass balance in the system.

Parameters

mbo reference to the MassBalance Object the adsorption is acting on

5.53.3.4 int UnsteadyAdsorption::setAdsorbIndices ()

Find and set the adsorbed species indices for each reaction object.

This function searches through the Reaction objects in UnsteadyAdsorption to find the solid species and their indices to set that information in the adsorb_index structure. That information will be used later to approximate maximum capacities and equilibrium parameters for use in a modified extended Langmuir type expression. Function will return 0 if successful and -1 on a failure.

```
5.53.3.5 int UnsteadyAdsorption::checkAqueousIndices ( )
```

Function to check and report errors in the aqueous species indices.

5.53.3.6 void UnsteadyAdsorption::setActivityModelInfo (int(*)(const Matrix< double > &logq, Matrix< double > &activity, const void *data) act, const void * act data)

Function to set the surface activity model and data pointer.

This function will setup the surface activity model based on the given pointer arguments. If no arguments are given, or are given as NULL, then the activity model will default to ideal solution assumption.

```
5.53.3.7 void UnsteadyAdsorption::setAqueousIndex ( int rxn_i, int species_i )
```

Set the primary aqueous species index for the ith reaction.

```
5.53.3.8 int UnsteadyAdsorption::setAqueousIndexAuto ( )
```

Automatically sets the primary aqueous species index based on reactions.

This function will go through all species and all reactions in the adsorption object and automatically set the primary aqueous species index based on the stoicheometry of the reaction. It will also check and make sure that the primary aqueous index species appears opposite of the adsorbed species in the reactions. Note: This function assumes that the adsorbed indices have already been set.

```
5.53.3.9 void UnsteadyAdsorption::setActivityEnum ( int act )
```

Set the surface activity enum value.

```
5.53.3.10 void UnsteadyAdsorption::setMolarFactor (int rxn_i, double m)
```

Set the molar factor for the ith reaction (mol/mol)

5.53.3.11 void UnsteadyAdsorption::setVolumeFactor (int i, double v)

Set the ith volume factor for the species list (cm³/mol)

5.53.3.12 void UnsteadyAdsorption::setAreaFactor (int i, double a)

Set the ith area factor for the species list (m^2/mol)

5.53.3.13 void UnsteadyAdsorption::setSpecificArea (double a)

Set the specific area for the adsorbent (m²/kg)

5.53.3.14 void UnsteadyAdsorption::setSpecificMolality (double a)

Set the specific molality for the adsorbent (mol/kg)

```
5.53.3.15 void UnsteadyAdsorption::setSurfaceCharge ( double c )
Set the surface charge of the uncomplexed ligands.
5.53.3.16 void UnsteadyAdsorption::setTotalMass ( double m )
Set the total mass of the adsorbent (kg)
5.53.3.17 void UnsteadyAdsorption::setTotalVolume ( double v )
Set the total volume of the system (L)
5.53.3.18 void UnsteadyAdsorption::setAreaBasisBool ( bool opt )
Set the basis boolean directly.
5.53.3.19 void UnsteadyAdsorption::setSurfaceChargeBool ( bool opt )
Set the boolean for inclusion of surface charging.
5.53.3.20 void UnsteadyAdsorption::setBasis ( std::string option )
Set the basis of the adsorption problem from the given string arg.
5.53.3.21 void UnsteadyAdsorption::setAdsorbentName ( std::string name )
Set the name of the adsorbent to the given string.
5.53.3.22 void UnsteadyAdsorption::updateActivities ( )
Set the old activities as the new activities before doing next time step.
5.53.3.23 void UnsteadyAdsorption::calculateAreaFactors ( )
Calculates the area factors used from the van der Waals volumes.
5.53.3.24 void UnsteadyAdsorption::calculateEquilibria ( double T )
Calculates all equilibrium parameters as a function of temperature.
5.53.3.25 void UnsteadyAdsorption::calculateRates ( double T )
Calculates all reaction rate parameters as a function of temperature.
5.53.3.26 void UnsteadyAdsorption::setChargeDensity ( const Matrix< double > & x )
Calculates and sets the current value of charge density.
```

5.53.3.27 void UnsteadyAdsorption::setlonicStrength (const Matrix < double > & x)

Calculates and sets the current value of ionic strength.

5.53.3.28 int UnsteadyAdsorption::callSurfaceActivity (const Matrix < double > & x)

Calls the activity model and returns an int flag for success or failure.

5.53.3.29 double UnsteadyAdsorption::calculateActiveFraction (const Matrix < double > & x)

Calculates the fraction of the surface that is active and available.

5.53.3.30 double UnsteadyAdsorption::calculateSurfaceChargeDensity (const Matrix < double > & x)

Function to calculate the surface charge density based on concentrations.

This function is used to calculate the surface charge density of the adsorbed species based on the charges and concentrations of the adsorbed species. The calculation is used to correct the adsorption equilibria constant based on a localized surface charge balance. This requires that you know the molality of the uncomplexed ligand species on the surface, as well as the specific surface area for the adsorbent.

Parameters

x matrix of the log(C) concentration values at the current non-linear step

5.53.3.31 double UnsteadyAdsorption::calculatePsi (double sigma, double T, double I, double rel_epsilon)

Function calculates the Psi (electric surface potential) given a set of arguments.

This function will calculate the electric surface potential of the adsorbent under the current conditions of charge density, temperature, ionic strength, and relative permittivity.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)

5.53.3.32 double UnsteadyAdsorption::calculateAqueousChargeExchange (int i)

Function to calculate the net exchange of charges of the aqeous species involved in a given reaction.

This function will look at all aqueous species involved in the ith adsorption reaction and sum up their stoicheometries and charges to see what the net change in charge is caused by the adsorption of charged species in solution. It is then used to adjust or correct the equilibrium constant for the given adsorption reaction.

Parameters

i index of the reaction of interest for the adsorption object

5.53.3.33 double UnsteadyAdsorption::calculateEquilibriumCorrection (double *sigma*, double *T*, double *I*, double *rel_epsilon*, int *i*)

Function to calculate the correction term for the equilibrium parameter.

This function calculates the correction term that gets applied to the equilibrium parameter to correct for surface charge and charge accumulation/depletion effects. It will call the psi approximation and charge exchange functions, therefore it needs to have those functions arguments passed to it as well.

Parameters

sigma	charge density of the surface (C/m^2)
T	temperature of the system in question (K)
1	ionic strength of the medium the surface is in (mol/L)
rel_epsilon	relative permittivity of the medium (Unitless)
i	index of the reaction of interest for the adsorption object

5.53.3.34 double UnsteadyAdsorption::Eval_Residual (const Matrix< double > & x, const Matrix< double > & a double a

Calculates the residual for the ith reaction in the system.

This function will provide a system residual for the ith reaction object involved in the Adsorption Reaction. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously. This function will also adjust the equilibrium parameter for the reaction

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
i	index of the reaction of interest for the adsorption object

5.53.3.35 double UnsteadyAdsorption::Eval_Residual (const Matrix < double > & x_new, const Matrix < double > & x_old, const Matrix < double > & gama_new, const Matrix < double > & gama_old, double T, double rel_perm, int i)

Calculates the unsteady residual for the ith reaction in the system.

This function will provide a system residual for the ith reaction object involved in the Unsteady Adsorption Reaction. The residual is fed into the SHARK solver to find the solution to solid and aqueous phase concentrations simultaneously. This function will also adjust the equilibrium parameter for the reaction

Parameters

x_new	matrix of the current log(C) concentration values at the current non-linear step
gama_new	matrix of current activity coefficients for each species at the current non-linear step
x_old	matrix of the old log(C) concentration values at the current non-linear step
gama_old	matrix of old activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
i	index of the reaction of interest for the adsorption object

5.53.3.36 double UnsteadyAdsorption::Eval_ReactionRate (const Matrix < double > & x, const Matrix < double > & gama, double T, double rel_perm, int i)

Function to calculate the explicit or implicit rate of reaction.

This function will calculate the rate/extent of the unsteady adsorption reaction given the log(C) concentrations and aqueous activities, as well as temperature and permittivity. The temperature and permittivity are used to make surface charge corrections to the equilibria and rate constants.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
i	index of the reaction of interest for the adsorption object

5.53.3.37 double UnsteadyAdsorption::Eval_IC_Residual (const Matrix< double > & x, int i)

Calculate the unsteady residual for initial conditions.

Setting the intial conditions for all variables in the system requires a speciation calculation. However, we want the unsteady variables to be set to their respective initial conditions. Using this residual function imposes an equality constraint on those non-linear, unsteady variables allowing the rest of the speciation problem to be solved via PJFNK iterations.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
i	index of the reaction of interest for the adsorption object

5.53.3.38 double UnsteadyAdsorption::Explicit_Eval (const Matrix < double > & x, const Matrix < double > & gama, double T, double rel_perm, int i)

Return an approximate explicit solution to our unsteady adsorption variable (mol/kg)

This function will approximate the concentration of the unsteady variables based on an explicit time discretization. The purpose of this function is to try to provide the PJFNK method with a good initial guess for the values of the non-linear, unsteady variables. If we do not provide a good initial guess to these variables, then the PJFNK method may not converge to the correct solution, because the unsteady problem is the most difficult to solve.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step
T	temperature of the system in question (K)
rel_perm	relative permittivity of the media (unitless)
i	index of the reaction of interest for the adsorption object

```
5.53.3.39 UnsteadyReaction UnsteadyAdsorption::getReaction (int i)
Return reference to the ith reaction object in the adsorption object.
5.53.3.40 double UnsteadyAdsorption::getMolarFactor ( int i )
Get the ith reaction's molar factor for adsorption (mol/mol)
5.53.3.41 double UnsteadyAdsorption::getVolumeFactor ( int i )
Get the ith volume factor (species not involved return zeros) (cm<sup>^</sup>3/mol)
5.53.3.42 double UnsteadyAdsorption::getAreaFactor ( int i )
Get the ith area factor (species not involved return zeros) (m<sup>2</sup>/mol)
5.53.3.43 double UnsteadyAdsorption::getActivity ( int i )
Get the ith activity factor for the surface species.
5.53.3.44 double UnsteadyAdsorption::getOldActivity ( int i )
Get the ith old activity factor for the surface species.
5.53.3.45 double UnsteadyAdsorption::getSpecificArea ( )
Get the specific area of the adsorbent (m<sup>2</sup>/kg) or (mol/kg)
5.53.3.46 double UnsteadyAdsorption::getSpecificMolality ( )
Get the specific molality of the adsorbent (mol/kg)
5.53.3.47 double UnsteadyAdsorption::getSurfaceCharge ( )
Get the surface charge of the adsorbent.
5.53.3.48 double UnsteadyAdsorption::getBulkDensity ( )
Calculate and return bulk density of adsorbent in system (kg/L)
5.53.3.49 double UnsteadyAdsorption::getTotalMass ( )
Get the total mass of adsorbent in the system (kg)
5.53.3.50 double UnsteadyAdsorption::getTotalVolume ( )
Get the total volume of the system (L)
```

```
5.53.3.51 double UnsteadyAdsorption::getChargeDensity ( )
Get the value of the surface charge density (C/m<sup>2</sup>)
5.53.3.52 double UnsteadyAdsorption::getlonicStrength ( )
Get the value of the ionic strength of solution (mol/L)
5.53.3.53 int UnsteadyAdsorption::getNumberRxns ( )
Get the number of reactions involved in the adsorption object.
5.53.3.54 int UnsteadyAdsorption::getAdsorbIndex (int i)
Get the index of the adsorbed species in the ith reaction.
5.53.3.55 int UnsteadyAdsorption::getAqueousIndex ( int i )
Get the index of the primary aqueous species in the ith reaction.
5.53.3.56 int UnsteadyAdsorption::getActivityEnum ( )
Return the enum representing the choosen activity function.
5.53.3.57 bool UnsteadyAdsorption::isAreaBasis ( )
Returns true if we are in the Area Basis, False if in Molar Basis.
5.53.3.58 bool UnsteadyAdsorption::includeSurfaceCharge ( )
Returns true if we are considering surface charging during adsorption.
5.53.3.59 std::string UnsteadyAdsorption::getAdsorbentName ( )
Returns the name of the adsorbent as a string.
5.53.4 Member Data Documentation
5.53.4.1 Matrix<double> UnsteadyAdsorption::activities_old [protected]
List of the old activities calculated by the activity model.
5.53.4.2 std::vector<UnsteadyReaction>UnsteadyAdsorption::ads_rxn [private]
List of reactions involved with adsorption.
The documentation for this class was generated from the following file:
```

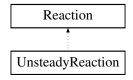
shark.h

5.54 UnsteadyReaction Class Reference

Unsteady Reaction Object (inherits from Reaction)

#include <shark.h>

Inheritance diagram for UnsteadyReaction:



Public Member Functions

• UnsteadyReaction ()

Default Constructor.

∼UnsteadyReaction ()

Default Destructor.

· void Initialize_Object (MasterSpeciesList &List)

Function to initialize the UnsteadyReaction object from the MasterSpeciesList.

void Display_Info ()

Display the unsteady reaction information.

void Set_Species_Index (int i)

Set the Unsteady species index by number.

void Set_Species_Index (std::string formula)

Set the Unsteady species index by formula.

• void Set_Stoichiometric (int i, double v)

Set the ith stoichiometric value (see Reaction object)

void Set_Equilibrium (double v)

Set the equilibrium constant (logK) (see Reaction object)

void Set_Enthalpy (double H)

Set the enthalpy of the reaction (J/mol) (see Reaction object)

void Set_Entropy (double S)

Set the entropy of the reaction (J/K/mol) (see Reaction object)

void Set_EnthalpyANDEntropy (double H, double S)

Set both the enthalpy and entropy (J/mol) & (J/K/mol) (see Reaction object)

void Set_Energy (double G)

Set the Gibb's free energy of reaction (J/mol) (see Reaction object)

void Set_InitialValue (double ic)

Set the initial value of the unsteady variable.

void Set_MaximumValue (double max)

Set the maximum value of the unsteady variable to a given value max (mol/L)

void Set_Forward (double forward)

Set the forward rate for the reaction (mol/L/hr)

void Set_Reverse (double reverse)

Set the reverse rate for the reaction (mol/L/hr)

· void Set ForwardRef (double Fref)

Set the forward reference rate (mol/L/hr)

void Set_ReverseRef (double Rref)

Set the reverse reference rate (mol/L/hr)

void Set_ActivationEnergy (double E)

Set the activation energy for the reaction (J/mol)

void Set Affinity (double b)

Set the temperature affinity parameter for the reaction.

void Set TimeStep (double dt)

Set the time step for the current simulation.

void checkSpeciesEnergies ()

Function to check MasterSpeciesList for species energy info (see Reaction object)

void calculateEnergies ()

Function to calculate the energy of the reaction (see Reaction object)

void calculateEquilibrium (double T)

Function to calculate the equilibrium constant (see Reaction object)

• void calculateRate (double T)

Function to calculate the rate constant based on given temperature.

bool haveEquilibrium ()

True if equilibrium constant is given or can be calculated (see Reaction object)

• bool haveRate ()

Function to return true if you have the forward or reverse rate calculated.

bool haveForwardRef ()

Function to return true if you have the forward reference rate.

• bool haveReverseRef ()

Function to return true if you have the reverse reference rate.

bool haveForward ()

Function to return true if you have the forward rate.

• bool haveReverse ()

Function to return true if you have the reverse rate.

int Get_Species_Index ()

Fetch the index of the Unsteady species.

• double Get Stoichiometric (int i)

Fetch the ith stoichiometric value.

double Get_Equilibrium ()

Fetch the equilibrium constant (logK)

double Get_Enthalpy ()

Fetch the enthalpy of the reaction.

• double Get_Entropy ()

Fetch the entropy of the reaction.

double Get_Energy ()

Fetch the energy of the reaction.

double Get InitialValue ()

Fetch the initial value of the variable.

• double Get MaximumValue ()

Fetch the maximum value of the variable.

double Get_Forward ()

Fetch the forward rate.

• double Get_Reverse ()

Fetch the reverse rate.

double Get_ForwardRef ()

Fetch the forward reference rate.

double Get_ReverseRef ()

Fetch the reverse reference rate.

double Get_ActivationEnergy ()

Fetch the activation energy for the reaction.

double Get_Affinity ()

Fetch the temperature affinity for the reaction.

double Get_TimeStep ()

Fetch the time step.

double Eval_ReactionRate (const Matrix< double > &x, const Matrix< double > &gama)

Calculate reation rate (dC/dt) from concentrations and activities.

double Eval_Residual (const Matrix< double > &x_new, const Matrix< double > &x_old, const Matrix
 double > &gama_new, const Matrix< double > &gama_old)

Calculate the unsteady residual for the reaction using and implicit time discretization.

• double Eval_Residual (const Matrix< double > &x, const Matrix< double > &gama)

Calculate the steady-state residual for this reaction (see Reaction object)

double Eval IC Residual (const Matrix< double > &x)

Calculate the unsteady residual for initial conditions.

double Explicit Eval (const Matrix < double > &x, const Matrix < double > &gama)

Return an approximate explicit solution to our unsteady variable (mol/L)

Protected Attributes

· double initial_value

Initial value given at t=0 (in mol/L)

double max_value

Maximum value plausible (in mol/L)

· double forward rate

Forward reaction rate constant (in (mol/L)^n/hr)

· double reverse_rate

Reverse reaction rate constant (in (mol/L)^n/hr)

double forward_ref_rate

Forward reference rate constant (in (mol/L)^n/hr)

double reverse_ref_rate

Reverse reference rate constant (in (mol/L)^n/hr)

· double activation_energy

Activation or barrier energy for the reaction (J/mol)

· double temperature_affinity

Temperature affinity parameter (dimensionless)

· double time_step

Time step size for current step.

bool HaveForward

True if can calculate, or was given the forward rate.

bool HaveReverse

True if can calculate, or was given the reverse rate.

· bool HaveForRef

True if given the forward reference rate.

bool HaveRevRef

True if given the reverse reference rate.

• int species_index

Index in MasterList of Unsteady Species.

Additional Inherited Members

5.54.1 Detailed Description

Unsteady Reaction Object (inherits from Reaction)

C++ style object that holds data and functions associated with unsteady chemical reactions...

```
i.e., aA + bB < -reverse: forward -> cC + dD
```

This is essentially the same as the steady reaction, but we now have a forward and reverse reaction rate to deal with. It should be noted that this is a very simple kinetic reaction model based on splitting an overall equilibrium reaction into an overall forward and reverse reaction model. Therefore, it is not expected that this representation of the reaction will provide high accuracy results for reaction kinetics, but should at least provide an overall idea of the process occurring.

```
5.54.2 Constructor & Destructor Documentation
```

```
5.54.2.1 UnsteadyReaction::UnsteadyReaction()
```

Default Constructor.

```
5.54.2.2 UnsteadyReaction:: ~UnsteadyReaction ( )
```

Default Destructor.

5.54.3 Member Function Documentation

5.54.3.1 void UnsteadyReaction::Initialize_Object (MasterSpeciesList & List)

Function to initialize the UnsteadyReaction object from the MasterSpeciesList.

```
5.54.3.2 void UnsteadyReaction::Display_Info ( )
```

Display the unsteady reaction information.

```
5.54.3.3 void UnsteadyReaction::Set_Species_Index ( int i )
```

Set the Unsteady species index by number.

This function will set the unsteady species index by the index i given. That given index must correspond to the index of the species in the MasterSpeciesList that is being considered as the unsteady species.

Parameters

```
i index of the unsteady species in the MasterSpeciesList
```

```
5.54.3.4 void UnsteadyReaction::Set_Species_Index ( std::string formula )
```

Set the Unsteady species index by formula.

This function will check the MasterSpeciesList for the molecule object that has the given formula, then set the unsteady species index based on the index of that species in the master list.

Parameters

formula | molecular formula of the unsteady species (see mola.h for standard formatting)

5.54.3.5 void UnsteadyReaction::Set_Stoichiometric (int i, double v)

Set the ith stoichiometric value (see Reaction object)

5.54.3.6 void UnsteadyReaction::Set_Equilibrium (double v)

Set the equilibrium constant (logK) (see Reaction object)

5.54.3.7 void UnsteadyReaction::Set_Enthalpy (double H)

Set the enthalpy of the reaction (J/mol) (see Reaction object)

5.54.3.8 void UnsteadyReaction::Set_Entropy (double S)

Set the entropy of the reaction (J/K/mol) (see Reaction object)

5.54.3.9 void UnsteadyReaction::Set_EnthalpyANDEntropy (double *H*, double *S*)

Set both the enthalpy and entropy (J/mol) & (J/K/mol) (see Reaction object)

5.54.3.10 void UnsteadyReaction::Set_Energy (double G)

Set the Gibb's free energy of reaction (J/mol) (see Reaction object)

5.54.3.11 void UnsteadyReaction::Set_InitialValue (double ic)

Set the initial value of the unsteady variable.

This function sets the initial concentration value for the unsteady species to the given value ic (mol/L). Only unsteady species need to be given an initial value. All other species initial values for the overall system is setup based on a speciation calculation performed while holding the unsteady variables constant at their respective initial values.

Parameters

ic initial concentration value for the unsteady object (mol/L)

5.54.3.12 void UnsteadyReaction::Set_MaximumValue (double max)

Set the maximum value of the unsteady variable to a given value max (mol/L)

This function will be called internally to help bound the unsteady variable to reasonable maximum values. That maximum is usually based on the mass balances for the current non-linear iteration.

Parameters

max maximum allowable value for the unsteady variable (mo	L)
---	----

5.54.3.13 void UnsteadyReaction::Set_Forward (double forward)

Set the forward rate for the reaction (mol/L/hr)

5.54.3.14 void UnsteadyReaction::Set_Reverse (double reverse)

Set the reverse rate for the reaction (mol/L/hr)

5.54.3.15 void UnsteadyReaction::Set_ForwardRef (double Fref)

Set the forward reference rate (mol/L/hr)

Unlike just setting the forward rate, this function sets a reference forward rate of the reaction that can be used to correct the overall forward rate based on system temperature and Arrhenius Rate Equation constants.

Parameters

5.54.3.16 void UnsteadyReaction::Set_ReverseRef (double Rref)

Set the reverse reference rate (mol/L/hr)

Unlike just setting the reverse rate, this function sets a reference reverse rate of the reaction that can be used to correct the overall reverse rate based on system temperature and Arrhenius Rate Equation constants.

Parameters

Rref reverse reference rate constant (mol/L/hr)

5.54.3.17 void UnsteadyReaction::Set_ActivationEnergy (double E)

Set the activation energy for the reaction (J/mol)

This function will set the activation energy for the reaction to the given value of E. Note that we will only set one value for activation energy, even though there are rates for forward and reverse reactions. This is because we use the ratio of the rates and the equilibrium constant to establish the other rate. Therefore, we only need either the forward or reverse rate and the equilibrium constant to set all the rates.

Parameters

E activation energy for the forward or reverse rate, depending on which was given

```
5.54.3.18 void UnsteadyReaction::Set_Affinity ( double b )
```

Set the temperature affinity parameter for the reaction.

This function will set the temperature affinity for the reaction to the given value of b. Note that we will only set one value for temperature affinity, even though there are rates for forward and reverse reactions. This is because we use the ratio of the rates and the equilibrium constant to establish the other rate. Therefore, we only need either the forward or reverse rate and the equilibrium constant to set all the rates.

Parameters

b temperature affinity for the forward or reverse rate, depending on which was given

```
5.54.3.19 void UnsteadyReaction::Set_TimeStep ( double dt )
```

Set the time step for the current simulation.

```
5.54.3.20 void UnsteadyReaction::checkSpeciesEnergies ( )
```

Function to check MasterSpeciesList for species energy info (see Reaction object)

```
5.54.3.21 void UnsteadyReaction::calculateEnergies ( )
```

Function to calculate the energy of the reaction (see Reaction object)

5.54.3.22 void UnsteadyReaction::calculateEquilibrium (double *T*)

Function to calculate the equilibrium constant (see Reaction object)

```
5.54.3.23 void UnsteadyReaction::calculateRate ( double T )
```

Function to calculate the rate constant based on given temperature.

This function will calculate and set either the forward or reverse rate for the unsteady reaction based on what information was given. If the forward rate information was given, then it sets the reverse rate and visa versa. If nothing was set correctly, an error will occur.

Parameters

T | temperature of the system in Kelvin

5.54.3.24 bool UnsteadyReaction::haveEquilibrium ()

True if equilibrium constant is given or can be calculated (see Reaction object)

5.54.3.25 bool UnsteadyReaction::haveRate ()

Function to return true if you have the forward or reverse rate calculated.

```
5.54.3.26 bool UnsteadyReaction::haveForwardRef()
Function to return true if you have the forward reference rate.
5.54.3.27 bool UnsteadyReaction::haveReverseRef ( )
Function to return true if you have the reverse reference rate.
5.54.3.28 bool UnsteadyReaction::haveForward ( )
Function to return true if you have the forward rate.
5.54.3.29 bool UnsteadyReaction::haveReverse ( )
Function to return true if you have the reverse rate.
5.54.3.30 int UnsteadyReaction::Get_Species_Index ( )
Fetch the index of the Unsteady species.
5.54.3.31 double UnsteadyReaction::Get_Stoichiometric ( int i )
Fetch the ith stoichiometric value.
5.54.3.32 double UnsteadyReaction::Get_Equilibrium ( )
Fetch the equilibrium constant (logK)
5.54.3.33 double UnsteadyReaction::Get_Enthalpy ( )
Fetch the enthalpy of the reaction.
5.54.3.34 double UnsteadyReaction::Get_Entropy ( )
Fetch the entropy of the reaction.
5.54.3.35 double UnsteadyReaction::Get_Energy ( )
Fetch the energy of the reaction.
5.54.3.36 double UnsteadyReaction::Get_InitialValue ( )
Fetch the initial value of the variable.
5.54.3.37 double UnsteadyReaction::Get_MaximumValue ( )
Fetch the maximum value of the variable.
```

```
5.54.3.38 double UnsteadyReaction::Get_Forward ( )
Fetch the forward rate.
5.54.3.39 double UnsteadyReaction::Get_Reverse ( )
Fetch the reverse rate.
5.54.3.40 double UnsteadyReaction::Get_ForwardRef()
Fetch the forward reference rate.
5.54.3.41 double UnsteadyReaction::Get_ReverseRef ( )
Fetch the reverse reference rate.
5.54.3.42 double UnsteadyReaction::Get_ActivationEnergy ( )
Fetch the activation energy for the reaction.
5.54.3.43 double UnsteadyReaction::Get_Affinity ( )
Fetch the temperature affinity for the reaction.
5.54.3.44 double UnsteadyReaction::Get_TimeStep ( )
Fetch the time step.
5.54.3.45 double UnsteadyReaction::Eval_ReactionRate ( const Matrix < double > & x, const Matrix < double > & gama )
```

Calculate reation rate (dC/dt) from concentrations and activities.

This function calculates the right hand side of the unsteady reaction equation based on the available rates, the

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step

current values of the non-linear variables (x=log(C)), and the activity coefficients (gama).

5.54.3.46 double UnsteadyReaction::Eval_Residual (const Matrix< double > & x_new, const Matrix< double > & x_old, const Matrix< double > & gama_new, const Matrix< double > & gama_old)

Calculate the unsteady residual for the reaction using and implicit time discretization.

This function uses the current time step and states of the non-linear variables and activities to form the residual contribution of the unsteady reaction. The time dependent functions are discretized using an implicit finite difference for best stability.

Parameters

x_new	matrix of the log(C) concentration values at the current non-linear step	
gama_new	matrix of activity coefficients for each species at the current non-linear step	
x_old	matrix of the log(C) concentration values at the previous non-linear step	
gama_old	matrix of activity coefficients for each species at the previous non-linear step	

5.54.3.47 double UnsteadyReaction::Eval_Residual (const Matrix < double > & x, const Matrix < double > & gama)

Calculate the steady-state residual for this reaction (see Reaction object)

5.54.3.48 double UnsteadyReaction::Eval_IC_Residual (const Matrix < double > & x)

Calculate the unsteady residual for initial conditions.

Setting the intial conditions for all variables in the system requires a speciation calculation. However, we want the unsteady variables to be set to their respective initial conditions. Using this residual function imposes an equality constraint on those non-linear, unsteady variables allowing the rest of the speciation problem to be solved via PJFNK iterations.

Parameters

x matrix of the log(C) concentration values at the current non-linear step

 $\textbf{5.54.3.49} \quad \textbf{double UnsteadyReaction::Explicit_Eval (const \ \textbf{Matrix} < \textbf{double} > \& \ \textbf{\textit{x}}, \ \textbf{const} \ \textbf{Matrix} < \textbf{double} > \& \ \textbf{\textit{gama}} \)$

Return an approximate explicit solution to our unsteady variable (mol/L)

This function will approximate the concentration of the unsteady variables based on an explicit time discretization. The purpose of this function is to try to provide the PJFNK method with a good initial guess for the values of the non-linear, unsteady variables. If we do not provide a good initial guess to these variables, then the PJFNK method may not converge to the correct solution, because the unsteady problem is the most difficult to solve.

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
gama	matrix of activity coefficients for each species at the current non-linear step

5.54.4 Member Data Documentation

5.54.4.1 double UnsteadyReaction::initial_value [protected]

Initial value given at t=0 (in mol/L)

5.54.4.2 double UnsteadyReaction::max_value [protected]

Maximum value plausible (in mol/L)

```
5.54.4.3 double UnsteadyReaction::forward_rate [protected]
Forward reaction rate constant (in (mol/L)^n/hr)
5.54.4.4 double UnsteadyReaction::reverse_rate [protected]
Reverse reaction rate constant (in (mol/L)^n/hr)
5.54.4.5 double UnsteadyReaction::forward_ref_rate [protected]
Forward reference rate constant (in (mol/L)^n/hr)
5.54.4.6 double UnsteadyReaction::reverse_ref_rate [protected]
Reverse reference rate constant (in (mol/L)^n/hr)
5.54.4.7 double UnsteadyReaction::activation_energy [protected]
Activation or barrier energy for the reaction (J/mol)
5.54.4.8 double UnsteadyReaction::temperature_affinity [protected]
Temperature affinity parameter (dimensionless)
5.54.4.9 double UnsteadyReaction::time_step [protected]
Time step size for current step.
5.54.4.10 bool UnsteadyReaction::HaveForward [protected]
True if can calculate, or was given the forward rate.
5.54.4.11 bool UnsteadyReaction::HaveReverse [protected]
True if can calculate, or was given the reverse rate.
5.54.4.12 bool UnsteadyReaction::HaveForRef [protected]
True if given the forward reference rate.
5.54.4.13 bool UnsteadyReaction::HaveRevRef [protected]
True if given the reverse reference rate.
5.54.4.14 int UnsteadyReaction::species_index [protected]
Index in MasterList of Unsteady Species.
The documentation for this class was generated from the following file:
```

• shark.h

5.55 ValueTypePair Class Reference

Value-Type Pair object to recognize data type of a string that was read.

```
#include <yaml_wrapper.h>
```

Public Member Functions

• ValueTypePair ()

Default constructor.

∼ValueTypePair ()

Default destructor.

ValueTypePair (const std::pair< std::string, int > &vt)

Constructor by pair.

• ValueTypePair (std::string value, int type)

Construction by string and int.

ValueTypePair (const ValueTypePair &vt)

Copy constructor.

ValueTypePair & operator= (const ValueTypePair &vt)

Equals operator overload.

void editValue (std::string value)

Edits value to pair with UNKOWN type.

void editPair (std::string value, int type)

Creates a paired Value-Type from the given args.

void findType ()

Determines the data type of the object.

void assertType (int type)

Forces a specific data type.

• void DisplayPair ()

Display the pair information.

• std::string getString ()

Returns the value of the pair as a string.

bool getBool ()

Returns the value of the pair as a bool.

double getDouble ()

Returns the value of the pair as a double.

• int getInt ()

Returns the value of the pair as an int.

• std::string getValue ()

Returns the value of the pair as it was given.

int getType ()

Returns the type of the pair.

std::pair< std::string, int > & getPair ()

Returns reference to the actual pair object.

Private Attributes

std::pair < std::string, int > Value_Type
 pair object holding the Value and Type info

int type

Type of the value.

5.55.1 Detailed Description

Value-Type Pair object to recognize data type of a string that was read.

C++ Object that creates a pair between a read in value as a string and an enum denoting what the data type of that string is. This object is primarily used in the other yaml_wrapper objects, but can also be used for any string that you want to parse to identify it's type. The supported types are denoted in the data_type enum and can be determined automatically by the findType() function or can be specified by the assertType() function.

```
5.55.2 Constructor & Destructor Documentation
5.55.2.1 ValueTypePair::ValueTypePair()
Default constructor.
5.55.2.2 ValueTypePair::~ValueTypePair()
Default destructor.
5.55.2.3 ValueTypePair::ValueTypePair ( const std::pair < std::string, int > & vt )
Constructor by pair.
5.55.2.4 ValueTypePair::ValueTypePair ( std::string value, int type )
Construction by string and int.
5.55.2.5 ValueTypePair::ValueTypePair ( const ValueTypePair & vt )
Copy constructor.
5.55.3 Member Function Documentation
5.55.3.1 ValueTypePair& ValueTypePair::operator= ( const ValueTypePair & vt )
Equals operator overload.
5.55.3.2 void ValueTypePair::editValue ( std::string value )
Edits value to pair with UNKOWN type.
5.55.3.3 void ValueTypePair::editPair ( std::string value, int type )
Creates a paired Value-Type from the given args.
5.55.3.4 void ValueTypePair::findType()
```

Determines the data type of the object.

```
5.55.3.5 void ValueTypePair::assertType (int type)
Forces a specific data type.
5.55.3.6 void ValueTypePair::DisplayPair ( )
Display the pair information.
5.55.3.7 std::string ValueTypePair::getString ( )
Returns the value of the pair as a string.
5.55.3.8 bool ValueTypePair::getBool ( )
Returns the value of the pair as a bool.
5.55.3.9 double ValueTypePair::getDouble ( )
Returns the value of the pair as a double.
5.55.3.10 int ValueTypePair::getInt()
Returns the value of the pair as an int.
5.55.3.11 std::string ValueTypePair::getValue ( )
Returns the value of the pair as it was given.
5.55.3.12 int ValueTypePair::getType ( )
Returns the type of the pair.
5.55.3.13 std::pair<std::string,int>& ValueTypePair::getPair()
Returns reference to the actual pair object.
5.55.4 Member Data Documentation
5.55.4.1 std::pair<std::string,int> ValueTypePair::Value_Type [private]
pair object holding the Value and Type info
5.55.4.2 int ValueTypePair::type [private]
Type of the value.
The documentation for this class was generated from the following file:
```

Generated by Doxygen

· yaml_wrapper.h

5.56 yaml_cpp_class Class Reference

Primary object used when reading and digitally storing yaml files.

```
#include <yaml_wrapper.h>
```

Public Member Functions

• yaml_cpp_class ()

Default constructor.

~yaml_cpp_class ()

Default destructor.

• int setInputFile (const char *file)

Set the input file to be read.

• int readInputFile ()

Reads through input file and stores into YamlWrapper.

• int cleanup ()

Deletes yaml_c objects and closes the input file.

int executeYamlRead (const char *file)

Runs the full execution of initialization, reading, and cleaning.

YamlWrapper & getYamlWrapper ()

Returns reference to the YamlWrapper Object.

void DisplayContents ()

Print out the contents of the read to the console window.

Private Attributes

• YamlWrapper yaml_wrapper

YamlWrapper object where digital file is stored.

FILE * input_file

Function pointer to the yaml formatted file.

• const char * file_name

Name of the file to be parsed and read.

• yaml_parser_t token_parser

C-YAML parser object for token based parsing.

yaml_token_t current_token

C-YAML token object for the current token in the file.

yaml_token_t previous_token

C-YAML token object for the previous token in the file.

5.56.1 Detailed Description

Primary object used when reading and digitally storing yaml files.

C++ Object that holds the YamlWrapper object and the C-YAML objects necessary for reading and parsing a yaml formatted file. This is the primary object that users are expected to work with when using yaml_wrapper.h to read input files. It contains functions necessary to setup a read instance, read and parse the input, place the parsed input results into the digital YamlWrapper object, and allow the user to query that object.

The two main functions that the typical user will need are: (i) executeYamlRead() and (ii) getYamlWrapper. Make sure that the read function was called prior to querying the YamlWrapper structure. Do not call the cleanup() function if using executeYamlRead(). That function will be automattically called after the read is complete.

```
5.56.2 Constructor & Destructor Documentation
5.56.2.1 yaml_cpp_class::yaml_cpp_class()
Default constructor.
5.56.2.2 yaml_cpp_class::~yaml_cpp_class()
Default destructor.
5.56.3 Member Function Documentation
5.56.3.1 int yaml_cpp_class::setInputFile ( const char * file )
Set the input file to be read.
5.56.3.2 int yaml_cpp_class::readInputFile()
Reads through input file and stores into YamlWrapper.
5.56.3.3 int yaml_cpp_class::cleanup()
Deletes yaml_c objects and closes the input file.
5.56.3.4 int yaml_cpp_class::executeYamlRead ( const char * file )
Runs the full execution of initialization, reading, and cleaning.
5.56.3.5 YamlWrapper& yaml_cpp_class::getYamlWrapper( )
Returns reference to the YamlWrapper Object.
5.56.3.6 void yaml_cpp_class::DisplayContents ( )
Print out the contents of the read to the console window.
5.56.4 Member Data Documentation
5.56.4.1 YamlWrapper yaml_cpp_class::yaml_wrapper [private]
YamlWrapper object where digital file is stored.
5.56.4.2 FILE* yaml_cpp_class::input_file [private]
Function pointer to the yaml formatted file.
```

```
5.56.4.3 const char* yaml_cpp_class::file_name [private]
Name of the file to be parsed and read.
5.56.4.4 yaml_parser_t yaml_cpp_class::token_parser [private]
C-YAML parser object for token based parsing.
5.56.4.5 yaml_token_t yaml_cpp_class::current_token [private]
C-YAML token object for the current token in the file.
5.56.4.6 yaml_token_t yaml_cpp_class::previous_token [private]
```

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

C-YAML token object for the previous token in the file.

• yaml_wrapper.h

5.57 YamlWrapper Class Reference

Object for the entire yaml file holding all documents, header, sub-headers, keys, and values.

```
#include <yaml_wrapper.h>
```

Public Member Functions

• YamlWrapper ()

Default constructor.

∼YamlWrapper ()

Default destructor.

YamlWrapper (const YamlWrapper &yaml)

Copy constructor.

YamlWrapper (std::string key, const Document &doc)

Constructor by a single document.

YamlWrapper & operator= (const YamlWrapper &yaml)

Equals overload.

• Document & operator() (const std::string key)

Return the Document reference at the given key.

• Document operator() (const std::string key) const

Return the Document at the given key.

- std::map< std::string, Document > & getDocMap ()

Return reference to the document map.

Document & getDocument (std::string key)

Return reference to the document at the key.

• std::map< std::string, Document >::const_iterator_end () const

Returns a const iterator pointing to the end of the list.

std::map< std::string, Document >::iterator end ()

Returns an iterator pointing to the end of the list.

• std::map< std::string, Document >::const_iterator begin () const

Returns a const iterator pointing to the begining of the list.

• std::map< std::string, Document >::iterator begin ()

Returns an iterator pointing to the begining of the list.

• void clear ()

Clear out the yaml object.

void resetKeys ()

Resets all the keys in DocumentMap to match document names.

void changeKey (std::string oldKey, std::string newKey)

Change a given oldKey in the map to the newKey given.

· void revalidateAllKeys ()

Resets and validates all keys in the structure.

void DisplayContents ()

Display the contents of the wrapper.

void addDocKey (std::string key)

Add a key to the document map.

void copyAnchor2Alias (std::string alias, Document &ref)

Find the anchor in the map, and copy to the Document reference given.

• int size ()

Return the size of the document map.

Document & getAnchoredDoc (std::string alias)

Return the reference to the document that is anchored with the given alias.

Document & getDocFromHeadAlias (std::string alias)

Return reference to the document that contains the header with the given alias.

Document & getDocFromSubAlias (std::string alias)

Return reference to the document that contains the subheader with the given alias.

Private Attributes

std::map< std::string, Document > Doc Map

Map of the documents contained within the wrapper.

5.57.1 Detailed Description

Object for the entire yaml file holding all documents, header, sub-headers, keys, and values.

C++ Object for the yaml file. This object holds a map of all Documents in the yaml file. Each document holds a map of Key-values and Headers. The Headers hold maps of Key-values and SubHeaders, and each SubHeader can hold more Key-values.

This object is used to represent a digital and queryable structure for all information contained within a yaml file. There are some limitations to what can be held here, and those limitations are based on the limitations in the C-YAML Library token parser. The main limitation is that the deepest level of allowable recursion in the file is Sub Header. Meaning that you are not allowed to have Sub-SubHeaders underneath a SubHeader object. This imposes a hard limit to number of nested lists that can be in a single Document object.

When using yaml_cpp_class, this object will generally be what you query into to get the information from your yaml input files. From this object, functions and operators are provided to give you the capability of querying down into any allowable level of the file by the keys that are were used in the file. Be sure that you are querying the correct objects by the correct keys, otherwise errors and exceptions will be thrown.

```
5.57.2 Constructor & Destructor Documentation
5.57.2.1 YamlWrapper::YamlWrapper()
Default constructor.
5.57.2.2 YamlWrapper::~YamlWrapper()
Default destructor.
5.57.2.3 YamlWrapper::YamlWrapper ( const YamlWrapper & yaml )
Copy constructor.
5.57.2.4 YamlWrapper::YamlWrapper ( std::string key, const Document & doc )
Constructor by a single document.
5.57.3 Member Function Documentation
5.57.3.1 YamlWrapper& YamlWrapper::operator= ( const YamlWrapper & yaml )
Equals overload.
5.57.3.2 Document& YamlWrapper::operator() ( const std::string key )
Return the Document reference at the given key.
5.57.3.3 Document YamlWrapper::operator() ( const std::string key ) const
Return the Document at the given key.
5.57.3.4 std::map<std::string, Document>& YamlWrapper::getDocMap()
Return reference to the document map.
5.57.3.5 Document & YamlWrapper::getDocument ( std::string key )
Return reference to the document at the key.
5.57.3.6 std::map<std::string, Document>::const_iterator YamlWrapper::end ( ) const
Returns a const iterator pointing to the end of the list.
5.57.3.7 std::map<std::string, Document>::iterator YamlWrapper::end ( )
Returns an iterator pointing to the end of the list.
```

```
5.57.3.8 std::map<std::string, Document>::const_iterator YamlWrapper::begin ( ) const
Returns a const iterator pointing to the begining of the list.
5.57.3.9 std::map<std::string, Document>::iterator YamlWrapper::begin ( )
Returns an iterator pointing to the begining of the list.
5.57.3.10 void YamlWrapper::clear ( )
Clear out the yaml object.
5.57.3.11 void YamlWrapper::resetKeys ( )
Resets all the keys in DocumentMap to match document names.
5.57.3.12 void YamlWrapper::changeKey ( std::string oldKey, std::string newKey )
Change a given oldKey in the map to the newKey given.
5.57.3.13 void YamlWrapper::revalidateAllKeys ( )
Resets and validates all keys in the structure.
5.57.3.14 void YamlWrapper::DisplayContents ( )
Display the contents of the wrapper.
5.57.3.15 void YamlWrapper::addDocKey ( std::string key )
Add a key to the document map.
5.57.3.16 void YamlWrapper::copyAnchor2Alias ( std::string alias, Document & ref )
Find the anchor in the map, and copy to the Document reference given.
5.57.3.17 int YamlWrapper::size ( )
Return the size of the document map.
5.57.3.18 Document& YamlWrapper::getAnchoredDoc ( std::string alias )
Return the reference to the document that is anchored with the given alias.
5.57.3.19 Document& YamlWrapper::getDocFromHeadAlias ( std::string alias )
Return reference to the document that contains the header with the given alias.
```

```
5.57.3.20 Document& YamlWrapper::getDocFromSubAlias ( std::string alias )
```

Return reference to the document that contains the subheader with the given alias.

5.57.4 Member Data Documentation

```
5.57.4.1 std::map<std::string, Document> YamlWrapper::Doc_Map [private]
```

Map of the documents contained within the wrapper.

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

· yaml_wrapper.h

6 File Documentation

6.1 dogfish.h File Reference

Diffusion Object Governing Fiber Interior Sorption History.

```
#include "finch.h"
#include "mola.h"
```

Classes

• struct DOGFISH PARAM

Data structure for species-specific parameters.

struct DOGFISH DATA

Primary data structure for running the DOGFISH application.

Functions

• void print2file_species_header (FILE *Output, DOGFISH_DATA *dog_dat, int i)

Function to print a species based header for the output file.

void print2file_DOGFISH_header (DOGFISH_DATA *dog_dat)

Function to print a time and space header for the output file.

void print2file DOGFISH result old (DOGFISH DATA *dog dat)

Function to print out the old time results for the output file.

void print2file_DOGFISH_result_new (DOGFISH_DATA *dog_dat)

Function to print out the new time results for the output file.

• double default_Retardation (int i, int I, const void *data)

Default function for the retardation coefficient.

double default_IntraDiffusion (int i, int I, const void *data)

Default function for the intraparticle diffusion coefficient.

• double default_FilmMTCoeff (int i, const void *data)

Default function for the film mass transfer coefficient.

double default_SurfaceConcentration (int i, const void *data)

Default function for the fiber surface concentration.

int setup_DOGFISH_DATA (FILE *file, double(*eval_R)(int i, int I, const void *user_data), double(*eval_\infty)
 DI)(int i, int I, const void *user_data), double(*eval_kf)(int i, const void *user_data), double(*eval_qs)(int i, const void *user_data), const void *user_data), const void *user_data)

Function will set up the memory and pointers for use in the DOGFISH simulations.

int DOGFISH_Executioner (DOGFISH_DATA *dog_dat)

Function to serially call all other functions need to solve the system at one time step.

int set DOGFISH ICs (DOGFISH DATA *dog dat)

Function called to evaluate the initial conditions for the time dependent problem.

• int set_DOGFISH_timestep (DOGFISH_DATA *dog_dat)

Function sets the time step size for the next step forward in the simulation.

int DOGFISH_preprocesses (DOGFISH_DATA *dog_dat)

Function to perform preprocess actions to be used before calling any solver.

int set_DOGFISH_params (const void *user_data)

Function to calculate the values of all parameters for all species at all nodes.

int DOGFISH_postprocesses (DOGFISH_DATA *dog_dat)

Function to perform post-solve actions such as printing out results.

int DOGFISH_reset (DOGFISH_DATA *dog_dat)

Function to reset the matrices and vectors and prepare for next time step.

int DOGFISH (DOGFISH_DATA *dog_dat)

Function performs all necessary steps to step the diffusion simulation through time.

• int DOGFISH TESTS ()

Running DOGFISH tests.

6.1.1 Detailed Description

Diffusion Object Governing Fiber Interior Sorption History.

dogfish.cpp

This set of objects and functions is used to numerically solve linear or non-linear diffusion physics of aqueous ions into cylindrical adsorbent fibers. Boundary conditions for this problem could be a film mass transfer, reaction, or dirichlet condition depending on the type of problem being solve.

Warning

Functions and methods in this file are still under construction.

Author

Austin Ladshaw

Date

04/09/2015

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6.1.2 Function Documentation

6.1.2.1 void print2file_species_header (FILE * Output, DOGFISH_DATA * dog_dat, int i)

Function to print a species based header for the output file.

6.1.2.2 void print2file_DOGFISH_header (DOGFISH_DATA * dog_dat)

Function to print a time and space header for the output file.

6.1.2.3 void print2file_DOGFISH_result_old (DOGFISH_DATA * dog_dat)

Function to print out the old time results for the output file.

6.1.2.4 void print2file_DOGFISH_result_new (DOGFISH_DATA * dog_dat)

Function to print out the new time results for the output file.

6.1.2.5 double default_Retardation (int i, int I, const void * data)

Default function for the retardation coefficient.

The default retardation coefficient for this problem is 1.0 for all time and space. Therefore, this function will only ever return a 1.

Parameters

i	index for the ith adsorbing species
1	index for the lth node in the domain
data	pointer to the DOGFISH_DATA structure

6.1.2.6 double default_IntraDiffusion (int i, int l, const void * data)

Default function for the intraparticle diffusion coefficient.

The default intraparticle diffusivity is to assume that each species i has a constant diffusivity. Therefore, this function returns the value of the parameter intraparticle_diffusion from the DOGFISH_PARAM structure for each adsorbing species i. Each species may have a different diffusivity.

Parameters

i	index for the ith adsorbing species
1	index for the lth node in the domain
data	pointer to the DOGFISH_DATA structure

6.1.2.7 double default_FilmMTCoeff (int i, const void * data)

Default function for the film mass transfer coefficient.

The default film mass transfer coefficient will be to assume that this value is a constant for each species i. Therefore, this function returns the parameter value of film_transfer_coeff from the DOGFISH_PARAM structure for each adsorbing species i.

Parameters

i	index for the ith adsorbing species	
data	pointer to the DOGFISH_DATA structure	

6.1.2.8 double default_SurfaceConcentration (int i, const void * data)

Default function for the fiber surface concentration.

The default fiber surface concentration will be to assume that this value is a constant for each species i. Therefore, this function returns the parameter value of surface_concentration from the DOGFISH_PARAM structure for each adsorbing species i.

Parameters

i	index for the ith adsorbing species
data	pointer to the DOGFISH_DATA structure

6.1.2.9 int setup_DOGFISH_DATA (FILE * file, double(*)(int i, int l, const void *user_data) eval_R, double(*)(int i, int l, const void *user_data) eval_R, double(*)(int i, const void *user_data) eval_kf, double(*)(int i, const void *user_data) eval_qs, const void * user_data, DOGFISH_DATA * dog_dat)

Function will set up the memory and pointers for use in the DOGFISH simulations.

The pointers to the output file, parameter functions, and data structures are passed into this function to setup the problem in memory. This function must always be called prior to calling any other DOGFISH routine and after the DOGFISH DATA structure has been initialized.

Parameters

file	pointer to the output file to print out results
eval_R	function pointer for the retardation coefficient function
eval_DI	function pointer for the intraparticle diffusion function
eval_kf	function pointer for the film mass transfer function
eval_qs	function pointer for the surface concentration function
user_data	pointer for the user's own data structure (only if using custom functions)
dog_dat	pointer for the DOGFISH_DATA structure

6.1.2.10 int DOGFISH_Executioner (DOGFISH_DATA * dog_dat)

Function to serially call all other functions need to solve the system at one time step.

This function will call the DOGFISH_preprocesses function, followed by the FINCH solver functions for each species i, then call the DOGFISH_postprocesses function. After completion, this would have solved the diffusion physics for a single time step.

```
6.1.2.11 int set_DOGFISH_ICs ( DOGFISH_DATA * dog_dat )
```

Function called to evaluate the initial conditions for the time dependent problem.

This function will use information in DOGFISH_DATA to setup the initial conditions, initial parameter values, and initial sorption averages for each species. This function always assumes a constant initial condition for the sorption of each species.

```
6.1.2.12 int set_DOGFISH_timestep ( DOGFISH_DATA * dog_dat )
```

Function sets the time step size for the next step forward in the simulation.

This function will set the next time step size based on the spatial discretization of the fiber. Maximum time step size is locked at 0.5 hours.

```
6.1.2.13 int DOGFISH_preprocesses ( DOGFISH_DATA * dog_dat )
```

Function to perform preprocess actions to be used before calling any solver.

This function will call all of the parameter functions in order to establish boundary condition parameter values prior to calling the FINCH solvers.

```
6.1.2.14 int set_DOGFISH_params ( const void * user_data )
```

Function to calculate the values of all parameters for all species at all nodes.

This function is passed to the FINCH_DATA data structure and set as the setparams function pointer. FINCH calls this function during it's solver routine to setup the non-linear form of the problem and solve the non-linear system.

Parameters

user data	this is actually the DOGF	ISH DATA structure, but is	passed anonymously to FINCH
-----------	---------------------------	----------------------------	-----------------------------

```
6.1.2.15 int DOGFISH_postprocesses ( DOGFISH_DATA * dog_dat )
```

Function to perform post-solve actions such as printing out results.

This function increments the total_steps counter in DOGFISH_DATA to keep a running total of all solver steps taken. Additionally, it prints out the results of the current time simulation to the output file.

```
6.1.2.16 int DOGFISH_reset ( DOGFISH_DATA * dog_dat )
```

Function to reset the matrices and vectors and prepare for next time step.

This function will reset the matrix and vector information of DOGFISH_DATA and FINCH_DATA to prepare for the next simulation step in time.

```
6.1.2.17 int DOGFISH ( DOGFISH_DATA * dog_dat )
```

Function performs all necessary steps to step the diffusion simulation through time.

This function calls the initial conditions, set time step, executioner, and reset functions to step the simulation through time. It will only exit when the simulation time is reached or if an error occurs.

6.2 eel.h File Reference 261

```
6.1.2.18 int DOGFISH_TESTS ( )
```

Running DOGFISH tests.

This function is called from the UI to run a test simulation of DOGFISH. Ouput is stored in a DOGFISH_Test

Output.txt file in a sub-directory "output" from the directory in which the executable was called.

6.2 eel.h File Reference

Easy-access Element Library.

```
#include <stdio.h>
#include <math.h>
#include <iostream>
#include <fstream>
#include <stdlib.h>
#include <vector>
#include <time.h>
#include <float.h>
#include <string>
#include "error.h"
```

Classes

· class Atom

Atom object to hold information about specific atoms in the periodic table (click Atom to go to function definitions)

class PeriodicTable

Class object that store a digitial copy of all Atom objects.

Functions

• int EEL_TESTS ()

Test function to exercise the class objects and check for errors.

6.2.1 Detailed Description

Easy-access Element Library.

eel.cpp

This file contains two C++ objects: (i) Atom and (ii) PeriodicTable.

The Atom class defines all relavent information necessary for dealing with actual atoms. However, this is not necessarilly all the information that one may need for any simulation dealing with atoms. Instead, it is really just a place holder used to construct Molecules and hold oxidation state and molecular/atomic wieght information.

The PeriodicTable class creates a digital version of a complete periodic table. Further development of this object can make it possible to query this structure for a particular atom upon user request.

Warning

The Atom class is mostly complete, but the PeriodicTable object is just a place holder.

Author

Austin Ladshaw

Date

02/23/2015

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6.2.2 Function Documentation

```
6.2.2.1 int EEL_TESTS ( )
```

Test function to exercise the class objects and check for errors.

6.3 egret.h File Reference

Estimation of Gas-phase pRopErTies.

```
#include "macaw.h"
```

Classes

struct PURE GAS

Data structure holding all the parameters for each pure gas spieces.

• struct MIXED_GAS

Data structure holding information necessary for computing mixed gas properties.

Macros

```
• #define Rstd 8.3144621
```

Gas Constant in J/K/mol (or) L*kPa/K/mol (Standard Units)

#define RE3 8.3144621E+3

Gas Constant in cm³*kPa/K/mol (Convenient for density calculations)

• #define Po 100.0

Standard state pressure (kPa)

• #define Cstd(p, T) ((p)/(Rstd*T))

Calculation of concentration/density from partial pressure (Cstd = mol/L)

#define CE3(p, T) ((p)/(RE3*T))

Calculation of concentration/density from partial pressure (CE3 = $mol/cm^{\land}3$)

#define Pstd(c, T) ((c)*Rstd*T)

Calculation of partial pressure from concentration/density (c = mol/L)

#define PE3(c, T) ((c)*RE3*T)

Calculation of partial pressure from concentration/density ($c = mol/cm^{\wedge} 3$)

• #define Nu(mu, rho) ((mu)/(rho))

Calculation of kinematic viscosity from dynamic viscosity and density (cm²/s)

#define PSI(T) (0.873143 + (0.000072375*T))

Calculation of temperature correction factor for dynamic viscosity.

#define Dp_ij(Dij, PT) ((PT*Dij)/Po)

Calculation of the corrected binary diffusivity (cm $^{\land}$ 2/s)

#define D_ij(MWi, MWj, rhoi, rhoj, mui, muj) ((4.0 / sqrt(2.0)) * pow(((1/MWi)+(1/MWj)),0.5)) / pow((pow((rhoi/(1.385*mui)),2.0)/MWi),0.25)+ pow((pow((rhoi/(1.385*mui)),2.0)/MWj),0.25)),2.0)

Calculation of binary diffusion based on MW, density, and viscosity info (cm²/s)

#define Mu(muo, To, C, T) (muo * ((To + C)/(T + C)) * pow((T/To),1.5))

Calculation of single species viscosity from Sutherland's Equ. (g/cm/s)

#define D_ii(rhoi, mui) (1.385*mui/rhoi)

Calculation of self-diffusivity (cm $^{\wedge}$ 2/s)

#define ReNum(u, L, nu) (u*L/nu)

Calculation of the Reynold's Number (-)

#define ScNum(nu, D) (nu/D)

Calculation of the Schmidt Number (-)

#define FilmMTCoeff(D, L, Re, Sc) ((D/L)*(2.0 + (1.1*pow(Re,0.6)*pow(Sc,0.3))))

Calculation of film mass transfer coefficient (cm/s)

Functions

int initialize_data (int N, MIXED_GAS *gas_dat)

int calculate_properties (MIXED_GAS *gas_dat)

Function to initialize the MIXED_GAS structure based on number of gas species.

- - Function to set the values of the parameters in the gas phase.

Function to calculate the gas properties based on information in MIXED_GAS.

• int EGRET TESTS ()

Function runs a series of tests for the EGRET file.

6.3.1 Detailed Description

Estimation of Gas-phase pRopErTies.

egret.cpp

This file is responsible for estimating various temperature, pressure, and concentration dependent parameters to be used in other models for gas phase adsorption, mass transfer, and or mass transport. The goal of this file is to eliminate redundancies in code such that the higher level programs operate more efficiently and cleanly. Calculations made here are based on kinetic theory of gases, ideal gas law, and some emperical models that were developed to account for changes in density and viscosity with changes in temperature between standard temperatures and up to 1000 K.

Author

Austin Ladshaw

Date

01/29/2015

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6.3.2 Macro Definition Documentation

6.3.2.1 #define Rstd 8.3144621

Gas Constant in J/K/mol (or) L*kPa/K/mol (Standard Units)

6.3.2.2 #define RE3 8.3144621E+3

Gas Constant in cm[^]3*kPa/K/mol (Convenient for density calculations)

6.3.2.3 #define Po 100.0

Standard state pressure (kPa)

6.3.2.4 #define Cstd(p, T) ((p)/(Rstd*T))

Calculation of concentration/density from partial pressure (Cstd = mol/L)

6.3.2.5 #define CE3(p, T) ((p)/(RE3*T))

Calculation of concentration/density from partial pressure (CE3 = mol/cm^3)

6.3.2.6 #define Pstd(c, T) ((c)*Rstd*T)

Calculation of partial pressure from concentration/density (c = mol/L)

```
6.3.2.7 #define PE3( c, T) ((c)*RE3*T)
Calculation of partial pressure from concentration/density (c = mol/cm^{3})
6.3.2.8 #define Nu( mu, rho) ((mu)/(rho))
Calculation of kinematic viscosity from dynamic viscosity and density (cm<sup>2</sup>/s)
6.3.2.9 #define PSI(T) (0.873143 + (0.000072375*T))
Calculation of temperature correction factor for dynamic viscosity.
6.3.2.10 #define Dp_ij( Dij, PT ) ((PT*Dij)/Po)
Calculation of the corrected binary diffusivity (cm<sup>2</sup>/s)
6.3.2.11 #define D_ij( MWi, MWj, rhoi, rhoj, mui, muj ) ( (4.0 / sqrt(2.0)) * pow(((1/MWi)+(1/MWj)),0.5) ) / pow(
         (pow((pow((rhoi/(1.385*mui)),2.0)/MWi),0.25)+ pow((pow((rhoi/(1.385*muj)),2.0)/MWj),0.25)),2.0)
Calculation of binary diffusion based on MW, density, and viscosity info (cm<sup>2</sup>/s)
6.3.2.12 #define Mu( muo, To, C, T) (muo * ((To + C)/(T + C)) * pow((T/To),1.5))
Calculation of single species viscosity from Sutherland's Equ. (g/cm/s)
6.3.2.13 #define D_ii( rhoi, mui ) (1.385*mui/rhoi)
Calculation of self-diffusivity (cm<sup>2</sup>/s)
6.3.2.14 #define ReNum( u, L, nu ) (u*L/nu)
Calculation of the Reynold's Number (-)
6.3.2.15 #define ScNum( nu, D) (nu/D)
Calculation of the Schmidt Number (-)
6.3.2.16 #define FilmMTCoeff( D, L, Re, Sc ) ((D/L)*(2.0 + (1.1*pow(Re,0.6)*pow(Sc,0.3))))
Calculation of film mass transfer coefficient (cm/s)
6.3.3 Function Documentation
```

6.3.3.1 int initialize_data (int N, MIXED GAS * gas_dat)

Function to initialize the MIXED_GAS structure based on number of gas species.

This function will initialize the sizes of all vector objects in the MIXED_GAS structure based on the number of gas species indicated by N.

6.3.3.2 int set_variables (double PT, double us, double L, std::vector< double > & y, MIXED_GAS * gas_dat)

Function to set the values of the parameters in the gas phase.

The gas phase properties are a function of total pressure, gas temperature, gas velocity, characteristic length, and the mole fractions of each species in the gas phase. Prior to calculating the gas phase properties, these parameters must be set and updated as they change.

Parameters

PT	total gas pressure in kPa
T	gas temperature in K
us	gas velocity in cm/s
L	characteristic length in cm (this depends on the particular system)
у	vector of gas mole fractions of each species in the mixture
gas_dat	pointer to the MIXED_GAS data structure

6.3.3.3 int calculate_properties (MIXED GAS * gas_dat)

Function to calculate the gas properties based on information in MIXED_GAS.

This function uses the kinetic theory of gases, combined with other semi-empirical models, to predict and approximate several properties of the mixed gas phase that might be necessary when running any gas dynamical simulation. This includes mass and energy transfer equations, as well as adsorption kinetics in porous adsorbents.

```
6.3.3.4 int EGRET_TESTS ( )
```

Function runs a series of tests for the EGRET file.

The test looks at a standard air with 5 primary species of interest and calculates the gas properties from 273 K to 373 K. This function can be called from the UI.

6.4 error.h File Reference

All error types are defined here.

```
#include <iostream>
```

Macros

#define mError(i)

Enumerations

enum error_type {
 generic_error, file_dne, indexing_error, magpie_reverse_error,
 simulation_fail, invalid_components, invalid_boolean, invalid_molefraction,
 invalid_gas_sum, invalid_solid_sum, scenario_fail, out_of_bounds,
 non_square_matrix, dim_mis_match, empty_matrix, opt_no_support,
 invalid_fraction, ortho_check_fail, unstable_matrix, no_diffusion,
 negative_mass, negative_time, matvec_mis_match, arg_matrix_same,
 singular_matrix, matrix_too_small, invalid_size, nullptr_func,
 invalid_norm, vector_out_of_bounds, zero_vector, tensor_out_of_bounds,
 non_real_edge, nullptr_error, invalid_atom, invalid_proton,
 invalid_neutron, invalid_electron, invalid_valence, string_parse_error,
 unregistered_name, rxn_rate_error, invalid_species, duplicate_variable,
 missing_information, invalid_type, key_not_found, anchor_alias_dne,
 initial_error, not_a_token, read_error, invalid_console_input }

List of names for error type.

6.4 error.h File Reference 267

Functions

· void error (int flag)

Error function customizes output message based on flag.

6.4.1 Detailed Description

All error types are defined here.

error.cpp

This file defines all the different errors that may occur in any simulation in any file. Those errors are recognized by an enum with is then passed through to the error.cpp file that customizes the error message to the console. A macro will also print out the file name and line number where the error occured.

Author

Austin Ladshaw

Date

04/28/2014

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6.4.2 Macro Definition Documentation

6.4.2.1 #define mError(*i*)

Value:

6.4.3 Enumeration Type Documentation

6.4.3.1 enum error_type

List of names for error type.

Enumerator

```
generic_error
file_dne
indexing_error
magpie_reverse_error
simulation_fail
```

invalid_components

invalid_boolean

invalid_molefraction

invalid_gas_sum

invalid_solid_sum

scenario_fail

out_of_bounds

non_square_matrix

dim_mis_match

empty_matrix

opt_no_support

invalid_fraction

ortho_check_fail

unstable_matrix

no_diffusion

negative_mass

negative_time

matvec_mis_match

arg_matrix_same

singular_matrix

matrix_too_small

invalid_size

nullptr_func

invalid_norm

vector_out_of_bounds

zero_vector

tensor_out_of_bounds

non_real_edge

nullptr_error

invalid_atom

invalid_proton

invalid_neutron

invalid_electron

invalid_valence

string_parse_error

unregistered_name

rxn_rate_error

invalid_species

duplicate_variable

missing_information

invalid_type

key_not_found

anchor_alias_dne

initial_error

not_a_token

read_error

invalid_console_input

6.5 finch.h File Reference 269

6.4.4 Function Documentation

```
6.4.4.1 void error (int flag)
```

Error function customizes output message based on flag.

This error function is reference in the error.cpp file, but is not called by any other file. Instead, all other files call the mError(i) macro that expands into this error function call plus prints out the file name and line number where the error occured.

6.5 finch.h File Reference

Flux-limiting Implicit Non-oscillatory Conservative High-resolution scheme.

```
#include "macaw.h"
#include "lark.h"
```

Classes

struct FINCH_DATA

Data structure for the FINCH object.

Enumerations

• enum finch_solve_type { FINCH_Picard, LARK_Picard, LARK_PJFNK }

List of enum options to define the solver type in FINCH.

enum finch_coord_type { Cartesian, Cylindrical, Spherical }

List of enum options to define the coordinate system in FINCH.

Functions

double max (std::vector< double > &values)

Function returns the maximum in a list of values.

double min (std::vector< double > &values)

Function returns the minimum in a list of values.

double minmod (std::vector< double > &values)

Function returns the result of the minmod function acting on a list of values.

• int uTotal (FINCH DATA *dat)

Function integrates the conserved quantity to return it's total in the domain.

int uAverage (FINCH_DATA *dat)

Function integrates the conserved quantity to reture it's average in the domain.

• int check_Mass (FINCH_DATA *dat)

Function checks the unp1 vector for negative values and will adjust if needed.

int l_direct (FINCH_DATA *dat)

Function solves the discretized FINCH problem directly by assuming it is linear.

• int lark picard step (const Matrix< double > &x, Matrix< double > &G, const void *data)

Function to perform the necessary LARK Picard iterative method (not typically used)

• int nl_picard (FINCH_DATA *dat)

Function to solve the discretized FINCH problem iteratively by assuming it is non-linear.

int setup_FINCH_DATA (int(*user_callroutine)(const void *user_data), int(*user_setic)(const void *user_data), int(*user_timestep)(const void *user_data), int(*user_preprocess)(const void *user_data), int(*user_solve)(const void *user_data), int(*user_setparams)(const void *user_data), int(*user_discretize)(const void *user_data), int(*user_bcs)(const void *user_data), int(*user_res)(const Matrix< double > &x, Matrix< double > &s, const void *user_data), int(*user_precon)(const Matrix< double > &b, Matrix< double > &p, const void *user_data), int(*user_postprocess)(const void *user_data), int(*user_reset)(const void *user_data), int(*user_reset)(const void *user_data), int(*user_reset)(const void *user_data)

Function to setup memory and set user defined functions into the FINCH object.

void print2file_dim_header (FILE *Output, FINCH_DATA *dat)

Function will print out a dimension header for FINCH output.

void print2file_time_header (FILE *Output, FINCH_DATA *dat)

Function will print out a time header for FINCH output.

void print2file_result_old (FILE *Output, FINCH_DATA *dat)

Function will print out the old results to the variable u.

void print2file_result_new (FILE *Output, FINCH_DATA *dat)

Function will print out the new results to the variable u.

• void print2file_newline (FILE *Output, FINCH_DATA *dat)

Function will force print out a blank line.

void print2file_tab (FILE *Output, FINCH_DATA *dat)

Function will force print out a tab.

• int default execution (const void *user data)

Default executioner function for FINCH.

int default_ic (const void *user_data)

Default initial conditions function for FINCH.

int default_timestep (const void *user_data)

Default time step function for FINCH.

int default_preprocess (const void *user_data)

Default preprocesses function for FINCH.

int default_solve (const void *user_data)

Default solve function for FINCH.

int default_params (const void *user_data)

Default params function for FINCH.

int minmod_discretization (const void *user_data)

Minmod Discretization function for FINCH.

• int vanAlbada discretization (const void *user data)

Van Albada Discretization function for FINCH.

int ospre_discretization (const void *user_data)

Ospre Discretization function for FINCH.

int default_bcs (const void *user_data)

Default boundary conditions function for FINCH.

int default_res (const Matrix< double > &x, Matrix< double > &res, const void *user_data)

Default residual function for FINCH.

int default_precon (const Matrix< double > &b, Matrix< double > &p, const void *user_data)

Default preconditioning function for FINCH.

- int default_postprocess (const void *user_data)
- int default reset (const void *user data)

Default reset function for FINCH.

• int FINCH_TESTS ()

Function runs a particular FINCH test.

6.5 finch.h File Reference 271

6.5.1 Detailed Description

Flux-limiting Implicit Non-oscillatory Conservative High-resolution scheme.

finch.cpp

This is a conservative finite differences scheme based on the Kurganov and Tadmoor (2000) MUSCL scheme for non-linear conservation laws. It can solve 1-D conservation law problems in three different coordinate systems: (i) Cartesian - axial, (ii) Cylindrical - radial, and (iii) Spherical - radial. It is the backbone algorithm behind all 1-D PDE problems in the ecosystem software.

The form of the general conservation law problem that FINCH solves is...

```
z^{\wedge}d*R*du/dt = d/dz(z^{\wedge}d*D*du/dz) - d/dz(z^{\wedge}d*v*u) - z^{\wedge}d*k*u + z^{\wedge}d*S
```

where R, D, v, k, and S are the parameters of the problem and d, z, and u are the coordinates, spatial dimension, and conserved quantities, respectively. The parameter R is a retardation coefficient, D is a diffusion coefficient, v is a velocity, k is a reaction coefficient, and S is a forcing function or source/sink term.

FINCH supports the use of both Dirichlet and Neuman boundary conditions as the input/inlet condition and uses the No Flux (or Natural) boundary condition for the output/outlet of the domain. For radial problems, the outlet is always taken to the center of the cylindrical or spherical particle. This enforces the symmetry of the problem. For axial problems, the outlet is determined by the sign of the velocity term and is therefore choosen by the routine based on the actual flow direction in the domain.

Parameters of the problem can be coupled to the variable u and also be functions of space and time. The coupling of the parameters with the variable forces the problem to become non-linear, which requires iteration to solve. The default iterative method is a built-in Picard's method. This method is equivalent to an inexact Newton method, because we use the Linear Solve of this system as a weak approximation to the non-linear solve. Generally, this method is sufficient and is the most efficient. However, if a problem is particularly difficult to solve, then we can call some of the non-linear solvers developed in LARK. If PJFNK is used, then the Linear Solve for the FINCH problem is used as the Preconditioner for the Linear Solve in PJFNK.

This algorithm comes packaged with three different slope limiter functions to stabilize the velocity term for highly advectively dominate problems. The available slope limiters are: (i) minmod, (ii) van Albada, and (iii) ospre. By default, the FINCH setup function will set the slope limiter to ospre, because this method provides a reasonable compromise between accuracy and efficiency.

Slope Limiter Stats:

minmod -> Highest Accuracy, Lowest Efficiency van Albada -> Lowest Accuracy, Highest Efficiency ospre -> Average Accuracy, Average Efficiency

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Date

01/29/2015

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6.5.2 Enumeration Type Documentation

6.5.2.1 enum finch_solve_type

List of enum options to define the solver type in FINCH.

Enumerator

FINCH_Picard LARK_Picard LARK_PJFNK

6.5.2.2 enum finch_coord_type

List of enum options to define the coordinate system in FINCH.

Enumerator

Cartesian

Cylindrical

Spherical

6.5.3 Function Documentation

6.5.3.1 double max (std::vector< double > & values)

Function returns the maximum in a list of values.

6.5.3.2 double min (std::vector< double > & values)

Function returns the minimum in a list of values.

6.5.3.3 double minmod (std::vector < double > & values)

Function returns the result of the minmod function acting on a list of values.

6.5.3.4 int uTotal (FINCH_DATA * dat)

Function integrates the conserved quantity to return it's total in the domain.

6.5.3.5 int uAverage (FINCH_DATA * dat)

Function integrates the conserved quantity to reture it's average in the domain.

6.5.3.6 int check_Mass (FINCH_DATA * dat)

Function checks the unp1 vector for negative values and will adjust if needed.

This function can be turned off or on in the FINCH_DATA structure. Typically, you will want to leave this on so that the routine does not return negative values for u. However, if you want to get negative values of u, then turn this option off.

6.5 finch.h File Reference 273

```
6.5.3.7 int I_direct ( FINCH_DATA * dat )
```

Function solves the discretized FINCH problem directly by assuming it is linear.

```
6.5.3.8 int lark_picard_step ( const Matrix < double > & x, Matrix < double > & G, const void * data )
```

Function to perform the necessary LARK Picard iterative method (not typically used)

```
6.5.3.9 int nl_picard ( FINCH_DATA * dat )
```

Function to solve the discretized FINCH problem iteratively by assuming it is non-linear.

Note

If the problem is actually linear, then this will solve it in one iteration. So it may be best to always assume the problem is non-linear.

6.5.3.10 int setup_FINCH_DATA (int(*)(const void *user_data) user_callroutine, int(*)(const void *user_data) user_setic, int(*)(const void *user_data) user_timestep, int(*)(const void *user_data) user_preprocess, int(*)(const void *user_data) user_setparams, int(*)(const void *user_data) user_discretize, int(*)(const void *user_data) user_bcs, int(*)(const Matrix< double > &x, Matrix< double > &res, const void *user_data) user_res, int(*)(const Matrix< double > &b, Matrix< double > &p, const void *user_data) user_precon, int(*)(const void *user_data) user_postprocess, int(*)(const void *user_data) user_reset, FINCH_DATA * dat, const void * param_data)

Function to setup memory and set user defined functions into the FINCH object.

This function MUST be called prior to running any FINCH based simulation. However, you are only every required to provide this function with the FINCH_DATA pointer. It is recommended, however, that you do provide the user ← setparams and param data pointers, as these will likely vary significantly from problem to problem.

After the problem is setup in memory, you do not technically have to have FINCH call all of it's own functions. You can write your own executioner, initial conditions, and other functions and decided how and when everything is called. Then just call the solve function in FINCH_DATA when you want to use the FINCH solver. This is how FINCH is used in SKUA, SCOPSOWL, DOGFISH, and MONKFISH.

Parameters

user_callroutine	function pointer the the call routine function
user_setic	function pointer to set initial conditions for problem
user_timestep	function pointer to set the next time step
user_preprocess	function pointer to setup a preprocess operation
user_solve	function pointer to solve the system of equations
user_setparams	function pointer to set the parameters in the problem (always override this)
user_discretize	function pointer to select discretization scheme for the problem
user_bcs	function pointer to evaluate boundary conditions for the problem
user_res	function pointer to evaluate non-linear residuals for the problem
user_precon	function pointer to perform a linear preconditioning operation
user_postprocess	function pointer to setup a postprocess operation
user_reset	function pointer to reset stateful data for next simulation
dat	pointer to the FINCH_DATA structure
param_data	user supplied pointer to a data structure needed in user_setparams

```
6.5.3.11 void print2file_dim_header ( FILE * Output, FINCH_DATA * dat )
```

Function will print out a dimension header for FINCH output.

```
6.5.3.12 void print2file_time_header ( FILE * Output, FINCH_DATA * dat )
```

Function will print out a time header for FINCH output.

```
6.5.3.13 void print2file_result_old ( FILE * Output, FINCH_DATA * dat )
```

Function will print out the old results to the variable u.

```
6.5.3.14 void print2file_result_new ( FILE * Output, FINCH_DATA * dat )
```

Function will print out the new results to the variable u.

```
6.5.3.15 void print2file_newline ( FILE * Output, FINCH DATA * dat )
```

Function will force print out a blank line.

```
6.5.3.16 void print2file_tab ( FILE * Output, FINCH_DATA * dat )
```

Function will force print out a tab.

```
6.5.3.17 int default_execution ( const void * user_data )
```

Default executioner function for FINCH.

The default executioner function for FINCH assumes the user_data parameter is the FINCH_DATA structure and calls the preprocesses, solve, postprocesses, checkMass, uTotal, and uAverage functions in that order.

```
6.5.3.18 int default_ic ( const void * user_data )
```

Default initial conditions function for FINCH.

The default initial condition function for FINCH assumes the user_data parameter is the FINCH_DATA structure and sets the initial values of all system parameters according to the given constants in that structure.

```
6.5.3.19 int default_timestep ( const void * user_data )
```

Default time step function for FINCH.

The default time step function for FINCH assumes the user_data parameter is the FINCH_DATA structure and sets the time step to 1/2 the mesh size or bases the time step off of the CFL condition if the problem is not being solved iteratively and involves an advective portion.

```
6.5.3.20 int default_preprocess ( const void * user_data )
```

Default preprocesses function for FINCH.

The default preprocesses function for FINCH assumes the user_data parameter is the FINCH_DATA structure and does nothing.

6.5 finch.h File Reference 275

```
6.5.3.21 int default_solve ( const void * user_data )
```

Default solve function for FINCH.

The default solve function for FINCH assumes the user_data parameter is the FINCH_DATA structure and calls the corresponding solution method depending on the users conditions.

```
6.5.3.22 int default_params ( const void * user_data )
```

Default params function for FINCH.

The default params function for FINCH assumes the user_data parameter is the FINCH_DATA structure and sets the values of all parameters at all nodes equal to the values of those parameters at the boundaries.

```
6.5.3.23 int minmod_discretization ( const void * user_data )
```

Minmod Discretization function for FINCH.

The minmod discretization function for FINCH assumes the user_data parameter is the FINCH_DATA structure and discretizes the time and space portion of the problem with 2nd order finite differences and uses the minmod slope limiter function to stabilize the advective physics.

```
6.5.3.24 int vanAlbada_discretization ( const void * user_data )
```

Van Albada Discretization function for FINCH.

The van Albada discretization function for FINCH assumes the user_data parameter is the FINCH_DATA structure and discretizes the time and space portion of the problem with 2nd order finite differences and uses the van Albada slope limiter function to stabilize the advective physics.

```
6.5.3.25 int ospre_discretization ( const void * user_data )
```

Ospre Discretization function for FINCH.

The ospre discretization function for FINCH assumes the user_data parameter is the FINCH_DATA structure and discretizes the time and space portion of the problem with 2nd order finite differences and uses the ospre slope limiter function to stabilize the advective physics. This is the default discretization function.

```
6.5.3.26 int default_bcs ( const void * user_data )
```

Default boundary conditions function for FINCH.

The default boundary conditions function for FINCH assumes the user_data parameter is the FINCH_DATA structure and sets the boundary conditions according to the type of problem requested. The input BCs will always be either Neumann or Dirichlet and the output BC will always be a zero flux Neumann BC.

```
6.5.3.27 int default_res ( const Matrix < double > & x, Matrix < double > & res, const void * user_data )
```

Default residual function for FINCH.

The default residual function for FINCH assumes the user_data parameter is the FINCH_DATA structure and calls the setparams function (passing the param_data structure), the discretization function, and the set BCs functions, in that order. It then forms the implicit and explicit side residuals that go into the iterative solver.

```
6.5.3.28 int default_precon ( const Matrix < double > & b, Matrix < double > & p, const void * user_data )
```

Default preconditioning function for FINCH.

The default preconditioning function for FINCH assumes the user_data parameter is the FINCH_DATA structure and performs a tridiagonal linear solve using a Modified Thomas Algorithm. This preconditioner will solve the linear problem exactly if there is no advective portion of the physics. Additionally, this preconditioner is also used as the basis for forming the default FINCH non-linear iterations and is sufficient for solving most problems.

```
6.5.3.29 int default_postprocess ( const void * user_data )
```

The default postprocesses function for FINCH assumes the user_data parameter is the FINCH_DATA structure and does nothing.

```
6.5.3.30 int default_reset ( const void * user_data )
```

Default reset function for FINCH.

The default reset function for FINCH assumes the user_data parameter is the FINCH_DATA structure and sets all old state parameters and variables to the new state.

```
6.5.3.31 int FINCH_TESTS ( )
```

Function runs a particular FINCH test.

The FINCH_TESTS function is used to exercise and test out the FINCH algorithms for correctness, efficiency, and accuracy. This test should never report a failure.

6.6 flock.h File Reference

FundamentaL Off-gas Collection of Kernels.

```
#include "macaw.h"
#include "egret.h"
#include "finch.h"
#include "lark.h"
#include "skua.h"
#include "scopsowl.h"
#include "gsta_opt.h"
#include "magpie.h"
#include "skua_opt.h"
#include "scopsowl_opt.h"
#include "yaml_wrapper.h"
```

6.6.1 Detailed Description

FundamentaL Off-gas Collection of Kernels.

This is just a .h file that holds all the includes necessary to develop and run simulations for adsorption and/or mass/energy transfer problems for gaseous systems. Include this file into any other project or source code that needs the methods below.

Files Included in FLOCK

macaw.h egret.h finch.h lark.h skua.h scopsowl.h gsta_opt.h magpie.h skua_opt.h scopsowl_opt.h yaml_wrapper.h

Author

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Date

04/28/2014

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6.7 gsta_opt.h File Reference

Generalized Statistical Thermodynamic Adsorption (GSTA) Optimization Routine.

```
#include "lmcurve.h"
#include <stdio.h>
#include <math.h>
#include <iostream>
#include <fstream>
#include <stdlib.h>
#include <vector>
#include <time.h>
#include <float.h>
#include <string>
#include "error.h"
```

Classes

struct GSTA_OPT_DATA

Data structure used in the GSTA optimization routines.

Macros

```
• #define Po 100.0
```

Standard State Pressure - Units: kPa.

#define R 8.3144621

```
Gas Constant - Units: J/(K*mol) = kB * Na.
```

• #define Na 6.0221413E+23

Avagadro's Number - Units: molecules/mol.

Functions

• int roundIt (double d)

Function rounds a double to an integer.

• int twoFifths (int m)

Function returns the rounded two-fifths result of int m.

• int orderMag (double x)

Function returns the order of magnitude for the parameter x.

int minValue (std::vector< int > &array)

Function returns the minimum integer in an array of integers.

int minIndex (std::vector< double > &array)

Function returns the index of the minimum integer in an array of integers.

int avgPar (std::vector< int > &array)

Function returns the average integer value in an array of integers.

double avgValue (std::vector< double > &array)

Function returns an average in an array of doubles.

double weightedAvg (double *enorm, double *x, int n)

Function returns a weighted average in an array.

double rSq (double *x, double *y, double slope, double vint, int m dat)

Function calculates the Coefficient of Determination (R Squared) for the temperature regression.

• bool isSmooth (double *par, void *data)

Function looks at the list of parameters to check if they are smoothly changing.

void orthoLinReg (double *x, double *y, double *par, int m dat, int n par)

Function performs an Orthogonal Linear Regression on a set of data.

void eduGuess (double *P, double *q, double *par, int k, int m_dat, void *data)

Function will formed an educated guess for the next set of parameters in the GSTA analysis.

double gstaFunc (double p, const double *K, double qmax, int n_par)

Function evaluates the result of the GSTA isotherm model.

• double gstaObjFunc (double *t, double *y, double *par, int m_dat, void *data)

Function to evaulate the GSTA objective function value.

void eval_GSTA (const double *par, int m_dat, const void *data, double *fvec, int *info)

Function to evaluate the GSTA model and feed into the Imfit routine.

• int gsta_optimize (const char *fileName)

Function to perform the GSTA optimization routine.

6.7.1 Detailed Description

Generalized Statistical Thermodynamic Adsorption (GSTA) Optimization Routine.

gsta opt.cpp

Optimization routine developed for the GSTA isotherm and data analysis. This algorithm was the primary subject of a publication made in Fluid Phase Equilibria. Please refer to the below paper for technical information about the algorithms.

Reference: Ladshaw, Yiacoumi, Tsouris, and DePaoli, Fluid Phase Equilibria, 388, 169-181, 2015.

The GSTA model was first introduced by Llano-Restrepo and Mosquera (2009). Please refer to the below reference for theoretical information about the model.

Reference: Llano-Restrepo and Mosquera, Fluid Phase Equilibria, 283, 73-88, 2009.

Author

Austin Ladshaw

Date

12/17/2013

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This software was designed and built at the Georgia Institute of Technology by Austin Ladshaw for PhD research in the area of adsorption and surface science. Copyright (c) 2015, all rights reserved.

6.7.2 Macro Definition Documentation

6.7.2.1 #define Po 100.0

Standard State Pressure - Units: kPa.

6.7.2.2 #define R 8.3144621

Gas Constant - Units: J/(K*mol) = kB * Na.

6.7.2.3 #define Na 6.0221413E+23

Avagadro's Number - Units: molecules/mol.

6.7.3 Function Documentation

6.7.3.1 int roundIt (double d)

Function rounds a double to an integer.

This function returns a rounded value of d. Rounding up for any decimal larger than 0.5 and down for all else.

6.7.3.2 int twoFifths (int m)

Function returns the rounded two-fifths result of int m.

This function is used to determine what the maximum number of parameters should be based on the number of data points m. It is designed to prevent the algorithms from "over fitting" the data.

6.7.3.3 int orderMag (double x)

Function returns the order of magnitude for the parameter x.

This function is used to help create initial guesses for the new GSTA parameters that are being optimized for. In order to make sure that those parameters are considered relavent in the optimization routine, we need to make the initial guesses to be around the same order of magnitude of the other GSTA parameters.

```
6.7.3.4 int minValue ( std::vector < int > & array )
```

Function returns the minimum integer in an array of integers.

This function is used to determine the minimum number of GSTA parameters that were required to adequately fit the isotherm data.

```
6.7.3.5 int minIndex ( std::vector < double > & array )
```

Function returns the index of the minimum integer in an array of integers.

This function identifies the index of the minimum number of parameters needed for the GSTA model to fit the data. This index is common for all vectors in the GSTA_OPT_DATA structure and is used to identify the most suitable solution.

```
6.7.3.6 int avgPar ( std::vector < int > & array )
```

Function returns the average integer value in an array of integers.

This function is used to identify the average number of parameters that all the data fitting needed for each GSTA analysis.

```
6.7.3.7 double avgValue ( std::vector < double > & array )
```

Function returns an average in an array of doubles.

```
6.7.3.8 double weightedAvg ( double * enorm, double * x, int n )
```

Function returns a weighted average in an array.

This averaging scheme is used to approximate the qmax parameter for the GSTA isotherm model, if that value is unknown. The weighting is based on the euclidean norms of all the fits of the data. Smaller norms are more heavily weighted since they represent a better fit of the data. Once averaging is complete and we have an estimate for qmax, the entire algorithm is re-run holding that qmax constant.

Parameters

enorm	array of euclidean norms from the fitting of the data
X	array of optimum qmax values to be averaged
n	the number of enorm and x values in the array

6.7.3.9 double rSq (double * x, double * y, double slope, double vint, int m_dat)

Function calculates the Coefficient of Determination (R Squared) for the temperature regression.

This function is used to determine the "fittness" of the linear regression performed on the temperature independent parameters of the GSTA isotherm. A good linear regression should return a value between 1.0 and 0.9.

X	observations in the x-axis
У	observations in the y-axis

Parameters

slope	slope of the linear regression
vint	intercept of the linear regression
m_dat	number of data points used in the linear regression

6.7.3.10 bool isSmooth (double * par, void * data)

Function looks at the list of parameters to check if they are smoothly changing.

This function takes the parameter array par and GSTA_OPT_DATA structure and checks to see if those parameters are changing smoothly. If they are erratic or non-smooth, then it could be an indication of "over fitting" of the data.

6.7.3.11 void orthoLinReg (double * x, double * y, double * par, int m_dat , int n_par)

Function performs an Orthogonal Linear Regression on a set of data.

This function takes an array of x and y observations and performs an orthogonal linear regression on that information to find optimum parameters for slope and intercept.

Parameters

Х	array of x-axis observations
У	array of y-axis observations
par	array of parameter results after regression
m_dat	number of data points or observations
n_par	number of parameters to seek (if n_par != 1 or 2, then par[0] = intercept and par[1] = slope)

6.7.3.12 void eduGuess (double * P, double * q, double * par, int k, int m dat, void * data)

Function will formed an educated guess for the next set of parameters in the GSTA analysis.

This function takes partial pressure and adsorption observations, P and q, and tries to give a decent initial guess to what the GSTA parameters, par, will be for the next iteration.

Parameters

Р	partial pressure observations in the data (kPa)
q	adsorption observations in the data (any units)
par	parameter array for the GSTA isotherm
k	index of the current number of parameters being considered
m_dat	number of pressure-adsorption observations in the isotherm
data	pointer to the GSTA_OPT_DATA data structure

6.7.3.13 double gstaFunc (double p, const double *K, double qmax, int n_par)

Function evaluates the result of the GSTA isotherm model.

This function will evaluate the GSTA model and return the adsorbed amount given the current partial pressure p and the equilibrium parameters K.

Parameters

р	current partial pressure (kPa)
K	array of equilibrium parameters (1/kPa^n)
qmax	the theorectical maximum capacity for the isotherm
n_par	the number of equilibrium parameters

6.7.3.14 double gstaObjFunc (double * t, double * y, double * par, int m_dat , void * data)

Function to evaulate the GSTA objective function value.

The objective function seeks to penalize the relative fittness of the model based on the number of parameters it took to minimize the euclidean norms. By penalizing the fittness of the model in this fashion, we can find the best solution to the system that required the least number of equilibrium parameters.

6.7.3.15 void eval_GSTA (const double * par, int m_dat, const void * data, double * fvec, int * info)

Function to evaluate the GSTA model and feed into the Imfit routine.

This function will formulate the residuals that go into the Levenberg-Marquardt's Algorithm for non-linear least squares regression. The form of this function is specific to how we interface with the Imfit routines.

6.7.3.16 int gsta_optimize (const char * fileName)

Function to perform the GSTA optimization routine.

This function is callable from the UI and is used to find the optimum parameters of the GSTA isotherm model given a particular set of isotherm data for single-component adsorption equilibria.

Parameters

fileName	name of the input file that holds the isotherm data
----------	---

Note

The input file for the GSTA optimization routine is a text file holding the necessary information and data needed to run the routine. That input file has a very specific format that is detailed below.

Number of Isotherm Curves

Theoretical Maximum Adsorption Capacity (if unknown, provide 0)

Temperature of the ith Isotherm (K)

Number of Data points for the ith Isotherm

Partial Pressure (kPa) [tab] Corresponding Adsorbed Amount (any units)

(2nd Line down is repeated for all isotherms you are optimizing on...)

Example:

2 21.0 298.15 4 0.000165483 2.77

```
0.000306379 2.75
0.00044922 5.00
0.000939259 10.40
313.15
4
0.000589636 2.75
0.001063584 3.70
0.001351836 4.2
0.001543464 4.6
```

The above example would be for 2 sets of isotherms at 298.15 and 313.15 K, respectively. Maximum adsorption capacity is given as 21 (which in this has units of wt%). Each isotherm has 4 data points, which are given in a list as p (kPa) and q (wt%) pairs. Units of adsorption don't matter as long as they are consistent. If you give maximum capacity in mol/kg, then the q's in the lists must also be in mol/kg.

6.8 lark.h File Reference

Linear Algebra Residual Kernels.

```
#include "macaw.h"
#include <float.h>
```

Classes

struct ARNOLDI_DATA

Data structure for the construction of the Krylov subspaces for a linear system.

struct GMRESLP DATA

Data structure for implementation of the Restarted GMRES algorithm with Left Preconditioning.

struct GMRESRP DATA

Data structure for the Restarted GMRES algorithm with Right Preconditioning.

struct PCG_DATA

Data structure for implementation of the PCG algorithms for symmetric linear systems.

struct BiCGSTAB_DATA

Data structure for the implementation of the BiCGSTAB algorithm for non-symmetric linear systems.

struct CGS DATA

Data structure for the implementation of the CGS algorithm for non-symmetric linear systems.

struct OPTRANS DATA

Data structure for implementation of linear operator transposition.

struct GCR DATA

Data structure for the implementation of the GCR algorithm for non-symmetric linear systems.

• struct GMRESR DATA

Data structure for the implementation of GCR with Nested GMRES preconditioning (i.e., GMRESR)

struct KMS_DATA

Data structure for the implemenation of the Krylov Multi-Space (KMS) Method.

struct QR DATA

Data structure for the implementation of a QR solver given some invertable linear operator.

struct PICARD_DATA

Data structure for the implementation of a Picard or Fixed-Point iteration for non-linear systems.

struct BACKTRACK_DATA

Data structure for the implementation of Backtracking Linesearch.

struct PJFNK_DATA

Data structure for the implementation of the PJFNK algorithm for non-linear systems.

struct NUM_JAC_DATA

Data structure to form a numerical jacobian matrix with finite differences.

Macros

• #define MIN_TOL 1e-15

Minimum Allowable Tolerance for linear and non-linear problems.

Enumerations

enum krylov_method {
 GMRESLP, PCG, BiCGSTAB, CGS,
 FOM, GMRESRP, GCR, GMRESR,
 KMS, QR }

Enum of definitions for linear solver types in PJFNK.

Functions

- int update_arnoldi_solution (Matrix< double > &x, Matrix< double > &x0, ARNOLDI_DATA *arnoldi_dat)

 Function to update the linear vector x based on the Arnoldi Krylov subspace.
- int arnoldi (int(*matvec)(const Matrix< double > &v, Matrix< double > &w, const void *data), int(*precon)(const Matrix< double > &b, Matrix< double > &p, const void *data), Matrix< double > &r0, ARNOLDI_DATA *arnoldi_dat, const void *matvec_data, const void *precon_data)

Function to factor a linear operator into an orthonormal basis and upper Hessenberg matrix.

int gmresLeftPreconditioned (int(*matvec)(const Matrix< double > &v, Matrix< double > &w, const void *data), int(*precon)(const Matrix< double > &b, Matrix< double > &p, const void *data), Matrix< double > &b, GMRESLP_DATA *gmreslp_dat, const void *matvec_data, const void *precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with GMRESLP.

int fom (int(*matvec)(const Matrix< double > &v, Matrix< double > &w, const void *data), int(*precon)(const Matrix< double > &b, Matrix< double > &b, GMRESLP_DATA *gmreslp_dat, const void *matvec_data, const void *precon_data)

Function to directly solve a non-symmetric, indefinite linear system with FOM.

int gmresRightPreconditioned (int(*matvec)(const Matrix< double > &v, Matrix< double > &w, const void *data), int(*precon)(const Matrix< double > &b, Matrix< double > &p, const void *data), Matrix< double > &b, GMRESRP_DATA *gmresrp_dat, const void *matvec_data, const void *precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with GMRESRP.

int pcg (int(*matvec)(const Matrix< double > &p, Matrix< double > &Ap, const void *data), int(*precon)(const Matrix< double > &r, Matrix< double > &z, const void *data), Matrix< double > &b, PCG_DATA *pcg_dat, const void *matvec_data, const void *precon_data)

Function to iteratively solve a symmetric, definite linear system with PCG.

int bicgstab (int(*matvec)(const Matrix< double > &p, Matrix< double > &Ap, const void *data), int(*precon)(const Matrix< double > &r, Matrix< double > &z, const void *data), Matrix< double > &b, BiCGSTAB_DATA *bicg_dat, const void *matvec_data, const void *precon_data)

Function to iteratively solve a non-symmetric, definite linear system with BiCGSTAB.

int cgs (int(*matvec)(const Matrix< double > &p, Matrix< double > &Ap, const void *data), int(*precon)(const Matrix< double > &r, Matrix< double > &z, const void *data), Matrix< double > &b, CGS_DATA *cgs_dat, const void *matvec_data, const void *precon_data)

Function to iteratively solve a non-symmetric, definite linear system with CGS.

int operatorTranspose (int(*matvec)(const Matrix< double > &v, Matrix< double > &Av, const void *data),
 Matrix< double > &r, Matrix< double > &u, OPTRANS_DATA *transpose_dat, const void *matvec_data)

Function that is used to perform transposition of a linear operator and results in a new vector A^T*r=u.

int gcr (int(*matvec)(const Matrix < double > &x, Matrix < double > &Ax, const void *data), int(*precon)(const Matrix < double > &r, Matrix < double > &Mr, const void *data), Matrix < double > &b, GCR_DATA *gcr_dat, const void *matvec_data, const void *precon_data)

Function to iteratively solve a non-symmetric, definite linear system with GCR.

• int gmresrPreconditioner (const Matrix< double > &r, Matrix< double > &Mr, const void *data)

Function used in conjunction with GMRESR to apply GMRESRP iterations as a preconditioner.

• int gmresr (int(*matvec)(const Matrix< double > &x, Matrix< double > &Ax, const void *data), int(*terminal_precon)(const Matrix< double > &r, Matrix< double > &Mr, const void *data), Matrix< double > &b, GMRESR_DATA *gmresr_dat, const void *matvec_data, const void *term_precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with GMRESR.

 $\bullet \ \, \text{int kmsPreconditioner (const Matrix} < \ \, \text{double} > \&r, \ \, \text{Matrix} < \ \, \text{double} > \&\text{Mr, const void} \ * \text{data}) \\$

Preconditioner function for the Krylov Multi-Space.

int krylovMultiSpace (int(*matvec)(const Matrix< double > &x, Matrix< double > &Ax, const void *data), int(*terminal_precon)(const Matrix< double > &r, Matrix< double > &Mr, const void *data), Matrix< double > &b, KMS_DATA *kms_dat, const void *matvec_data, const void *term_precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with KMS.

int QRsolve (int(*matvec)(const Matrix< double > &x, Matrix< double > &Ax, const void *data), Matrix
 double > &b, QR_DATA *qr_dat, const void *matvec_data)

Function to solve a dense linear operator system using QR factorization.

• int picard (int(*res)(const Matrix< double > &x, Matrix< double > &r, const void *data), int(*evalx)(const Matrix< double > &x0, Matrix< double > &x, const void *data), Matrix< double > &x, PICARD_DATA *picard_dat, const void *res_data, const void *evalx_data)

Function to iteratively solve a non-linear system using the Picard or Fixed-Point method.

int jacvec (const Matrix< double > &v, Matrix< double > &Jv, const void *data)

Function to form a linear operator of a Jacobian matrix used along with the PJFNK method.

int backtrackLineSearch (int(*feval)(const Matrix< double > &x, Matrix< double > &F, const void *data),
 Matrix< double > &pk, double normFk, BACKTRAC←
 K DATA *backtrack dat, const void *feval data)

Function to perform a Backtracking Line Search operation to smooth out convergence of PJFNK.

int pjfnk (int(*res)(const Matrix< double > &x, Matrix< double > &F, const void *data), int(*precon)(const Matrix< double > &r, Matrix< double > &x, PJFNK_DATA *pjfnk← _dat, const void *res_data, const void *precon_data)

Function to perform the PJFNK algorithm to solve a non-linear system of equations.

int NumericalJacobian (int(*Func)(const Matrix< double > &x, Matrix< double > &F, const void *user_←
data), const Matrix< double > &x, Matrix< double > &J, int Nx, int Nf, NUM_JAC_DATA *jac_dat, const void
*user_data)

Function to form a full numerical Jacobian matrix from a given non-linear function.

• int LARK TESTS ()

Function that runs a variety of tests on all the functions in LARK.

6.8.1 Detailed Description

Linear Algebra Residual Kernels.

lark.cpp

The functions contained within are designed to solve generic linear and non-linear square systems of equations given a function argument and data from the user. Optionally, the user can also provide a function to return a preconditioning result that will be applied to the system.

Having the user define how the preconditioning is carried out provides two major advantages: (1) we do not need to store and large, sparse preconditioning matrices and instead only store the preconditioned vector result and (2) this allows the user to use any kind of preconditioner they see fit for their problem.

The Arnoldi function is typically not called by the user, but can be if desired. It accepts the function arguments and a residual vector to form an orthonormal basis of the Krylov subspace using the Modified Gram-Schmidt process (aka Arnoldi Iteration). This function is called by GMRES to iteratively solve a linear system of equations. Note that

you can use this function to directly solve the linear system as long as that system is not too large. Construction of the basis is expensive, which is why this is used as a sub-function of an iterative method.

The Restarted GMRES function will accept function arguments for a linear system and attempt to solve said system iteratively by constructing an orthonormal basis from the Krylov function. Note that this GMRES function does support restarting and will use restarting by default if the linear system is too large.

Also included is a GMRES algorithm without restarting. This will directly solve the linear system within residual tolerance using a Full Orthogonal basis set of that system. It is equivalent to calling the Krylov method with the k parameter equal to N (i.e. the number of equations). This method is nick-named the Full Othogonalization Method (FOM), although the true FOM algorithm in literature is slightly different.

The PJFNK function will accept function arguments for a square, non-linear system of equations and attempt to solve it iteratively using both the GMRES and Krylov functions with Newton's method to convert the non-linear system into a linear system.

Also built here is a PCG implementation for solving symmetric linear systems. Can also be called by PJFNK if we know that the linear system (i.e. the Jacobian) is symmetric. This algorithm is significantly more efficient than GMRES, but is only valid if the system of equations is symmetric.

Other linear solvers implemented in this work are the BiCGSTAB and CGS algorithms for non-symmetric, positive definite matrices. These algorithms are significantly more computationally efficient than GMRES or FOM. However, they can both break down if the linear system is poorly conditioned. In general, you only want to use these methods if you have preconditioning available and your linear system is very, very large. Otherwise, you will be better suited to using GMRES or FOM.

There is also an implementation of the Generalized Conjugate Residual (GCR) method with and without restarting. This is a GMRES-like method that should give the exact solution within N iterations, where N is the original size of the matrix. Built ontop of the GCR method is a GMRESR (or GMRES Recursive) algorithm that uses GCR as the base method and performs GMRESRP iterations as a preconditioner at each iteration of GCR. This is the only linear solver that has built-in preconditioning. As a result, it may be slower than other algorithms for simple problems, but generally will have much better convergence behavior and will almost always give better residual reduction, even for hard to solve problems.

We have also developed a novel/experimental iterative method based on the idea of recursively preconditioning a Krylov Subspace with more Krylov Subspaces. We have called with algorithm the Krylov Multi-Space (KMS) method. This algorithm is based on publications from Vorst and Vuik (1991) and Saad (1993). The idea is too use the FGMRES algorithm developed by Saad (1993) and precondition it with more FGMRES steps, i.e., nesting the iterations as Vorst and Vuik (1991) had proposed. In this way, we have created a generalized Krylov Subspace method that has it's own variable preconditioner that can be adjusted depending on the user's desired complexity and convergence rate. If the levels of recursion requested is zero, then this algorithm is exactly equal to GMRES with right preconditioning. If the level is one, then it is FGMRES with a GMRES preconditioner. However, we allow the levels of recursion to reach up to 5, thus allowing us to precondition the preconitioners with more GMRES steps. This can result is significantly faster convergence rates, but is typically only necessary for very large or difficult to solve problems.

NOTE: There are three GMRES implementations: (i) gmresLP, (ii) fom, and (iii) gmresRP. GMRESLP is a restarted GMRES implementation that is left preconditioned and only checks the residual on the outer loops. This may be less efficient than GMRESRP, which can check both outer and inner loop residuals. However, GMRESRP has to use right preconditioning, which also slightly changes the convergence behavior of the linear system. GMRES with left preconditioning and without restarting will just build the full subspace by default, thus solving the system exactly, but may require too much memory. You can do a GMRESRP unrestarted by specifying that the restart parameter be equal to the size of the problem.

Basic Implementation Details:

```
Linear Solvers -> Solve Ax=b for x

Non-Linear Solvers -> Solve F(x)=0 for x

All implementations require system size to be 2 or greater
```

Author

Austin Ladshaw

Date

10/14/2014

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6.8.2 Macro Definition Documentation

6.8.2.1 #define MIN_TOL 1e-15

Minimum Allowable Tolerance for linear and non-linear problems.

6.8.3 Enumeration Type Documentation

6.8.3.1 enum krylov_method

Enum of definitions for linear solver types in PJFNK.

Enum delineates the available Krylov Subspace methods that can be used to solve the linear sub-problem at each non-linear iteration in a Newton method.

Enumerator

GMRESLP

PCG

BiCGSTAB

CGS

FOM

GMRESRP

GCR

GMRESR

KMS

QR

6.8.4 Function Documentation

6.8.4.1 int update_arnoldi_solution (Matrix < double > & x, Matrix < double > & x0, ARNOLDI_DATA * arnoldi_dat)

Function to update the linear vector x based on the Arnoldi Krylov subspace.

This function will update a solution vector x based on the previous solution x0 given the orthonormal basis and upper Hessenberg matrix formed in the Arnoldi algorithm. Updating is automatically called by the GMRESLP function. It is expected that the Arnoldi algorithm has already been called prior to calling this function.

Parameters

X	matrix that will hold the new updated solution to the linear system
x0	matrix that holds the previous solution to the linear system
arnoldi_dat	pointer to the ARNOLDI_DATA data structure

```
6.8.4.2 int arnoldi ( int(*)(const Matrix < double > &v, Matrix < double > &w, const void *data) matvec, int(*)(const Matrix < double > &b, Matrix < double > &p, const void *data) precon, Matrix < double > & r0, ARNOLDI_DATA * arnoldi_dat, const void * matvec_data, const void * precon_data)
```

Function to factor a linear operator into an orthonormal basis and upper Hessenberg matrix.

This function performs the Arnoldi algorithm to factor a linear operator into an orthonormal basis and upper Hessenberg matrix. Each orthonormal vector is formed using a Modified Gram-Schmidt procedure. When used in conjunction with GMRESLP, user may supply a preconditioning operator to improve convergence of the linear system. However, this function can be used by itself to factor the user's linear operator.

Parameters

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
r0	user supplied vector to serve as the first basis vector in the orthonormal basis
arnoldi_dat	pointer to the ARNOLDI_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.3 int gmresLeftPreconditioned (int(*)(const Matrix < double > &v, Matrix < double > &w, const void *data) matvec, int(*)(const Matrix < double > &b, Matrix < double > &b, const void *data) precon, Matrix < double > & b, GMRESLP_DATA * gmreslp_dat, const void * matvec_data, const void * precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with GMRESLP.

This function iteratively solves a non-symmetric, indefinite linear system using the Generalized Minimum RE ← Sidual method with Left Preconditioning (GMRESLP). It calls the Arnoldi algorithm to factor a linear operator into an

orthonormal basis and upper Hessenberg matrix, then uses that factorization to form an approximation to the linear system. Because this algorithm uses left-side preconditioning, it can only check the linear residuals at the outer iterations.

Parameters

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
gmreslp_dat	pointer to the GMRESLP_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.4 int fom (int(*)(const Matrix< double > &v, Matrix< double > &w, const void *data) matvec, int(*)(const Matrix< double > &b, Matrix< double > &b, GMRESLP_DATA * gmreslp_dat, const void * matvec_data, const void * precon_data)

Function to directly solve a non-symmetric, indefinite linear system with FOM.

This function directly solves a non-symmetric, indefinite linear system using the Full Orthogonalization Method (F← OM). This algorithm is exactly equivalent to GMRESLP without restarting. Therefore, it uses the GMRESLP_DATA structure and calls the GMRESLP algorithm without using restarts. As a result, it never checks linear residuals. However, this should give the exact solution upon completion, assuming the linear operator is not singular.

Parameters

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
gmreslp_dat	pointer to the GMRESLP_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

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This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and

anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

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6.8.4.5 int gmresRightPreconditioned (int(*)(const Matrix< double > &v, Matrix< double > &w, const void *data) matvec, int(*)(const Matrix< double > &b, Matrix< double > &b, const void *data) precon, Matrix< double > & b, GMRESRP_DATA * gmresrp_dat, const void * matvec_data, const void * precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with GMRESRP.

This function iteratively solves a non-symmetric, indefinite linear system using the Generalized Minimum RESidual method with Right Preconditioning (GMRESRP). Because this algorithm uses right preconditioning, it is able to check the linear residuals at both the outer and inner iterations. This may be much for efficient compared to $G \leftarrow MRESLP$. In order to check inner residuals, this algorithm has to perform it's own internal Modified Gram-Schmidt procedure and will not call the Arnoldi algorithm.

Parameters

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
gmresrp_dat	pointer to the GMRESRP_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.6 int pcg (int(*)(const Matrix < double > &p, Matrix < double > &Ap, const void *data) matvec, int(*)(const $\textbf{Matrix} < \textbf{double} > \textbf{\&r}, \textbf{Matrix} < \textbf{double} > \textbf{\&z}, \textbf{const void} * \textbf{data}) \textit{precon}, \textbf{ Matrix} < \textbf{double} > \textbf{\&} \textit{b}, \textbf{ PCG_DATA} * \textbf{Acceptable} = \textbf{Accep$ pcg_dat, const void * matvec_data, const void * precon_data)

Function to iteratively solve a symmetric, definite linear system with PCG.

This function iteratively solves a symmetric, definite linear system using the Preconditioned Conjugate Gradient (PCG) method. The PCG algorithm is optimal in terms of efficiency and residual reduction, but only if the linear system is symmetric. PCG will fail if the linear operator is non-symmetric!

Parameters

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
pcg_dat	pointer to the PCG_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix < double > & b, Matrix < double > & Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.7 int bicgstab (int(*)(const Matrix < double > &p, Matrix < double > &Ap, const void *data) matvec, int(*)(const Matrix < double > &r, Matrix < double > &z, const void *data) precon, Matrix < double > & b, BiCGSTAB_DATA * bicg_dat, const void * matvec_data, const void * precon_data)

Function to iteratively solve a non-symmetric, definite linear system with BiCGSTAB.

This function iteratively solves a non-symmetric, definite linear system using the Bi-Conjugate Gradient STABilized (BiCGSTAB) method. This is a highly efficient algorithm for solving non-symmetric problems, but will occassionally breakdown and fail. Most common failures are caused by poor preconditioning. Works very well for grid-based linear systems.

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
bicg_dat	pointer to the BiCGSTAB_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
С орредео Н <u>р</u> у <i>Ф</i> руудс	nuser supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

.....

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix < double > & b, Matrix < double > & Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.8 int cgs (int(*)(const Matrix < double > &p, Matrix < double > &Ap, const void *data) matvec, int(*)(const Matrix < double > &r, Matrix < double > &z, const void *data) precon, Matrix < double > & b, CGS_DATA * cgs_dat, const void * matvec_data, const void * precon_data)

Function to iteratively solve a non-symmetric, definite linear system with CGS.

This function iteratively solves a non-symmetric, definite linear system using the Conjugate Gradient Squared (CGS) method. This is an extremely efficient algorithm for solving non-symmetric problems, but will often breakdown and fail. Most common failures are caused by poor or no preconditioning. Works very will for grid-based linear systems.

Parameters

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
cgs_dat	pointer to the CGS_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.9 int operatorTranspose (int(*)(const Matrix < double > &v, Matrix < double > &Av, const void *data) matvec,

Matrix < double > & r, Matrix < double > & u, OPTRANS_DATA * transpose_dat, const void * matvec_data)

Function that is used to perform transposition of a linear operator and results in a new vector A^T*r=u.

This function takes a user supplied linear operator and forms the result of that operator transposed and multiplied by a given vector r ($A^T*r=u$). Transposition is accomplised by reordering the transpose operator and multiplying the non-transposed operator by a complete set of orthonormal vectors. The end result gives the ith component of the vector u for each operation ($u_i = r^T*A*i$). Here, i is a vector made from the ith column of the identity matrix. If the linear system if sufficiently large, then this operation may take some time.

Parameters

matvec	user supplied linear operator given as an int function
r	vector to be multiplied by the transpose of the operator
и	vector to store the result of the operator transposition (u=A^T*r)
transpose_dat	pointer to the OPTRANS_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

6.8.4.10 int gcr (int(*)(const Matrix< double > &x, Matrix< double > &Ax, const void *data) matvec, int(*)(const Matrix< double > &r, Matrix< double > &Mr, const void *data) precon, Matrix< double > & b, GCR_DATA * gcr_dat, const void * matvec_data, const void * precon_data)

Function to iteratively solve a non-symmetric, definite linear system with GCR.

This function iteratively solves a non-symmetric, definite linear system using the Generalized Conjugate Residual (GCR) method. Similar to GMRESRP, this algorithm will construct a growing orthonormal basis set that will eventually form the exact solution to the linear system. However, this algorithm is less efficient than GMRESRP and can suffer breakdowns if the linear system is indefinite.

matvec	user supplied linear operator given as an int function
precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
gcr_dat	pointer to the GCR_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.11 int gmresrPreconditioner (const Matrix < double > & r, Matrix < double > & Mr, const void * data)

Function used in conjunction with GMRESR to apply GMRESRP iterations as a preconditioner.

This function is required to take the form of the user supplied preconditioning functions for other iterative methods. However, it cannot be used in conjunction with any other Krylov method. It is only called by the GMRESR function when the preconditioner needs to be applied.

Parameters

r	vector supplied to the preconditioner to operate on
Mr	vector to hold the result of the preconditioning operation
data	void pointer to the GMRESR_DATA data structure

6.8.4.12 int gmresr (int(*)(const Matrix< double > &x, Matrix< double > &Ax, const void *data) matvec, int(*)(const Matrix< double > &r, Matrix< double > &Mr, const void *data) terminal_precon, Matrix< double > & b, GMRESR DATA * gmresr dat, const void * matvec data, const void * term precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with GMRESR.

This function iteratively solves a non-symmetric, indefinite linear system using the Generalized Minimum RE← Sidual Recursive (GMRESR) method. This algorithm actually uses GCR at the outer iterations, but stabilizes GCR with GMRESRP inner iterations to implicitly form a variable preconditioner to the linear system. As such, this is one of only two methods that inherently includes preconditioning (the other is KMS), without any user supplied preconditioning operator. However, this algorithms is signficantly more computationally expensive than GCR or GMRESRP separately. It should only be used for solving very large or very hard to solve linear systems.

matvec	user supplied linear operator given as an int function
terminal_precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
gmresr_dat	pointer to the GMRESR_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
term_precon_data	user supplied void pointer to a data structure needed for the precondtioning operator

Note

```
int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)
```

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*terminal precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.13 int kmsPreconditioner (const Matrix < double > & r, Matrix < double > & Mr, const void * data)

Preconditioner function for the Krylov Multi-Space.

This function is required to take the form of the user supplied preconditioning functions for other iterative methods. However, it cannot be used in conjunction with any other Krylov method. It is only called by the KMS function when the preconditioner needs to be applied.

Parameters

r	vector supplied to the preconditioner to operate on
Mr	vector to hold the result of the preconditioning operation
data	void pointer to the KMS_DATA data structure

6.8.4.14 int krylovMultiSpace (int(*)(const Matrix < double > &x, Matrix < double > &Ax, const void *data) matvec, int(*)(const Matrix< double > &r, Matrix< double > &Mr, const void *data) terminal_precon, Matrix< double > & b, KMS_DATA * kms_dat, const void * matvec_data, const void * term_precon_data)

Function to iteratively solve a non-symmetric, indefinite linear system with KMS.

This function iteratively solves a non-symmetric, indefinite linear system using the Krylov Multi-Space (KMS) method. This algorithm uses GMRESRP at both outer and inner iterations to implicitly form a variable preconditioner to the linear system. As such, this is one of only two methods that inherently includes preconditioning, without any user supplied preconditioning operator (the other being GMRESR). The advantage to this method over GMRESR is that this method is GMRES at its core, and will therefore never breakdown or need to be stabilized. Additionally, you can call this method and set it's max_level parameter (see KMS_DATA) to 0, which will make this algorithm exactly equal to GMRESRP. If the max_level is set to 1, then this algorithm is exactly FGMRES (Saad, 1993) with the GMRES algorithm as a preconditioner. However, you can set max_level higher to precondition the preconditioners with more preconditioners. Thus creating a method with any desired complexity or rate of convergence.

Parameters

matvec	user supplied linear operator given as an int function
terminal_precon	user supplied preconditioning operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
kms_dat	pointer to the KMS_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator
term_precon_data	user supplied void pointer to a data structure needed for the precondtioning operator
matvec_data	user supplied void pointer to a data structure needed for the linear of

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Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

int (*terminal precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the above linear operator function and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the original linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.15 int QRsolve (int(*)(const Matrix < double > &x, Matrix < double > &Ax, const void *data) matvec, Matrix < double > & b, QR_DATA * qr_dat , const void * $matvec_data$)

Function to solve a dense linear operator system using QR factorization.

This function is used to solve a dense linear system using QR factorization. It should only be used if iterative methods are unstable or if the linear system is very dense. There will likely be memory limitations to using this method, since it is assumed that the matrix/operator is dense. This method may also be less efficient because it has to extract the matrix elements from the linear operator. So if the linear operator is large, then the setup cost for this method is high.

Factorization is carried out using Householder Reflections. Each reflection matrix is iteratively applied to the operator and the vector b to convert the linear system to upper triangular. Then, the system is solved using backwards substitution.

Parameters

matvec	user supplied linear operator given as an int function
b	matrix of boundary conditions in the linear system Ax=b
qr_dat	pointer to the QR_DATA data structure
matvec_data	user supplied void pointer to a data structure needed for the linear operator

Note

int (*matvec) (const Matrix<double>& v, Matrix<double> &Av, const void *data)

This is a user supplied function for a linear operator. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix v that will act on the linear operator a modified the matrix entries of Av to form the result of a matrix-vector product. Void pointer data is used to pass any user data structure that the function may need in order to perform the linear operation.

6.8.4.16 int picard ($int(*)(const\ Matrix< double> &x,\ Matrix< double> &x,\ const\ void *data) res,\ int(*)(const\ Matrix< double> &x,\ int(*)(const\ Ma$

Function to iteratively solve a non-linear system using the Picard or Fixed-Point method.

This function iteratively solves a non-linear system using the Picard method. User supplies a residual function and a weak solution form function. The weak form function is used to approximate the next solution vector for the non-linear system and the residual function is used to determine convergence. User also supplies an initial guess to the non-linear system as a matix x, which will also be used to store the solution. This algorithm is very simple and may not be sufficient to solve complex non-linear systems.

Parameters

res	user supplied function for the non-linear residuals of the system
evalx	user supplied function for the weak form to estimate the next solution
X	user supplied matrix holding the initial guess to the non-linear system
picard_dat	pointer to the PICARD_DATA data structure
res_data	user supplied void pointer to a data structure used for residual evaluations
evalx_data	user supplied void pointer to a data structure used for evaluation of weak form

Note

int (*res) (const Matrix<double>& x, Matrix<double> &F, const void *data)

This is a user supplied function for the non-linear residuals. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix x representing the current non-linear variables. Those variables are used to evaluate the users functions and return the residuals in the matrix F. The void pointer data is a data structure provided by the user to hold information the function may need in order to form the residuals.

int (*evalx) (const Matrix<double>& x0, Matrix<double> &x, const void *data)

This is a user supplied function to approximate the next solution vector x based on the previous solution vector x0. The x0 matrix is passed to this function and must be used to edit the entries of x based on the weak form of the problem. The user is free to define any weak form approximation. Void pointer data is the users data structure that may be used to pass additional information into this function in order to evaluate the weak form.

Example Residual: $F(x) = x^2 + x - 1$ Goal is to make this function equal zero

Example Weak Form: $x = 1 - x0^2$ Rearrage residual to form a weak solution

6.8.4.17 int jacvec (const Matrix < double > & v, Matrix < double > & Jv, const void * data)

Function to form a linear operator of a Jacobian matrix used along with the PJFNK method.

This function is used in conjunction with the PJFNK routine to form a linear operator that a Krylov method can operate on. This linear operator is formed from the current residual vector of the non-linear iteration in PJFNK using a finite difference approximation.

Jacobian Linear Operator: J*v = (F(x k + eps*v) - F(x k)) / eps

V	vector to be multiplied by the Jacobian matrix
Jv	storage vector for the result of the Jacobi-vector product
data	void pointer to the PJFNK_DATA data structure holding solver information

6.8.4.18 int backtrackLineSearch (int(*)(const Matrix< double > &x, Matrix< double > &F, const void *data)

feval, Matrix< double > & Fkp1, Matrix< double > & xkp1, Matrix< double > & pk, double normFk,

BACKTRACK DATA * backtrack_dat, const void * feval_data)

Function to perform a Backtracking Line Search operation to smooth out convergence of PJFNK.

This function performs a simple backtracking line search operation on the residuals from the PJFNK method. The step size of the non-linear iteration is checked against a level of tolerance for residual reduction, then adjusted down if necessary. This method always starts out with the maximum allowable step size. If the largest step size is fine, then the algorithm does nothing. Otherwise, it iteratively adjusts the step size down, until a suitable step is found. In the case that no suitable step is found, this algorithm will report failure to the PJFNK method and PJFNK will decide whether to continue trying to find a global minimum or report that it is stuck in a local minimum.

Parameters

feval	user supplied residual function for the non-linear system
Fkp1	vector holding the residuals for the next non-linear step
xkp1	vector holding the solution for the next non-linear step
pk	vector holding the current non-linear search direction
normFk	value of the current non-linear residual
backtrack_dat	pointer to the BACKTRACK_DATA data structure
feval_data	user supplied void pointer to the data structure needed for residual evaluation

Note

int (*feval) (const Matrix<double>& x, Matrix<double> &F, const void *data)

This is a user supplied function for the non-linear residuals. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix x representing the current non-linear variables. Those variables are used to evaluate the users functions and return the residuals in the matrix F. The void pointer data is a data structure provided by the user to hold information the function may need in order to form the residuals.

6.8.4.19 int pjfnk (int(*)(const Matrix < double > &x, Matrix < double > &F, const void *data) res, int(*)(const Matrix < double > &r, Matrix < double > &x, PJFNK_DATA * pjfnk_dat, const void * res_data, const void * precon_data)

Function to perform the PJFNK algorithm to solve a non-linear system of equations.

This function solves a non-linear system of equations using the Preconditioned Jacobian- Free Newton-Krylov (P← JFNK) algorithm. Each non-linear step of this method results in a linear sub-problem that is solved iteratively with one of the Krylov methods in the krylov_method enum. User must supplied a residual function that computes the non-linear residuals of the system given the current state of the variables x. Additionally, the user must also supplied an initial guess to the non-linear system. Optionally, the user may supply a preconditioning function for the linear sub-problem.

Basic Steps: (i) Calc $F(x_k)$, (ii) Solve $J(x_k)*s_k=-F(x_k)$ for s_k , (iii) Form $x_kp1=x_k+s_k$

Parameters

res	user supplied residual function for the non-linear system
precon	user supplied preconditioning function for the linear sub-problems
X	user supplied initial guess and storage location of the solution
pjfnk_dat	pointer to the PJFNK_DATA data structure
res_data	user supplied void pointer to data structure used in residual function
precon_data	user supplied void pointer to data structure used in preconditioning function

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Note

int (*res) (const Matrix<double>& x, Matrix<double> &F, const void *data)

This is a user supplied function for the non-linear residuals. User's function must return an int of 0 upon success and anything else denotes a failure. The function accepts a matrix x representing the current non-linear variables. Those variables are used to evaluate the users functions and return the residuals in the matrix

F. The void pointer data is a data structure provided by the user to hold information the function may need in order to form the residuals.

int (*precon) (const Matrix<double>& b, Matrix<double> &Mb, const void *data)

This is a user supplied function for a preconditioning operator. It has the same form as the linear operators from the Krylov methods and should have all the same properties. The only difference is that this function must form an approximate matrix inversion on the jacvec linear operator and modify the entries of Mb to represent the result of that approximate matrix inversion. The matrix b is given as the vector that this operator is acting on and the void pointer data is for any user data structure that the operator may need.

6.8.4.20 int Numerical Jacobian (int(*)(const Matrix < double > &x, Matrix < double > &F, const void *user_data) Func, const Matrix < double > & x, Matrix < double > & J, int Nx, int Nf, NUM_JAC_DATA * jac_dat , const void * $user_data$)

Function to form a full numerical Jacobian matrix from a given non-linear function.

This function uses finite differences to form a full rank Jacobian matrix for a user supplied non-linear function. The Jacobian matrix will be formed at the current state of the non-linear variables x and stored in a full matrix J. Integers Nx and Nf are used to determine the size of the Jacobian matrix.

Parameters

Func	user supplied function for evaluation of the non-linear system
X	matrix holding the current value of the non-linear variables
J	matrix that will store the numerical Jacobian result
Nx	number of non-linear variables in the system
Nf	number of non-linear functions in the system
jac_dat	pointer to the NUM_JAC_DATA data structure
user_data	user supplied void pointer to a data structure used in the non-linear function

6.8.4.21 int LARK_TESTS ()

Function that runs a variety of tests on all the functions in LARK.

This function runs a variety of tests on the linear and non-linear methods developed in LARK. It can be called from the UI.

6.9 macaw.h File Reference

MAtrix CAlculation Workspace.

```
#include <stdio.h>
#include <math.h>
#include <iostream>
#include <fstream>
#include <stdlib.h>
#include <vector>
#include <time.h>
#include <float.h>
#include <string>
#include <exception>
#include "error.h"
```

Classes

class Matrix< T >

Templated C++ Matrix Class Object (click Matrix to go to function definitions)

Macros

#define M_PI 3.14159265358979323846264338327950288
 Value of PI with double precision.

Functions

• int MACAW_TESTS ()

Function to run the MACAW tests.

6.9.1 Detailed Description

MAtrix CAlculation Workspace.

macaw.cpp

This is a small C++ library that facilitates the use and construction of real matrices using vector objects. The Matrix class is templated so that users are able to work with matrices of any type including, but not limited to: (i) doubles, (ii) ints, (iii) floats, and (iv) even other matrices! Routines and functions are defined for Dense matrix operations. As a result, we typically only use Column Matrices (or Vectors) when doing any actual simulations. However, the development of this class was integral to the development and testing of the Sparse matrix operators in lark.h.

While the primary goal of this object was to define how to operate on real matrices, we could extend this idea to complex matrices as well. For this, we could develop objects that represent imaginary and complex numbers and then create a Matrix of those objects. For this reason, the matrix operations here are all templated to abstract away the specificity of the type of matrix being operated on.

Author

Austin Ladshaw

Date

01/07/2014

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6.9.2 Macro Definition Documentation

6.9.2.1 #define M_PI 3.14159265358979323846264338327950288

Value of PI with double precision.

6.9.3 Function Documentation

```
6.9.3.1 int MACAW_TESTS ( )
```

Function to run the MACAW tests.

This function is callable from the UI and is used to run several algorithm tests for the Matrix objects. This test should never report any errors.

6.10 magpie.h File Reference

Multicomponent Adsorption Generalized Procedure for Isothermal Equilibria.

```
#include "lmcurve.h"
#include <stdio.h>
#include <math.h>
#include <iostream>
#include <fstream>
#include <stdlib.h>
#include <vector>
#include <time.h>
#include <string>
#include "error.h"
#include "lark.h"
```

Classes

struct GSTA_DATA

GSTA Data Structure.

struct mSPD_DATA

MSPD Data Structure.

struct GPAST_DATA

GPAST Data Structure.

struct SYSTEM_DATA

System Data Structure.

struct MAGPIE_DATA

MAGPIE Data Structure.

Macros

#define DBL_EPSILON 2.2204460492503131e-016

Machine precision value used for approximating gradients.

#define Z 10.0

Surface coordination number used in the MSPD activity model.

#define A 3.13E+09

Corresponding van der Waals standard area for our coordination number (cm²/mol)

#define V 18.92

Corresponding van der Waals standard volume for our coordination number (cm^{\(\circ\)} 3/mol)

• #define Po 100.0

Standard State Pressure - Units: kPa.

#define R 8.3144621

Gas Constant - Units: J/(K*mol) = kB * Na.

#define Na 6.0221413E+23

Avagadro's Number - Units: molecules/mol.

#define kB 1.3806488E-23

Boltzmann's Constant - Units: J/K.

#define shapeFactor(v i) (((Z - 2) * v i) / (Z * V)) + (2 / Z)

This macro replaces all instances of shapeFactor(#) with the following single line calculation.

#define InKo(H, S, T) -(H / (R * T)) + (S / R)

This macro calculates the natural log of the dimensionless isotherm parameter.

#define He(qm, K1, m) (qm * K1) / (m * Po)

This macro calculates the Henry's Coefficient for the ith component.

Functions

• double qo (double po, const void *data, int i)

Function computes the result of the GSTA isotherm for the ith species.

double dq_dp (double p, const void *data, int i)

Function computes the derivative of the GSTA model with respect to partial pressure.

double q_p (double p, const void *data, int i)

Function computes the ratio between the adsorbed amount and partial pressure for the GSTA isotherm.

double PI (double po, const void *data, int i)

Function computes the spreading pressure integral of the ith species.

double Qst (double po, const void *data, int i)

Function computes the heat of adsorption based on the ith species GSTA parameters.

double eMax (const void *data, int i)

Function to approximate the maximum lateral energy term for the ith species.

• double Inact mSPD (const double *par, const void *data, int i, volatile double PI)

Function to evaluate the MSPD activity coefficient for the ith species.

double grad_mSPD (const double *par, const void *data, int i)

Function to approximate the derivative of the MSPD activity model with spreading pressure.

double qT (const double *par, const void *data)

Function to calculate the total adsorbed amount (mol/kg) for the mixed surface phase.

void initialGuess_mSPD (double *par, const void *data)

Function to provide an initial guess to the unknown parameters being solved for in GPAST.

void eval po PI (const double *par, int m dat, const void *data, double *fvec, int *info)

Function used with Imfit to evaluate the reference state pressure of a species based on spreading pressure.

void eval_po_qo (const double *par, int m_dat, const void *data, double *fvec, int *info)

Function used with Imfit to evaluate the reference state pressure of a species based on that species isotherm.

void eval_po (const double *par, int m_dat, const void *data, double *fvec, int *info)

Function used with Imfit to evaluate the reference state pressure of a species based on a sub-system.

• void eval_eta (const double *par, int m_dat, const void *data, double *fvec, int *info)

Function used with Imfit to evaluate the binary interaction parameters for each unique species pair.

void eval_GPAST (const double *par, int m_dat, const void *data, double *fvec, int *info)

Function used with Imfit to solve the GPAST system of equations.

int MAGPIE (const void *data)

Function to call all sub-routines to solve a MAGPIE/GPAST problem at a given temperature and pressure.

int MAGPIE_SCENARIOS (const char *inputFileName, const char *sceneFileName)

Function to perform a series of MAGPIE simulations based on given input files.

6.10.1 Detailed Description

Multicomponent Adsorption Generalized Procedure for Isothermal Equilibria.

magpie.cpp

This file contains all functions and routines associated with predicting isothermal adsorption equilibria from only single component isotherm information. The basis of the model is the Adsorbed Solution Theory developed by Myers and Prausnitz (1965). Added to that base model is a procedure by which we can predict the non-idealities present at the surface phase by solving a closed system of equations involving the activity model.

For more details on this procedure, check out our publication in AIChE where we give a fully feature explaination of our Generalized Predictive Adsorbed Solution Theory (GPAST).

Reference: Ladshaw, A., Yiacoumi, S., and Tsouris, C., "A generalized procedure for the prediction of multicomponent adsorption equilibria", AIChE J., vol. 61, No. 8, p. 2600-2610, 2015.

MAGPIE represents a special case of the more general GPAST procedure, wherin the isotherm for each species is respresent by the GSTA isotherm (see gsta_opt.h) and the activity model for non-ideality at the adsorbent surface is a Modified Spreading Pressure Dependent (MSPD) model. See the above paper reference for more details.

Author

Austin Ladshaw

Date

12/17/2013

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6.10.2 Macro Definition Documentation

6.10.2.1 #define DBL_EPSILON 2.2204460492503131e-016

Machine precision value used for approximating gradients.

6.10.2.2 #define Z 10.0

Surface coordination number used in the MSPD activity model.

6.10.2.3 #define A 3.13E+09

Corresponding van der Waals standard area for our coordination number (cm^2/mol)

6.10.2.4 #define V 18.92

Corresponding van der Waals standard volume for our coordination number (cm^3/mol)

6.10.2.5 #define Po 100.0

Standard State Pressure - Units: kPa.

6.10.2.6 #define R 8.3144621

Gas Constant - Units: J/(K*mol) = kB * Na.

6.10.2.7 #define Na 6.0221413E+23

Avagadro's Number - Units: molecules/mol.

6.10.2.8 #define kB 1.3806488E-23

Boltzmann's Constant - Units: J/K.

6.10.2.9 #define shapeFactor(v_i)(((Z-2) * v_i)/(Z * V))+(2/Z)

This macro replaces all instances of shapeFactor(#) with the following single line calculation.

6.10.2.10 #define lnKo(H, S, T)-(H/(R * T))+(S/R)

This macro calculates the natural log of the dimensionless isotherm parameter.

6.10.2.11 #define He(qm, K1, m)(qm * K1)/(m * Po)

This macro calculates the Henry's Coefficient for the ith component.

6.10.3 Function Documentation

6.10.3.1 double qo (double po, const void * data, int i)

Function computes the result of the GSTA isotherm for the ith species.

This function just computes the result of the GSTA isotherm model for the ith species given the partial pressure po.

Parameters

ро	partial pressure in kPa at which to evaluate the GSTA model
data	void pointer to the MAGPIE_DATA data structure
i	index of the gas species for which the GSTA model is being evaluated

6.10.3.2 double dq_dp (double p, const void * data, int i)

Function computes the derivative of the GSTA model with respect to partial pressure.

This function just computes the result of the derivative of GSTA isotherm model for the ith species at the given the partial pressure p.

Parameters

р	partial pressure in kPa at which to evaluate the GSTA model
data	void pointer to the MAGPIE_DATA data structure
i	index of the gas species for which the GSTA model is being evaluated

6.10.3.3 double q_p (double p, const void * data, int i)

Function computes the ratio between the adsorbed amount and partial pressure for the GSTA isotherm.

This function just computes the ratio between the adsorbed amount q (mol/kg) and the partial pressure p (kPa) at the given partial pressure. If p == 0, then this function returns the Henry's Law constant for the isotherm of the ith species.

Parameters

р	partial pressure in kPa at which to evaluate the GSTA model
data	void pointer to the MAGPIE_DATA data structure
i	index of the gas species for which the GSTA model is being evaluated

6.10.3.4 double PI (double po, const void * data, int i)

Function computes the spreading pressure integral of the ith species.

This function uses an analytical solution to the spreading pressure integral with the GSTA isotherm to evaluate and return the value computed by that integral equation.

Parameters

ро	partial pressure in kPa at which to evaluate the lumped spreading pressure
data	void pointer to the MAGPIE_DATA data structure
i	index of the gas species for which the GSTA model is being evaluated

6.10.3.5 double Qst (double po, const void * data, int i)

Function computes the heat of adsorption based on the ith species GSTA parameters.

This function computes the isosteric heat of adsorption (J/mol) for the GSTA parameters of the ith species.

Parameters

ро	partial pressure in kPa at which to evaluate the heat of adsorption
data	void pointer to the MAGPIE_DATA data structure
i	index of the gas species for which the GSTA model is being evaluated

6.10.3.6 double eMax (const void * data, int i)

Function to approximate the maximum lateral energy term for the ith species.

The function attempts to approximate the maximum lateral energy term for the ith species. This is not a true maximum, but a cheaper estimate. Value being computed is used to shift the geometric mean and formulate the average cross-lateral energy term between species i and j.

6.10.3.7 double lnact_mSPD (const double * par, const void * data, int i, volatile double PI)

Function to evaluate the MSPD activity coefficient for the ith species.

This function will return the natural log of the ith species activity coefficient using the Modified Spreading Pressure Dependent (MSPD) activity model. The par argument holds the variable values being solved for by GPAST and their contents will change depending on whether we are doing a forward or reverse evaluation. This function should not be called by the user and will only be called when needed in the GPAST routine.

Parameters

par	list of parameters representing variables to be solved for in GPAST
data	void pointer for the MAGPIE_DATA data structure
i	ith species that we want to calculate the activity coefficient for
PI	lumped spreading pressure term used in gradient estimations

6.10.3.8 double grad_mSPD (const double * par, const void * data, int i)

Function to approximate the derivative of the MSPD activity model with spreading pressure.

This function returns a 2nd order, finite different approximation of the derivative of the MSPD activity model with the spreading pressure. The par argument will either hold the current iterates estimate of spreading pressure or should be passed as null. User does not need to call this function. GPAST will call automatically when needed.

Parameters

par	list of parameters representing variables to be solved for in GPAST
data	void pointer for the MAGPIE_DATA data structure
i	ith species for which we will approximate the activty model gradient

6.10.3.9 double qT (const double * par, const void * data)

Function to calculate the total adsorbed amount (mol/kg) for the mixed surface phase.

This function will uses the obtained system parameters from par and estimate the total amount of gases adsorbed to the surface in mol/kg. The user does not need to call this function, since this result will be stored in the SYST DATA structure.

Parameters

par	list of parameters representing variables to be solved for in GPAST
data	void pointer for the MAGPIE_DATA data structure

6.10.3.10 void initialGuess_mSPD (double * par, const void * data)

Function to provide an initial guess to the unknown parameters being solved for in GPAST.

This function intends to provide an initial guess for the unknown values being solved for in the GPAST system. Depending on what type of solve is requested, this algorithm will provide a guess for the adsorbed or gas phase composition.

Parameters

par	list of parameters representing variables to be solved for in GPAST
data	void pointer for the MAGPIE_DATA data structure

6.10.3.11 void eval_po_PI (const double * par, int m_dat, const void * data, double * fvec, int * info)

Function used with Imfit to evaluate the reference state pressure of a species based on spreading pressure.

This function is used inside of the MSPD activity model to calculate the reference state pressure of a particular species at a given spreading pressure for the system. User does not need to call this function. GPAST will call automatically when needed.

Parameters

par	list of parameters representing variables to be solved for in GPAST
m_dat	number of functions/variables in the GPAST system of equations
data	void pointer for the MAGPIE_DATA data structure
fvec	list of residuals formed by the functions in GPAST
info	integer flag variable used in the Imfit routine

6.10.3.12 void eval_po_qo (const double * par, int m_d at, const void * data, double * fvec, int * info)

Function used with Imfit to evaluate the reference state pressure of a species based on that species isotherm.

This function is used to evaluate the partial pressure or reference state pressure for a particular species given single-component adsorbed amount. User does not need to call this function. GPAST will call automatically when needed.

par	list of parameters representing variables to be solved for in GPAST
m_dat	number of functions/variables in the GPAST system of equations

Parameters

data	void pointer for the MAGPIE_DATA data structure
fvec	list of residuals formed by the functions in GPAST
info	integer flag variable used in the Imfit routine

6.10.3.13 void eval_po (const double * par, int m_dat , const void * data, double * fvec, int * info)

Function used with Imfit to evaluate the reference state pressure of a species based on a sub-system.

This function is used to approximate reference state pressures based on the spreading pressure of a sub-system in GPAST. The sub-system will be one of the unique binary systems that exist in the overall mixed gas system. User does not need to call this function. GPAST will call automatically when needed.

Parameters

par	list of parameters representing variables to be solved for in GPAST
m_dat	number of functions/variables in the GPAST system of equations
data	void pointer for the MAGPIE_DATA data structure
fvec	list of residuals formed by the functions in GPAST
info	integer flag variable used in the Imfit routine

6.10.3.14 void eval_eta (const double * par, int m_dat, const void * data, double * fvec, int * info)

Function used with Imfit to evaluate the binary interaction parameters for each unique species pair.

This function is used to estimate the binary interaction parameters for all species pairs in a given sub-system. Those parameters are then stored for later used when evaluating the activity coefficients for the overall mixture. User does not need to call this function. GPAST will call automatically when needed.

Parameters

par	list of parameters representing variables to be solved for in GPAST
m_dat	number of functions/variables in the GPAST system of equations
data	void pointer for the MAGPIE_DATA data structure
fvec	list of residuals formed by the functions in GPAST
info	integer flag variable used in the Imfit routine

6.10.3.15 void eval_GPAST (const double * par, int m_{-} dat, const void * data, double * fvec, int * info)

Function used with Imfit to solve the GPAST system of equations.

This function is used after having calculated and stored all necessary information to solve a closed form GPAST system of equations. User does not need to call this function. GPAST will call automatically when needed.

par	list of parameters representing variables to be solved for in GPAST
m_dat	number of functions/variables in the GPAST system of equations

Parameters

data	void pointer for the MAGPIE_DATA data structure
fvec	list of residuals formed by the functions in GPAST
info	integer flag variable used in the Imfit routine

6.10.3.16 int MAGPIE (const void * data)

Function to call all sub-routines to solve a MAGPIE/GPAST problem at a given temperature and pressure.

This is the function that a typical user will want to incorporate into their own codes when evaluating adsorption of a gas mixture. Prior to calling this function, all required structures and information in the MAGPIE_DATA structure must have been properly initialized. After this function has completed it's operations, it will return an integer used to denote a success or failure of the routine. Integers 0, 1, 2, and 3 all denote success. Anything else is considered a failure.

To setup the MAGPIE_DATA structure correctly, you must reserve space for all vector objects based on the number of gas species in the mixture. In general, you only need to reserve space for the adsorbing species. However, you can also reserve space for non-adsorbing species, but you MUST give a gas/adsorbed mole fraction of the non-adsorbing species 0.0 so that the routine knows to ignore them (very important)!

After setting up the memory for the vector objects, you can intialize information specific to the simulation you want to request. The number of species (N), total pressure (PT) and gas temperature (T) must always be given. You can neglect the non-idealities of the surface phase by setting the Ideal bool to true. This will result in faster calculations, because MAGPIE will just revert down to the Ideal Adsorbed Solution Theory (IAST).

The Recover bool will denote whether we are doing a forward or reverse GPAST evaluation. Forward evaluation is for solving for the composition of the adsorbed phase given the composition of the gas phase (Recover = false). Reverse evaluation is for solve for the composition of the gas phase given the composition of the adsorbed phase (Recover = true).

For a reverse evaluation (Recover = true) you will also need to stipulate whether or not there is a carrier gas (Carrier = true or false). A carrier gas is considered any non-adsorbing species that may be present in the gas phase and contributing to the total pressure in the system.

The parameters that must be initialized for all species include all GSTA_DATA parameters and the van der Waals volume parameter (v) in the mSPD_DATA structure. For non-adsorbing species, you can ignore these parameters, but need to set the sites (m) from GSTA_DATA to 1. GPAST cannot run any evaluations without these parameters being set properly AND set in the same order for all species (i.e., make sure that gpast_dat[i].qmax corresponds to mspd_dat[i].v and so on).

Lastly, you need to give either the gas phase or adsorbed phase mole fractions, depending on whether you are going to run a forward or reverse evaluation, respectively. For a forward evaluation, provide the gas mole fractions (y) in GPAST_DATA for each species (non-adsorbing species should have this value set to 0.0). For a reverse evaluation, provide the adsorbed mole fractions (x) in GPAST_DATA for each species, as well as the total adsorbed amount (qT) in SYSTEM_DATA. Again, non-adsorbing species should have their respective phase mole fractions set to 0.0 to exclude them from the simulation. Additionally, if there are non-adsorbing species present, then the Carrier bool in SYSTEM_DATA must be set to true.

data	void pointer for the MAGPIE_DATA data structure holding all necessary information
------	---

```
6.10.3.17 int MAGPIE_SCENARIOS ( const char * inputFileName, const char * sceneFileName )
```

Function to perform a series of MAGPIE simulations based on given input files.

This function is callable from the UI and is used to perform a series of isothermal equilibria evaluations using the MAGPIE routines. There are two input files that must be provided: (i) inputFileName - containing parameter information for the species and (ii) sceneFileName - containing information for each MAGPIE simulation. Each of these files have a specific structure (see below). NOTE: this may change in future versions.

inputFileName Text File Structure:

```
Integer for Number of Adsorbing Species van der Waals Volume (cm^3/mol) of ith species GSTA adsorption capacity (mol/kg) of ith species Number of GSTA parameters of ith species Enthalpy (J/mol) of nth site [tab] Entropy of nth site (J/K/mol) of ith species (repeat above for all n sites in species i) (repeat above for all species i)
```

Example Input File:

```
5
17.1
5.8797
-20351.9 -81.8369
16.2
5.14934
-16662.7 -74.4766
19.7
9.27339
-46597.5 -53.6994
-125024 -221.073
-193619 -356.728
-272228 -567.459
13.25
4.59144
1
-13418.5 -84.888
18.0
10.0348
-20640.4 -72.6119
(The above input file gives the parameter information for 5 adsorbing species)
```

sceneFileName Text File Structure:

```
Integer Flag to mark Forward (0) or { Reverse (1) evaluations }

Number of Simulations to Run

Total Pressure (kPa) [tab] Temperature (K) { [tab] Total Adsorption (mol/kg) [tab] Carrier Gas Flag (0=false, 1=true) }

Gas/Adsorbed Mole Fractions for each species in the order given in prior file (tab separated) (repeat above for all simulations desired)

NOTE: only provide the Total Adsorption and Carrier Flag if doing Reverse evaluations!
```

Example Scenario File 1:

```
0
4
0.65 303.15
0.364 0.318 0.318
3.25 303.15
0.371 0.32 0.309
6.85 303.15
0.388 0.299 0.313
13.42 303.15
0.349 0.326 0.325
```

(The above scenario file is for 4 forward evaluations/simulations for a 3-adsorbing species system)

Example Scenario File 2:

```
1

4

0.65 303.15 5.4 0

0.364 0.318 0.318

3.25 303.15 7.7 0

0.371 0.32 0.309

6.85 303.15 9.8 0

0.388 0.299 0.313

13.42 303.15 10.4 0

0.349 0.326 0.325
```

(The above scenario file is for 4 reverse evaluations/simulations for a 3-adsorbing species system and no carrier gas)

6.11 mola.h File Reference

Molecule Object Library from Atoms.

```
#include <ctype.h>
#include "eel.h"
```

Classes

· class Molecule

C++ Molecule Object built from Atom Objects (click Molecule to go to function definitions)

Macros

- #define M_PI 3.14159
- #define SphereVolume(r) ((4.0/3.0)*M_PI*r*r*r)
- #define SphereArea(r) (4.0*M_PI*r*r)

Enumerations

enum valid_phase {
 SOLID, LIQUID, AQUEOUS, GAS,
 PLASMA, ADSORBED, OTHER }

Functions

• int MOLA_TESTS ()

Function to run the MOLA tests.

6.11.1 Detailed Description

Molecule Object Library from Atoms.

mola.cpp

This file contains a C++ Class for creating Molecule objects from the Atom objects that were defined in eel.←
h. Molecules can be created and registered from basic information or can be registered from a growing list of pre-registered molecules that are accessible by name/formula.

Registered Molecules are are known and defined prior to runtime. They have a charge, energy characteristics, phase, name, and formula that they are recongized by. The formula is used to create the atoms that they are made from. If some information is incomplete, it must be specified as to what information is missing (i.e. denote whether the formation energies are known).

Formation energies are used to determine stability/dissociation/acidity equilibrium constants during runtime. If the formation energies are unknown, then the equilibrium constants must be given to a reaction object on when it is initialized.

The molecule formula's are given as strings which are parsed in the constructor to determine what atoms from the EEL files will be registered and used. Note, you will be able to build molecules from an input file, but the library molecules here are ready to be used in applications and require no more input other that the molecule's formula.

List of Currently Registered Molecules

Ag (s)

Ag + (aq)

AgBr (s)

AgCI (s)

Agl (s)

Ag2S (s)

AgOH (aq)

Ag(OH)2 - (aq)

AgCl (aq)

AgCl2 - (aq)

Al (s)

Al 3+ (aq)

AIOH 2+ (aq)

AI(OH)2 + (aq)

AI(OH)3 (aq)

Al(OH)4 - (aq)

Al2O3 (s)

AIOOH (s)

Al(OH)3 (s)

Al2Si2(OH)4 (s)

As (s)

AsO4 3- (aq)

Ba 2+ (aq)

BaSO4 (s)

BaCO3 (s)

Be 2+ (aq)

- Be(OH)2 (s)
- Be3(OH)3 3+ (aq)
- B(OH)4 (aq)
- Br2 (I)
- Br2 (aq)
- Br (aq)
- BrO (aq)
- CO3 2- (aq)
- CI (aq)
- CaCl2 (aq)
- CaAl2Si2O8 (s)
- C (s)
- CO2 (g)
- CH4 (g)
- CH4 (aq)
- CH3OH (aq)
- CN (aq)
- CH3COOH (aq)
- CH3COO (aq)
- C2H5OH (aq)
- Ca 2+ (aq)
- CaOH + (aq)
- Ca(OH)2 (aq)
- Ca(OH)2 (s)
- CaCO3 (s)
- CaMg(CO3)2 (s)
- CaSiO3 (s)
- CaSO4 (s)
- CaSO4(H2O)2 (s)
- Ca5(PO4)3OH (s)
- Cd 2+ (aq)
- Cd(OH) + (aq)
- Cd(OH)3 (aq)
- Cd(OH)4 2- (aq)
- Cd(OH)2 (aq)
- CdO(s)
- Cd(OH)2 (s)
- CdCl + (aq)
- CdCl2 (aq)
- CdCl3 (aq)
- CdCO3 (s)
- Cl2 (g)
- Cl2 (aq)
- CIO (aq)
- CIO2 (aq)
- CIO2 (aq)
- CIO3 (aq)
- CIO4 (aq)
- Co(s)
- Co 2+ (aq)
- Co 3+ (aq)
- CoOH + (aq)
- Co(OH)2 (aq)
- Co(OH)3 (aq)
- Co(OH)2 (s)
- CoO (s)
- Co3O4 (s)
- Cr (s)

Cr 2+ (aq)

Cr 3+ (aq)

CrOH 2+ (aq)

Cr(OH)2 + (aq)

Cr(OH)3 (aq)

Cr(OH)4 - (aq)

Cr2O3 (s)

CrO4 2- (aq)

Cr2O7 2- (aq)

Cu (s)

Cu + (aq)

Cu 2+ (aq)

CuOH + (aq)

Cu(OH)2 (aq)

Cu(OH)3 - (aq)

Cu(OH)4 2- (aq)

CuS (s)

Cu2S (s)

CuO (s)

CuCO3Cu(OH)2 (s)

(CuCO3)2Cu(OH)2 (s)

F2 (g)

F - (aq)

Fe (s)

Fe 2+ (aq)

FeOH + (aq)

Fe(OH)2 (aq)

Fe(OH)3 - (aq)

Fe 3+ (aq)

FeOH 2+ (aq)

Fe(OH)2 + (aq)

Fe(OH)3 (aq)

Fe(OH)4 - (aq)

Fe2(OH)2 4+ (aq)

FeS2 (s)

FeO (s)

Fe(OH)2 (s)

Fe2O3 (s)

Fe3O4 (s)

FeOOH (s)

Fe(OH)3 (s)

FeCO3 (s)

Fe2SiO4 (s)

H2O (I)

H + (aq)

H2CO3 (aq)

HCO3 - (aq)

HNO3 (aq)

HCI (aq)

H3AsO4 (aq)

H2AsO4 - (aq)

HAsO4 2- (aq)

H2AsO3 - (aq) H3BO3 (aq)

HBrO (aq)

HCOOH (aq)

HCOO - (aq)

HCN (aq)

- HCIO (aq)
- HCoO2 (aq)
- HCrO4 (aq)
- HCuO2 (aq)
- HF (aq)
- HF2 (aq)
- H2 (g)
- H2 (aq)
- H2O2 (aq)
- HO2 (aq)
- H2O (g)
- Hg (I)
- Hg2 2+ (aq)
- Hg 2+ (aq)
- HgOH + (aq)
- Hg(OH)2 (aq)
- Hg(OH)3 (aq)
- Hg2Cl2 (s)
- HgO (s)
- HgS (s)
- Hgl2 (s)
- HgCl + (aq)
- HgCl2 (aq)
- HgCl3 (aq)
- HgCl4 2- (aq)
- HgOH + (aq)
- Hg(OH)2 (aq)
- HgO2 (aq)
- HIO (aq)
- HIO3 (aq)
- HNO2 (aq)
- HPO4 2- (aq)
- H2PO4 (aq)
- H3PO4 (aq)
- H2S (g)
- H2S (aq)
- HS (aq)
- HSO3 (aq)
- H2SO3 (aq)
- HSO4 (aq)
- H2SO4 (aq)
- HSeO3 (aq)
- H2SeO3 (aq)
- HSeO4 (aq)
- H4SiO4 (aq)
- HV2O5 (aq)
- H4VO4 + (aq)
- H3VO4 (aq)
- H2VO4 (aq)
- HVO4 2- (aq)
- H4VO4(C2O4)2 3- (aq)
- H4VO4C2O4 (aq)
- H2V10O28 4- (aq)
- HV10O28 5- (aq)
- HV2O7 3- (aq)
- 12 (s)
- 12 (aq)
- I (aq)

13 - (aq)

10 - (aq)

IO3 - (aq)

KAl3Si3O10(OH)2 (s)

K + (aq)

Mg(OH)2 (aq)

Mg5Al2Si3O10(OH)8 (s)

Mg (s)

Mg 2+ (aq)

MgOH + (aq)

Mg(OH)2 (s)

Mn (s)

Mn 2+ (aq)

Mn(OH)2 (s)

Mn3O4 (s)

MnOOH (s)

MnO2 (s)

MnCO3 (s)

MnS (s)

MnSiO3 (s)

NaHCO3 (aq)

Narioos (aq)

NaCO3 - (aq)

Na + (aq)

NaCl (aq)

NaOH (aq)

NO3 - (aq)

NH3 (aq)

NaAlSiO3O8 (s)

NH2CH2COOH (aq)

NH2CH2COO - (aq)

N2 (g)

N2O (g)

NH3 (g)

NH4 + (aq)

NO2 - (aq)

Ni 2+ (aq)

NiOH + (aq)

Ni(OH)2 (aq)

Ni(OH)3 - (aq)

NiO (s)

NiS (s)

OH - (aq)

O2 (g)

O2 (aq)

O3 (g)

P (s)

PO4 3- (aq)

Pb (s)

Pb 2+ (aq)

PbOH + (aq)

Pb(OH)2 (aq)

Pb(OH)3 - (aq)

Pb(OH)4 2- (aq) Pb(OH)2 (s)

PbO (s)

PbO2 (s)

Pb3O4 (s)

PbS (s)

- PbSO4 (s)
- PbCO3 (s)
- S (s)
- SO2 (g)
- SO3 (g)
- S 2- (aq)
- SO3 2- (aq)
- SO4 2- (aq)
- Se (s)
- SeO3 2- (aq)
- SeO4 2- (aq)
- Si (s)
- SiO2 (s)
- Sr 2+ (aq)
- SrOH + (aq)
- SrCO3 (s)
- SrSO4 (s)
- UO2 2+ (aq)
- UO2NO3 + (aq)
- UO2(NO3)2 (aq)
- UO2OH + (aq)
- UO2(OH)2 (aq)
- UO2(OH)3 (aq)
- UO2(OH)4 2- (aq)
- (UO2)2OH 3+ (aq)
- (UO2)2(OH)2 2+ (aq)
- (UO2)3(OH)4 2+ (aq)
- (UO2)3(OH)5 + (aq)
- (UO2)3(OH)7 (aq)
- (UO2)4(OH)7 + (aq)
- UO2CO3 (aq)
- UO2(CO3)2 2- (aq)
- UO2(CO3)3 4- (aq)
- UO2CI + (aq)
- UO2CI2 (aq)
- UO2Cl3 (aq)
- UO2SO4 (aq)
- UO2(SO4)2 2- (aq)
- VO 2+ (aq)
- VOOH + (aq)
- VO(OH)2 (s)
- V2O4 (s)
- (VO)2(OH)2 + (aq)
- VOF + (aq)
- VOF2 (aq)
- VOF3 (aq)
- VOF4 2- (aq)
- VOCI + (aq)
- VOSO4 (aq)
- VO(C2O4)2 2- (aq)
- VOOHC2O4 (aq)
- VOCH3COO + (aq)
- VO(CH3COO)2 (aq)
- VOCO3 (aq)
- VOOHCO3 (aq)
- V4O9 2- (aq)
- VO2 + (aq)
- VO4 3- (aq)

V2O5 (s) V10O28 6- (aq) V2O7 4- (aq) V4O12 4- (aq) VO2SO4 - (aq) VO2OHCO3 2- (aq) VO2(CO3)2 3- (aq) Zn (s) Zn 2+ (aq) ZnOH + (aq) Zn(OH)2 (aq) Zn(OH)3 - (aq) Zn(OH)4 2- (aq) Zn(OH)2 (s) ZnCl + (aq) ZnCl2 (aq) ZnCl3 - (aq) ZnCl4 2- (aq)

Those registered molecules follow a strict naming convention by which they can be recognized (see below)...

Naming Convention

ZnCO3 (s)

Plus (+) and minus (-) charges are denoted by the numeric value of the charge followed by a + or - sign, respectively (e.g. UO2(CO3)3 4- (aq))

The phase is always denoted last and will be marked as (I) for liquid, (s) for solid, (aq) for aqueous, and (g) for gas (see above).

When registering a molecule that is not in the library, you must also provide a linear formula during construction or registration. This is needed so that the string parsing is easier to handle when the molecule subsequently registers the necessary atoms. (e.g. UO2(CO3)3 = UO2C3O9 or UO11C3).

Author

Austin Ladshaw

Date

02/24/2014

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```
6.11.2 Macro Definition Documentation
```

6.11.2.1 #define M_PI 3.14159

6.11.2.2 #define SphereVolume(r) ((4.0/3.0)*M_PI*r*r*r)

6.11.2.3 #define SphereArea(r) (4.0*M_PI*r*r)

6.11.3 Enumeration Type Documentation

6.11.3.1 enum valid_phase

Enumerator

SOLID

LIQUID

AQUEOUS

GAS

PLASMA

ADSORBED

OTHER

6.11.4 Function Documentation

```
6.11.4.1 int MOLA_TESTS ( )
```

Function to run the MOLA tests.

This function is callable from the UI and is used to run several algorithm tests for the Molecule objects. This test should never report any errors.

6.12 monkfish.h File Reference

Multi-fiber wOven Nest Kernel For Interparticle Sorption History.

```
#include "dogfish.h"
```

Classes

struct MONKFISH_PARAM

Data structure for species specific information and parameters.

struct MONKFISH_DATA

Primary data structure for running MONKFISH.

Functions

• double default_porosity (int i, int I, const void *user_data)

Default porosity function for MONKFISH.

double default_density (int i, int I, const void *user_data)

Default density function for MONKFISH.

• double default_interparticle_diffusion (int i, int I, const void *user_data)

Default interparticle diffusion function.

• double default_monk_adsorption (int i, int I, const void *user_data)

Default adsorption strength function.

double default_monk_equilibrium (int i, int I, const void *user_data)

Default equilibirium adsorption function in mg/g.

• double default_monkfish_retardation (int i, int I, const void *user_data)

Default retardation coefficient function.

• double default_exterior_concentration (int i, const void *user_data)

Default exterior concentratio function.

double default_film_transfer (int i, const void *user_data)

Default film mass transfer function.

int setup_MONKFISH_DATA (FILE *file, double(*eval_porosity)(int i, int I, const void *user_data), double(*eval_density)(int i, int I, const void *user_data), double(*eval_ext_diff)(int i, int I, const void *user_data), double(*eval_retard)(int i, int I, const void *user_data), double(*eval_ext_film)(int i, const void *user_data), double(*eval_ext_film)(int i, const void *user_data), double(*dog_diffusion)(int i, int I, const void *user_data), double(*dog_ext_film)(int i, const void *user_data), double(*dog_surf_conc)(int i, const void *user_data), const void *user_data, MONKFISH_D ATA *monk dat)

Setup function to allocate memory and setup function pointers for the MONKFISH simulation.

• int MONKFISH TESTS ()

Function to run tests on the MONKFISH algorithms.

6.12.1 Detailed Description

Multi-fiber wOven Nest Kernel For Interparticle Sorption History.

monkfish.cpp

This file contains structures and functions associated with modeling the sorption characteristics of woven fiber bundles used to recover uranium from seawater. It is coupled with the DOGFISH kernel that determines the sorption of individual fibers. This kernel will resolve the interparticle diffusion between bundles of individual fibers in a woven ball-like domain.

Warning

Functions and methods in this file are still under construction.

Author

Austin Ladshaw

Date

04/14/2015

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6.12.2 Function Documentation

6.12.2.1 double default_porosity (int i, int I, const void * user_data)

Default porosity function for MONKFISH.

This function assumes a linear relationship between the maximum porosity at the center of the woven fibers and the minimum porosity at the edge of the woven fiber bundle.

Parameters

i	index for the ith adsorbing species	
1	index for the lth node in the domain	
user_data	pointer to the MONKFISH_DATA structure	

6.12.2.2 double default_density (int i, int l, const void * user_data)

Default density function for MONKFISH.

This function calls the porosity function and uses the single fiber density to provide an estimate of the bulk fiber density locally in the woven fiber bundle.

Parameters

i	index for the ith adsorbing species
1	index for the lth node in the domain
user_data	pointer to the MONKFISH_DATA structure

6.12.2.3 double default_interparticle_diffusion (int i, int I, const void * user_data)

Default interparticle diffusion function.

This function assumes that the interparticle diffusivity is a contant and returns that diffusivity multiplied by the domain porosity to form the effective diffusion coefficient in the domain.

Parameters

i	index for the ith adsorbing species
1	index for the lth node in the domain
user_data	pointer to the MONKFISH_DATA structure

6.12.2.4 double default_monk_adsorption (int i, int l, const void * $user_data$)

Default adsorption strength function.

This function will either use the default equilibrium function or the DOGFISH simulation result to produce the approximate adsorption strength using perturbation theory.

Parameters

i	index for the ith adsorbing species
1	index for the lth node in the domain
user_data	pointer to the MONKFISH_DATA structure

6.12.2.5 double default_monk_equilibrium (int i, int l, const void * user_data)

Default equilibirium adsorption function in mg/g.

This function uses the exterior species' concentration (mol/L), the species' molecular weight (g/mol), and the bulk fiber density (g/L) to calculate the adsorption equilibrium in mg/g. It assumes that the exterior concentration represents the moles of species per liter of solution that is being sorbed.

Parameters

i	index for the ith adsorbing species
1	index for the lth node in the domain
user_data	pointer to the MONKFISH_DATA structure

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6.12.2.6 double default_monkfish_retardation (int i, int l, const void * user_data)

Default retardation coefficient function.

This function calls the porosity, density, and adsorption functions to evaluate the retardation coefficient of the diffusing material.

Parameters

i	index for the ith adsorbing species
1	index for the lth node in the domain
user_data	pointer to the MONKFISH_DATA structure

6.12.2.7 double default_exterior_concentration (int i, const void * user_data)

Default exterior concentratio function.

This function assumes that the exterior concentration for sorption is just equal to the value of exterior_concentration given in MONKFISH_PARAM.

Parameters

i	index for the ith adsorbing species
user_data	pointer to the MONKFISH_DATA structure

6.12.2.8 double default_film_transfer (int i, const void * user_data)

Default film mass transfer function.

This function assumes that the film mass transfer coefficient is just equal to the value of the film_transfer_coeff in MONKFISH PARAM.

Parameters

i	index for the ith adsorbing species
user_data	pointer to the MONKFISH_DATA structure

6.12.2.9 int setup_MONKFISH_DATA (FILE * file, double(*)(int i, int I, const void *user_data) eval_porosity, double(*)(int i, int I, const void *user_data) eval_ext_diff, double(*)(int i, int I, const void *user_data) eval_ext_diff, double(*)(int i, int I, const void *user_data) eval_ext_double(*)(int i, const void *user_data) eval_ext_conc, double(*)(int i, const void *user_data) eval_ext_film, double(*)(int i, int I, const void *user_data) dog_diffusion, double(*)(int i, const void *user_data) dog_ext_film, double(*)(int i, const void *user_data) dog_surf_conc, const void *user_data, MONKFISH_DATA * monk_dat)

Setup function to allocate memory and setup function pointers for the MONKFISH simulation.

This function will allocate memory and setup the MONKFISH problem. To specify use of the default functions in MONKFISH, pass NULL args for all function pointers and the user_data data structure. Otherwise, pass in your own custom arguments. The MONKFISH DATA pointer must always be passed to this function.

Parameters

file	pointer to the output file to print out results
eval_porosity	function pointer for the bulk domain porosity function
eval_density	function pointer for the bulk domain density function
eval_ext_diff	function pointer for the interparticle diffusion function
eval_adsorb	function pointer for the adsorption strength function
eval_retard	function pointer for the retardation coefficient function
eval_ext_conc	function pointer for the external concentration function
eval_ext_film	function pointer for the external film mass transfer function
dog_diffusion	function pointer for the DOGFISH diffusion function (see dogfish.h)
dog_ext_film	function pointer for the DOGFISH film mass transfer (see dogfish.h)
dog_surf_conc	function pointer for the DOGFISH surface concentration (see dogfish.h)
user_data	pointer for the user's own data structure (only if using custom functions)
monk_dat	pointer for the MONKFISH_DATA structure

6.12.2.10 int MONKFISH_TESTS ()

Function to run tests on the MONKFISH algorithms.

This function currently does nothing and is not callable from the UI.

6.13 sandbox.h File Reference

Coding Test Area.

```
#include "flock.h"
#include "school.h"
```

Functions

• int RUN_SANDBOX ()

Function to run the methods implemented in the Sandbox.

6.13.1 Detailed Description

Coding Test Area.

sandbox.cpp

This file contains a series of simple tests for routines used in other files and algorithms. Before any code or methods are used, they are tested here to make sure that they are useful. The tests in the sandbox are callable from the UI to make it easier to alter existing sandbox code and run tests on new proposed methods or algorithms.

Warning

Functions and methods in this file are not meant to be used anywhere else.

Author

Austin Ladshaw

Date

04/11/2015

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6.13.2 Function Documentation

```
6.13.2.1 int RUN_SANDBOX ( )
```

Function to run the methods implemented in the Sandbox.

This function is callable from the UI and is used to observe results from the tests of newly developed algorithms. Edit header and source files here to test out your own routines or functions. Then you can run those functions by rebuilding the Ecosystem executable and running the sandbox tests.

6.14 school.h File Reference

Seawater Codes from a Highly Object-Oriented Library.

```
#include "eel.h"
#include "mola.h"
#include "shark.h"
#include "dogfish.h"
#include "monkfish.h"
#include "yaml_wrapper.h"
```

6.14.1 Detailed Description

Seawater Codes from a Highly Object-Oriented Library.

This file contains include statements for all files used in the aqueous adsorption problems, primarily targeted at Seawater simulations. Include this file into any other project or source code that needs the methods below.

Files Included in SCHOOL

eel.h mola.h shark.h dogfish.h monkfish.h yaml_wrapper.h

Note

- (1) shark.h also includes methods from macaw.h and lark.h
- (2) dogfish.h also includes methods from finch.h

Author

Austin Ladshaw

Date

02/23/2015

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6.15 scopsowl.h File Reference

Simultaneously Coupled Objects for Pore and Surface diffusion Operations With Linear systems.

```
#include "egret.h"
#include "skua.h"
```

Classes

struct SCOPSOWL PARAM DATA

Data structure for the species' parameters in SCOPSOWL.

struct SCOPSOWL_DATA

Primary data structure for SCOPSOWL simulations.

Macros

- #define SCOPSOWL_HPP_
- #define Dp(Dm, ep) (ep*ep*Dm)

Estimate of Pore Diffusivity (cm²/s)

#define Dk(rp, T, MW) (9700.0*rp*pow((T/MW),0.5))

Estimate of Knudsen Diffusivity (cm²/s)

#define avgDp(Dp, Dk) (pow(((1/Dp)+(1/Dk)),-1.0))

Estimate of Average Pore Diffusion (cm[^]2/s)

Functions

void print2file_species_header (FILE *Output, SCOPSOWL_DATA *owl_dat, int i)

Function to print out the main header for the output file.

void print2file_SCOPSOWL_time_header (FILE *Output, SCOPSOWL_DATA *owl_dat, int i)

Function to print out the time and space header for the output file.

void print2file_SCOPSOWL_header (SCOPSOWL_DATA *owl_dat)

Function to call the species and time header functions.

void print2file_SCOPSOWL_result_old (SCOPSOWL_DATA *owl_dat)

Function to print out the old time results to the output file.

• void print2file SCOPSOWL result new (SCOPSOWL DATA *owl dat)

Function to print out the new time results to the output file.

double default adsorption (int i, int I, const void *user data)

Default function for evaluating adsorption and adsorption strength.

• double default retardation (int i, int I, const void *user data)

Default function for evaluating retardation coefficient.

double default_pore_diffusion (int i, int I, const void *user_data)

Default function for evaluating pore diffusivity.

• double default surf diffusion (int i, int I, const void *user data)

Default function for evaluating surface diffusion for HOMOGENEOUS pellets.

double zero_surf_diffusion (int i, int I, const void *user_data)

Zero function for evaluating no surface diffusion in HOMOGENEOUS pellets.

double default effective diffusion (int i, int I, const void *user data)

Default function for evaluating effective diffusivity for HOMOGENEOUS pellets.

double const_pore_diffusion (int i, int I, const void *user_data)

Constant pore diffusion function for homogeneous or heterogeneous pellets.

double default_filmMassTransfer (int i, const void *user_data)

Default function for evaluating the film mass transfer coefficient.

double const_filmMassTransfer (int i, const void *user_data)

Constant film mass transfer coefficient function.

• int setup_SCOPSOWL_DATA (FILE *file, double(*eval_sorption)(int i, int I, const void *user_data), double(*eval_retardation)(int i, int I, const void *user_data), double(*eval_pore_diff)(int i, int I, const void *user_data), double(*eval_surface_diff)(int i, int I, const void *user_data), double(*eval_surface_diff)(int i, int I, const void *user_data), const void *user_data, MIXED_GAS *gas_data, SCOPSOWL_DATA *owl_data)

Setup function to allocate memory and setup function pointers for the SCOPSOWL simulation.

int SCOPSOWL Executioner (SCOPSOWL DATA *owl dat)

SCOPSOWL executioner function to solve a time step.

int set_SCOPSOWL_ICs (SCOPSOWL_DATA *owl_dat)

Function to set the initial conditions for a SCOPSOWL simulation.

• int set_SCOPSOWL_timestep (SCOPSOWL_DATA *owl_dat)

Function to set the timestep of the SCOPSOWL simulation.

• int SCOPSOWL preprocesses (SCOPSOWL DATA *owl dat)

Function to perform all preprocess SCOPSOWL operations.

• int set SCOPSOWL params (const void *user data)

Function to set the values of all non-linear system parameters during simulation.

• int SCOPSOWL postprocesses (SCOPSOWL DATA *owl dat)

Function to perform all postprocess SCOPSOWL operations.

int SCOPSOWL_reset (SCOPSOWL_DATA *owl_dat)

Function to reset all stateful information to prepare for next simulation.

• int SCOPSOWL (SCOPSOWL DATA *owl dat)

Function to progress the SCOPSOWL simulation through time till complete.

int SCOPSOWL_SCENARIOS (const char *scene, const char *sorbent, const char *comp, const char *sorbate)

Function to perform a SCOPSOWL simulation based on a set of parameters given in input files.

• int SCOPSOWL_TESTS ()

Function to run a SCOPSOWL test simulation.

6.15.1 Detailed Description

Simultaneously Coupled Objects for Pore and Surface diffusion Operations With Linear systems.

scopsowl.cpp

This file contains structures and functions associated with modeling adsorption in commercial, bi-porous adsorbents such as zeolites and mordenites. The pore diffusion and mass transfer equations are coupled with adsorption and surface diffusion through smaller crystals embedded in a binder matrix. However, you can also direct this simulation to treat the adsorbent as homogeneous (instead of heterogeneous) in order to model an even greater variety of gaseous adsorption kinetic problems. This object is coupled with either MAGPIE, SKUA, or BOTH depending on the type of simulation requested.

Author

Austin Ladshaw

Date

01/29/2015

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```
6.15.2 Macro Definition Documentation
```

6.15.2.1 #define SCOPSOWL_HPP_

6.15.2.2 #define Dp(Dm, ep) (ep*ep*Dm)

Estimate of Pore Diffusivity (cm²/s)

6.15.2.3 #define Dk(rp, T, MW) (9700.0*rp*pow((T/MW),0.5))

Estimate of Knudsen Diffusivity (cm²/s)

6.15.2.4 #define avgDp(Dp, Dk) (pow(((1/Dp)+(1/Dk)),-1.0))

Estimate of Average Pore Diffusion (cm²/s)

6.15.3 Function Documentation

```
6.15.3.1 void print2file_species_header ( FILE * Output, SCOPSOWL DATA * owl_dat, int i )
```

Function to print out the main header for the output file.

```
6.15.3.2 void print2file_SCOPSOWL_time_header ( FILE * Output, SCOPSOWL_DATA * owl_dat, int i )
```

Function to print out the time and space header for the output file.

```
6.15.3.3 void print2file_SCOPSOWL_header ( SCOPSOWL_DATA * owl_dat )
```

Function to call the species and time header functions.

```
6.15.3.4 void print2file_SCOPSOWL_result_old ( SCOPSOWL_DATA * owl_dat )
```

Function to print out the old time results to the output file.

```
6.15.3.5 void print2file_SCOPSOWL_result_new ( SCOPSOWL_DATA * owl_dat )
```

Function to print out the new time results to the output file.

```
6.15.3.6 double default_adsorption ( int i, int I, const void * user_data )
```

Default function for evaluating adsorption and adsorption strength.

This function is called in the preprocesses and postprocesses to estimate the strength of adsorption in the macroscale problem from perturbations. It will use perturbations in either the MAGPIE simulation or SKUA simulation, depending on the type of problem the user is solving.

Parameters

i	index for the ith species in the system
1	index for the lth node in the macro-scale domain
user_data	pointer for the SCOSPOWL_DATA structure

6.15.3.7 double default_retardation (int i, int I, const void * user_data)

Default function for evaluating retardation coefficient.

This function is called in the preprocesses and postprocesses to estimate the retardation coefficient for the simulation. It is recalculated at every time step to keep track of all changing conditions in the simulation.

Parameters

i	index for the ith species in the system
1	index for the lth node in the macro-scale domain
user_data	pointer for the SCOSPOWL_DATA structure

6.15.3.8 double default_pore_diffusion (int i, int l, const void * user_data)

Default function for evaluating pore diffusivity.

This function is called during the evaluation of non-linear residuals to more accurately represent non-linearities in the pore diffusion behavior. The pore diffusion is calculated based on kinetic theory of gases (see egret.h) and is adjusted according to the Knudsen Diffusion model and the porosity of the binder material.

Parameters

i	index for the ith species in the system
1	index for the lth node in the macro-scale domain
user_data	pointer for the SCOSPOWL_DATA structure

6.15.3.9 double default_surf_diffusion (int i, int l, const void * user_data)

Default function for evaluating surface diffusion for HOMOGENEOUS pellets.

This function is ONLY used if the pellet is determined to be homogeneous. Otherwise, this is replaced by the surface diffusion function for the SKUA simulation. The diffusivity is calculated based on the Arrhenius rate expression and then adjusted by the outside partial pressure of the adsorbing species.

Parameters

i index for the ith species in the systemI index for the lth node in the macro-scale domain	

6.15.3.10 double zero_surf_diffusion (int i, int l, const void * user_data)

Zero function for evaluating no surface diffusion in HOMOGENEOUS pellets.

This function is ONLY used if the pellet is determined to be homogeneous and we want to specify that there is no surface diffusion.

Parameters

i	index for the ith species in the system
1	index for the lth node in the macro-scale domain
user_data	pointer for the SCOSPOWL_DATA structure

6.15.3.11 double default_effective_diffusion (int i, int l, const void * $user_data$)

Default function for evaluating effective diffusivity for HOMOGENEOUS pellets.

This function is ONLY used if the pellet is determined to be homogeneous. Otherwise, this is replaced by the pore diffusion function. The effective diffusivity is determined by the combination of pore diffusivity and surface diffusivity with adsorption strength in an homogeneous pellet.

Parameters

i	index for the ith species in the system index for the lth node in the macro-scale domain	
1		
user_data	pointer for the SCOSPOWL_DATA structure	

6.15.3.12 double const_pore_diffusion (int i, int I, const void * user_data)

Constant pore diffusion function for homogeneous or heterogeneous pellets.

This function should be used if the user wants to specify a constant pore diffusivity. The value of pore diffusion is then set equal to the value of pore_diffusion in the SCOPSOWL_PARAM_DATA structure.

Parameters

i	index for the ith species in the system index for the lth node in the macro-scale domain	
1		
user_data	pointer for the SCOSPOWL_DATA structure	

6.15.3.13 double default_filmMassTransfer (int i, const void * user_data)

Default function for evaluating the film mass transfer coefficient.

This function is called during the setup of the boundary conditions and is used to estimate the film mass transfer coefficient for the macro-scale problem. The coefficient is calculated according to the kinetic theory of gases (see egret.h).

Parameters

i	index for the ith species in the system	
user_data	pointer for the SCOSPOWL_DATA structure	

6.15.3.14 double const_filmMassTransfer (int i, const void * user_data)

Constant film mass transfer coefficient function.

This function is used when the user wants to specify a constant value for film mass transfer. The value of that coefficient is then set equal to the value of film_transfer in the SCOPSOWL_PARAM_DATA structure.

Parameters

i	index for the ith species in the system	
user_data	pointer for the SCOSPOWL_DATA structure	

6.15.3.15 int setup_SCOPSOWL_DATA (FILE * file, double(*)(int i, int I, const void *user_data) eval_sorption, double(*)(int i, int I, const void *user_data) eval_pore_diff, double(*)(int i, const void *user_data) eval_surface_diff, const void * user_data, MIXED_GAS * gas_data, SCOPSOWL_DATA * owl_data)

Setup function to allocate memory and setup function pointers for the SCOPSOWL simulation.

This function sets up the memory and function pointers used in SCOPSOWL simulations. User can provide NULL in place of functions for the function pointers and the setup will automatically use just the default settings. However, the user is required to pass the necessary data structure pointers for MIXED_GAS and SCOPSOWL_DATA.

Parameters

file	pointer to the output file to print out results	
eval_sorption	pointer to the adsorption evaluation function	
eval_retardation	pointer to the retardation evaluation function	
eval_pore_diff	pointer to the pore diffusion function	
eval_filmMT	pointer to the film mass transfer function	
eval_surface_diff	pointer to the surface diffusion function (required)	
user_data	pointer to the user's data structure used for the parameter functions	
gas_data	pointer to the MIXED_GAS structure used to evaluate kinetic gas theory	
owl_data	pointer to the SCOPSOWL_DATA structure	

6.15.3.16 int SCOPSOWL_Executioner (SCOPSOWL_DATA * owl_dat)

SCOPSOWL executioner function to solve a time step.

This function will call the preprocess, solver, and postprocess functions to evaluate a single time step in a simulation. All simulation conditions must be set prior to calling this function. This function will typically be the one called from other simulations that will involve a SCOPSOWL evaluation to resolve kinetic coupling.

Parameters

owl_	_dat	pointer to the SCOPSOWL_	DATA structure (must be initialized)
------	------	--------------------------	--------------------------------------

6.15.3.17 int set_SCOPSOWL_ICs (SCOPSOWL_DATA * owl_dat)

Function to set the initial conditions for a SCOPSOWL simulation.

This function will setup the initial conditions of the simulation based on the initial temperature, pressure, and adsorbed molefractions. It assumes that the initial conditions are constant throughout the domain of the problem. This function should only be called once during a simulation.

Parameters

 owl_dat
 pointer to the SCOPSOWL_DATA structure (must be initialized)

6.15.3.18 int set_SCOPSOWL_timestep (SCOPSOWL_DATA * owl_dat)

Function to set the timestep of the SCOPSOWL simulation.

This function is used to set the next time step to be used in the SCOPSOWL simulation. A constant time step based on the size of the pellet discretization will be used. Users may want to use a custom time step to ensure that coupled-multi-scale systems are all in sync.

Parameters

|--|

6.15.3.19 int SCOPSOWL_preprocesses (SCOPSOWL_DATA * owl_dat)

Function to perform all preprocess SCOPSOWL operations.

This function will update the boundary conditions and simulation conditions based on the current temperature, pressure, and gas phase composition, which may all vary in time. Since this function is called by the SCOPSOW← L_Executioner, it does not need to be called explicitly by the user.

Parameters

owl_dat pointer to the SCOPS	OWL_DATA structure (must be initialized)
--------------------------------	--

6.15.3.20 int set_SCOPSOWL_params (const void * user_data)

Function to set the values of all non-linear system parameters during simulation.

This is the function override for the FINCH setparams function (see finch.h). It will update the values of non-linear parameters in the residuals so that all variables in a species' system are fully coupled.

Parameters

user_data	pointer to the SCOPSOWL_DATA structure (must be initialized)

6.15.3.21 int SCOPSOWL_postprocesses (SCOPSOWL_DATA * owl_dat)

Function to perform all postprocess SCOPSOWL operations.

This function will update the retardation coefficients based on newly obtained simulation results for the current time step and calculate the average and total amount of adsorption of each species in the domain. Additionally, this function will call the print functions to store simulation results in the output file.

Parameters

owl_dat	pointer to the SCOPSOWL_DATA structure (must be initialized)
---------	--

6.15.3.22 int SCOPSOWL_reset (SCOPSOWL_DATA * owl_dat)

Function to reset all stateful information to prepare for next simulation.

This function will update the stateful information used in SCOPSOWL to prepare the system for the next time step in the simulation. However, because updating the states erases the old state, the user must be absolutely sure that

the simulation is ready to be updated. other simulations it is very important.	For just running standard simulations,	, this is not an issue, but in coupling w	ith

Parameters

owl_dat pointer to the SCOPSOWL_DATA structure (must be initial

6.15.3.23 int SCOPSOWL (SCOPSOWL_DATA * owl_dat)

Function to progress the SCOPSOWL simulation through time till complete.

This function will call the initial conditions, then progressively call the executioner, time step, and reset functions to propagate the simulation in time. As such, this function is primarily used when running a SCOPSOWL simulation by itself and not when coupling it to an other problem.

Parameters

owl_dat	pointer to the SCOPSOWL_DATA structure (must be initialized)
---------	--

6.15.3.24 int SCOPSOWL_SCENARIOS (const char * scene, const char * sorbent, const char * comp, const char * sorbate)

Function to perform a SCOPSOWL simulation based on a set of parameters given in input files.

This is the primary function to be called when running a stand-alone SCOPSOWL simulation. Parameters and system information for the simulation are given in a series of input files that come in as character arrays. These inputs are all required to call this function.

Parameters

scene	Sceneario Input File
sorbent	Adsorbent Input File
comp	Component Input File
sorbate	Adsorbate Input File

Note

Each input file has a particular format that must be strictly adhered to in order for the simulation to be carried out correctly. The format for each input file, and an example, is provided below...

Scenario Input Format

System Temperature (K) [tab] Total Pressure (kPa) [tab] Gas Velocity (cm/s)

Simulation Time (hrs) [tab] Print Out Time (hrs)

BC Type (0 = Neumann, 1 = Dirichlet)

Number of Gas Species

Initial Total Adsorption (mol/kg)

Name of ith Species [tab] Adsorbable? (0 = false, 1 = true) [tab] Gas Phase Molefraction [tab] Initial Sorbed Molefraction

(repeat above for all species)

Example Scenario Input

353.15 101.35 0.36 4.0 0.05 0 5 0.0 N2 0 0.7634 0.0 O2 0 0.2081 0.0 Ar 0 0.009 0.0 CO2 0 0.0004 0.0 H2O 1 0.0191 0.0

Above example is for a 5-component mixture of N2, O2, Ar, CO2, and H2O, but we are only considering the H2O as adsorbable.

Adsorbent Input File

```
Heterogeneous Pellet? (0 = false, 1 = true) [tab] Surface Diffusion Included? (0 = false, 1 = true) Macro Coord. (2 = spherical, 1 = cylindrical) { [tab] Char. Length (cm) (i.e., cylinder length) } (NOTE: Char. Length is only needed if problem is not spherical) Pellet Radius (cm) [tab] Pellet Density (kg/L) [tab] Porosity (vol. void / vol. binder) [tab] Pore Radius (cm) (Below is only needed if pellet is Heterogeneous) Micro Coord. (2 = spherical, 1 = cylindrical) { [tab] Char. Length (cm) (i.e., cylinder length) } Crystal Radius (um) [tab] Binder Fraction (vol. binder / vol. pellet)
```

Example Adsorbent Input

1 1 2 0.118 1.69 0.272 3.5E-6 2 2.0 0.175

Above example is for a heterogeneous adsorbent with surface diffusion. The pellet and crystals are both considered spherical. Pellet radius is 0.118 cm, density is 1.69 kg/L, porosity is 0.272, and pore size is 3.5e-6 cm. The pellet is made up of 17.5 % binder material and contains crystals roughly 2.0 um in radius.

Component Input File

Molar Weight of ith species (g/mol) [tab] Specific Heat of ith species (J/g/K) Sutherland Viscosity (g/cm/s) [tab] Sutherland Temperature (K) [tab] Sutherland Constant (K) of ith species (repeat above for all species in same order they appeared in the Scenario Input File)

Example Component Input

28.016 1.04 0.0001781 300.55 111.0 32.0 0.919 0.0002018 292.25 127.0 39.948 0.522 0.0002125 273.11 144.4 44.009 0.846

```
0.000148 293.15 240.0
18.0 1.97
0.0001043 298.16 784.72
```

Above example is a continuation of the Scenario Input example wherein each grouping represents parameters that are associated with N2, O2, Ar, CO2, and H2O, respectively. The order is VERY important!

Adsorbate Input File

```
{ Type of Surface Diffusion Function (0 = constant, 1 = simple Darken, 2 = theoretical Darken) } (NOTE: The above option is only given IF the pellet was specified as Heterogeneous!) Reference Diffusivity (um^2/hr) [tab] Activation Energy (J/mol) of ith adsorbable species Reference Temperature (K) [tab] Affinity Constant (-) of ith adsorbable species van der Waals Volume (cm^3/mol) of ith species GSTA adsorption capacity (mol/kg) of ith species Number of GSTA parameters of ith species Enthalpy (J/mol) of nth site [tab] Entropy of nth site (J/K/mol) of ith species (repeat enthalpy and entropy for all n sites in species i) (repeat above for all species i)
```

Example Adsorbate Input

```
0
0.8814 0.0
267.999 0.0
13.91
11.67
-46597.5 -53.6994
-125024 -221.073
-193619 -356.728
-272228 -567.459
1.28 540.1
374.99 0.01
3.01
1.27
2
-46597.5 -53.6994
-125024 -221.073
```

Above example would be for a simulation involving two adsorbable species using a constant surface diffusion function. Each adsorbable species has it's own set of kinetic and equilibrium parameters that must be given in the same order as the species appeared in the Scenario Input. Note: we do not need to supply this information for non-adsorbable species.

```
6.15.3.25 int SCOPSOWL_TESTS ( )
```

Function to run a SCOPSOWL test simulation.

This function runs a test of the SCOPSOWL physics and prints out results to a text file. It is callable from the UI.

6.16 scopsowl_opt.h File Reference

Optimization Routine for Surface Diffusivities in SCOPSOWL.

```
#include "scopsowl.h"
```

Classes

struct SCOPSOWL OPT DATA

Data structure for the SCOPSOWL optmization routine.

Functions

int SCOPSOWL_OPT_set_y (SCOPSOWL_OPT_DATA *owl_opt)

Function to set the rest of the gas phase mole fractions based on current mole fraction of adsorbing gas.

int initial_guess_SCOPSOWL (SCOPSOWL_OPT_DATA *owl_opt)

Function to set up an initial guess for the surface diffusivity parameter in SCOPSOWL.

- void eval_SCOPSOWL_Uptake (const double *par, int m_dat, const void *data, double *fvec, int *info)
 - Function that works in conjunction with the Imfit routine to minimize the euclidean norm between function and data.
- int SCOPSOWL_OPTIMIZE (const char *scene, const char *sorbent, const char *comp, const char *sorbate, const char *data)

Function called to perform the optimization routine given a specific set of information and data.

6.16.1 Detailed Description

Optimization Routine for Surface Diffusivities in SCOPSOWL.

scopsowl_opt.cpp

This file contains structures and functions associated with performing non-linear least-squares optimization of the SCOPSOWL simulation results against actual kinetic adsorption data. The optimization routine here allows you to run data comparisons and optimizations in three forms: (i) Rough optimizations - cheaper operations, but less accurate, (ii) Exact optmizations - much more expensive, but greater accuracy, and (iii) data/model comparisons - no optimization, just using system parameters to compare simulation results agains a set of data.

Depending on the level of optimization desired, this routine could take several minutes or several hours. The optimization/comparisons are printed out in two files: (i) a parameter file, which contains the simulation partial pressures and temperatures and the optimized diffusivities with the euclidean norm of the fitting and (ii) a comparison file that shows the model value and data value at each time step for each kinetic curve.

The optimized diffusion parameters are given for each individual kinetic data curve. Each data curve will have a different pairing of partial pressure and temperature. Because of this, you will get a list of different diffusivities for each data curve. To get the optimum kinetic parameters from this list of diffusivities, you must fit the diffusion parameter values to the following diffusion function model...

$$D_opt = D_ref * exp(-E / (R*T)) * pow(p, (T_ref/T) - B)$$

where D_ref is the Reference Diffusivity (um^2/hr), E is the activation energy for adsorption (J/mol), R is the gas law constant (J/K/mol), T is the system temperature (K), p is the partial pressure of the adsorbing species (kPa), T_ref is the Reference Temperature (K), and B is the Affinity constant. This algorithm does not automatically produce these parameters for you, but gives you everything you need to produce them yourself.

This routine allows you to optimize multiple kinetic curves at one time. However, all data must be for the same adsorbent-adsorbate system. In other words, the adsorbent and adsorbate pair must be the same for each kinetic curve analyzed. Also, each experiment must have been done in a thin bed or continuous flow system where the adsorbents were exposed to a nearly constant outside partial pressure for all time steps and the gas velocity of that system is assumed constant for all experiments. This experimental setup is very typical for studying adsorption kinetics for gas-solid systems.

Author

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Date

05/14/2015

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6.16.2 Function Documentation

```
6.16.2.1 int SCOPSOWL_OPT_set_y ( SCOPSOWL_OPT_DATA * owl_opt )
```

Function to set the rest of the gas phase mole fractions based on current mole fraction of adsorbing gas.

This function takes the current mole fraction of the adsorbing gas and calculates the gas mole fractions of the other gases in the sytem based on the standard inlet gas composition given in the scenario file.

```
6.16.2.2 int initial_guess_SCOPSOWL ( SCOPSOWL OPT DATA * owl_opt )
```

Function to set up an initial guess for the surface diffusivity parameter in SCOPSOWL.

This function performs the Rough optimization on the surface diffusivity based on the idea of reducing or eliminating function bias between data and simulation. A positive function bias means that the simulation curve is "higher" than the data curve and a negative function bias means that the simulation curve is "lower" than the data curve. We use this information to incrementally adjust the rate of surface diffusion until this bias is near zero. When bias is near zero, the simulation is nearly optimized, but further refinement may be necessary to find the true minimum solution.

```
6.16.2.3 void eval_SCOPSOWL_Uptake ( const double * par, int m_dat, const void * data, double * fvec, int * info )
```

Function that works in conjunction with the Imfit routine to minimize the euclidean norm between function and data.

This function will run the SCOPSOWL simulation at a given value of surface diffusivity and produce residuals that feed into the Levenberg-Marquardt's algorithm for non-linear least-squares regression. The form of this function is specific to the format required by the Imfit routine.

Parameters

par	array of parameters that are to be optimized
m_dat	number of data points or functions to evaluate
data	user supplied data structure holding information necessary to form the residuals
fvec	array of residuals computed at the current parameter values
info	integer pointer denoting whether or not the user requests to end a particular simulation

6.16.2.4 int SCOPSOWL_OPTIMIZE (const char * scene, const char * sorbent, const char * comp, const char * sorbate, const char * data)

Function called to perform the optimization routine given a specific set of information and data.

This is the function that is callable by the UI. The user must provide 5 input files to the routine in order to establish simulation conditions, adsborbent properties, component properties, adsorbate equilibrium parameters, and the set of data that we are comparing the simulations to. Each input file has a very specific structure and order to the information that it contains. The structure here is DIFFERENT than the structure for just running standard SCOP ← SOWL simulations (see scopsowl.h).

Parameters

scene	Sceneario Input File
sorbent	Adsorbent Input File
comp	Component Input File
sorbate	Adsorbate Input File
data	Kinetic Adsorption Data File
uaia	Minetic Ausorption Data File

Note

Much of the structure of these input files are "similar" to that of the input files used in SCOPSOWL_SCENA ← RIOS (see scopsowl.h), but with some notable differences. Below gives the format for each input file with an example. Make sure your input files follow this format before calling this routine from the UI.

Scenario Input File

```
Optimization? (0 = false, 1 = true) [tab] Rough Optimization? (0 = false, 1 = true)

Surf. Diff. (0 = constant, 1 = simple Darken, 2 = theoretical Darken) [tab] BC Type (0 = Neumann, 1 = Dirichlet)

Total Pressure (kPa) [tab] Gas Velocity (cm/s)

Number of Gaseous Species

Initial Adsorption Total (mol/kg)
```

Name [tab] Adsorbable? (0 = false, 1 = true) [tab] Inlet Gas Mole Fraction [tab] Initial Adsorbed Mole Fraction (NOTE: The above line is repeated for all species in gas phase. Also, this algorithm only allows you to consider one adsorbable gas component. Inlet gas mole fractions must be non-zero for all non-adsorbing gases and must sum to 1.)

Example Scenario Input

```
1 0
0 0
101.35 0.36
5
0.0
N2 0 0.7825 0.0
O2 0 0.2081 0.0
Ar 0 0.009 0.0
CO2 0 0.0004 0.0
H2O 1 0.0 0.0
```

Above example is for running optimizations on data collected with a gas stream at 0.36 cm/s with 5 gas species in the mixture, only H2O of which is adsorbing. The "base line" or "inlet gas" without H2O has a composition of N2 at 0.7825, O2 at 0.2081, Ar at 0.009, and CO2 at 0.0004.

Adsorbent Input File

```
Heterogeneous Pellet? (0 = false, 1 = true)

Macro Coord. (2 = spherical, 1 = cylindrical) { [tab] Char. Length (cm) (i.e., cylinder length) }

(NOTE: Char. Length is only needed if problem is not spherical)

Pellet Radius (cm) [tab] Pellet Density (kg/L) [tab] Porosity (vol. void / vol. binder) [tab] Pore Radius (cm)

(Below is only needed if pellet is Heterogeneous)

Micro Coord. (2 = spherical, 1 = cylindrical) { [tab] Char. Length (cm) (i.e., cylinder length) }

Crystal Radius (um) [tab] Binder Fraction (vol. binder / vol. pellet)
```

Example Adsorbent Input

```
1
2
0.118 1.69 0.272 3.5E-6
2
2.0 0.175
```

Above example is nearly identical to the file given in the SCOPSOWL_SCENARIO example (see scopsowl.h). However, here we do not give an integer flag denoting whether or not we are considering surface diffusion as a mechanism. This is because we automatically assume that surface diffusion is a mechanism in the system, since that is the unknown parameter that we are performing the optimizations for.

Component Input File

Molar Weight of ith species (g/mol) [tab] Specific Heat of ith species (J/g/K) Sutherland Viscosity (g/cm/s) [tab] Sutherland Temperature (K) [tab] Sutherland Constant (K) of ith species (repeat above for all species in same order they appeared in the Scenario Input File)

Example Component Input

28.016 1.04 0.0001781 300.55 111.0 32.0 0.919 0.0002018 292.25 127.0 39.948 0.522 0.0002125 273.11 144.4 44.009 0.846 0.000148 293.15 240.0 18.0 1.97 0.0001043 298.16 784.72

Above example is exactly the same as in the SCOPSOWL_SCENARIO example (see scopsowl.h). There is no difference in the input file formats for this input. Keep in mind that the order is VERY important! All species information must be in the same order that the species appeared in the Scenario input file.

Adsorbate Input File

Reference Diffusivity (um^2/hr) [tab] Activation Energy (J/mol) of ith adsorbable species Reference Temperature (K) [tab] Affinity Constant (-) of ith adsorbable species van der Waals Volume (cm^3/mol) of ith species GSTA adsorption capacity (mol/kg) of ith species Number of GSTA parameters of ith species Enthalpy (J/mol) of nth site [tab] Entropy of nth site (J/K/mol) of ith species (repeat enthalpy and entropy for all n sites in species i) (repeat above for all species i)

Example Adsorbate Input

11.67

11

-46597.5 -53.6994

-125024 -221.073

-193619 -356.728

-272228 -567.459

Above example gives the equilibrium parameters associated with the H2O-MS3A single component adsorption system. Note that the kinetic parameters (Ref. Diff., Act. Energy, Ref. Temp., and Affinity) were all given a value of zero. These values are irrelavent if we are running an optimization because they will be replaced with a single estimate for the diffusivity that is being optimization for. However, if we wanted to run this routine with comparisons and not do any optimization, then you would need to provide non-zero values for these parameters (at least for Ref. Diff.).

Data Input File

```
Number of Kinetic Data Curves
Number of data points in the ith curve
Temperature (K) [tab] Partial Pressure (kPa) [tab] Equilibrium Adsorption (mol/kg) all of ith curve
Time point 1 (hrs) [tab] Adsorption 1 (mol/kg) of ith curve
Time point 1 (hrs) [tab] Adsorption 2 (mol/kg) of ith curve
... (Repeat for all time-adsorption data points)
(Repeat above for all curves i)
```

Example Data Input

```
40
2990
298.15 0.000310922 2.9
0 0
0.1666666667 0.001834419
0.333611111 0.004880247
0.5 0.008306803
...
2789
298.15 0.00055189 5
0 0
0.166944444 0.003350185
0.333611111 0.007418267
0.5 0.009930906
0.6666666667 0.014597236
0.833611111 0.021377373
```

Above is a partial example for a data set of 40 kinetic curves. The first curve contains 2990 data points and has temperature of 298.15 K, partial pressure of 0.000310922 kPa, and an equilibrium adsorption of 2.9. Each first time point should start from 0 hours and each initial adsorption should correspond to the value of initial adsorption indicated in the Scenario input file. Then, this structure is repeated for all adsorptio curves.

6.17 shark.h File Reference

Speciation-object Hierarchy for Adsorption Reactions and Kinetics.

```
#include "mola.h"
#include "macaw.h"
#include "lark.h"
#include "yaml_wrapper.h"
#include "dogfish.h"
```

Classes

· class MasterSpeciesList

Master Species List Object.

class Reaction

Reaction Object.

· class MassBalance

Mass Balance Object.

class UnsteadyReaction

Unsteady Reaction Object (inherits from Reaction)

· class AdsorptionReaction

Adsorption Reaction Object.

· class UnsteadyAdsorption

Unsteady Adsorption Reaction Object.

class MultiligandAdsorption

Multi-ligand Adsorption Reaction Object.

· class ChemisorptionReaction

Chemisorption Reaction Object.

· class MultiligandChemisorption

Multi-ligand Chemisorption Reaction Object.

struct SHARK_DATA

Data structure for SHARK simulations.

Macros

• #define Rstd 8.3144621

Gas Law Constant in J/K/mol (or) L*kPa/K/mol (Standard Units)

• #define Na 6.0221413E+23

Avagadro's Number - Units: molecules/mol.

#define kB 1.3806488E-23

Boltzmann's Constant - Units: J/K or C*V/K.

• #define e 1.6021766208E-19

Elementary Electric Charge - Units: C.

• #define Faraday 96485.33289

Faraday's Constant - C/mol.

#define VolumeSTD 15.17

Standard Segment Volume - cm[^] 3/mol.

• #define AreaSTD 2.5E5

Standard Segment Area - m²/mol.

• #define CoordSTD 10

Standard Coordination Number.

#define LengthFactor(z, r, s) (((z/2.0)*(r-s)) - (r-1.0))

Calculation of the Length Factor Parameter in UNIQUAC.

• #define VacuumPermittivity 8.8541878176E-12

Vacuum Permittivity Constant - F/m or C/V/m.

• #define WaterRelPerm 80.1

Approximate Relative Permittivity for water - Unitless.

#define AbsPerm(Rel) (Rel*VacuumPermittivity)

Calculation of Absolute Permittivity of a medium - F/m or C/V/m.

Typedefs

typedef struct SHARK_DATA SHARK_DATA

Data structure for SHARK simulations.

Enumerations

enum valid_mb { BATCH, CSTR, PFR }

Enumeration for the list of valid activity models for non-ideal solutions.

enum valid_act { IDEAL, DAVIES, DEBYE_HUCKEL, SIT, PITZER }

Enumeration for the list of valid activity models for non-ideal solutions.

enum valid_surf_act { IDEAL_ADS, FLORY_HUGGINS, UNIQUAC_ACT }

Enumeration for the list of valid surface activity models for non-ideal adsorption.

Functions

void print2file_shark_info (SHARK_DATA *shark_dat)

Function to print out simulation conditions and options to the output file.

void print2file_shark_header (SHARK_DATA *shark_dat)

Function to print out the head of species and time stamps to the output file.

void print2file_shark_results_new (SHARK_DATA *shark_dat)

Function to print out the simulation results for the current time step.

void print2file_shark_results_old (SHARK_DATA *shark_dat)

Function to print out the simulation results for the previous time step.

double calculate_ionic_strength (const Matrix< double > &x, MasterSpeciesList &MasterList)
 Function to calculate the ionic strength of the solution.

• int FloryHuggins (const Matrix< double > &x, Matrix< double > &F, const void *data)

Surface Activity function for simple non-ideal adsorption (for adsorption reaction object)

• int FloryHuggins_unsteady (const Matrix< double > &x, Matrix< double > &F, const void *data)

Surface Activity function for simple non-ideal adsorption (for unsteady adsorption object)

 $\bullet \ \, \text{int FloryHuggins_multiligand (const Matrix} < \ \, \text{double} > \&x, \ \, \text{Matrix} < \ \, \text{double} > \&F, \ \, \text{const void } * \ \, \text{data}) \\$

Surface Activity function for simple non-ideal adsorption (for multiligand adsorption object)

• int FloryHuggins_chemi (const Matrix< double > &x, Matrix< double > &F, const void *data) Surface Activity function for simple non-ideal adsorption (for chemisorption reaction object)

• int FloryHuggins_multichemi (const Matrix< double > &x, Matrix< double > &F, const void *data)

Surface Activity function for simple non-ideal adsorption (for multiligand chemisorption object)

int UNIQUAC (const Matrix< double > &x, Matrix< double > &F, const void *data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for adsorption reaction object)

 $\bullet \ \, \text{int UNIQUAC_unsteady (const Matrix} < \ \, \text{double} > \&x, \ \, \text{Matrix} < \ \, \text{double} > \&F, \ \, \text{const void} * \ \, \text{data}) \\$

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for unsteady adsorption object)

• int UNIQUAC_multiligand (const Matrix< double > &x, Matrix< double > &F, const void *data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for multiligand adsorption object)

int UNIQUAC_chemi (const Matrix< double > &x, Matrix< double > &F, const void *data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for chemisorption reaction object)

int UNIQUAC_multichemi (const Matrix < double > &x, Matrix < double > &F, const void *data)
 Surface Activity function for the UNIQUAC model for non-ideal adsorption (for multiligand chemisorption object)

- int ideal_solution (const Matrix< double > &x, Matrix< double > &F, const void *data)

Activity function for Ideal Solution.

• int Davies_equation (const Matrix< double > &x, Matrix< double > &F, const void *data)

**Activity function for Davies Equation.

• int DebyeHuckel_equation (const Matrix< double > &x, Matrix< double > &F, const void *data)

**Activity function for Debye-Huckel Equation.

int surf_act_choice (const std::string &input)

First test of SIT Model.

int act_choice (const std::string &input)

Function takes a given string and returns a flag denoting which activity model was choosen.

int reactor_choice (const std::string &input)

Function takes a give string and returns a flag denoting which type of reactor was choosen for the system.

bool linesearch_choice (const std::string &input)

Function returns a bool to determine the form of line search requested.

int linearsolve_choice (const std::string &input)

Function returns the linear solver flag for the PJFNK method.

int Convert2LogConcentration (const Matrix< double > &x, Matrix< double > &logx)

Function to convert the given values of variables (x) to the log of those variables (logx)

int Convert2Concentration (const Matrix< double > &logx, Matrix< double > &x)

Function to convert the given log values of variables (logx) to the values of those variables (x)

int read_scenario (SHARK_DATA *shark_dat)

Function to go through the yaml object for the scenario document.

int read multiligand scenario (SHARK DATA *shark dat)

Function to go through the yaml object to setup memory space for multiligand objects.

• int read_multichemi_scenario (SHARK_DATA *shark_dat)

Function to go through the yaml object to setup memory space for multiligand chemisorption objects.

int read options (SHARK DATA *shark dat)

Function to go through the yaml object for the solver options document.

• int read species (SHARK DATA *shark dat)

Function to go through the yaml object for the master species document.

int read massbalance (SHARK DATA *shark dat)

Function to go through the yaml object for the mass balance document.

int read equilrxn (SHARK DATA *shark dat)

Function to go through the yaml object for the equilibrium reaction document.

int read unsteadyrxn (SHARK DATA *shark dat)

Function to go through the yaml object for the unsteady reaction document.

int read_adsorbobjects (SHARK_DATA *shark_dat)

Function to go through the yaml object for each Adsorption Object.

• int read unsteadyadsorbobjects (SHARK DATA *shark dat)

Function to go through the yaml object for each Unsteady Adsorption Object.

int read_multiligandobjects (SHARK_DATA *shark_dat)

Function to go through the yaml object for each MultiligandAdsorption Object.

int read_chemisorbobjects (SHARK_DATA *shark_dat)

Function to go through the yaml object for each Chemisorption Object.

• int read multichemiobjects (SHARK DATA *shark dat)

Function to go through the yaml object for each MultiligandChemisorption Object.

int setup_SHARK_DATA (FILE *file, int(*residual)(const Matrix< double > &x, Matrix< double > &res, const void *data), int(*activity)(const Matrix< double > &x, Matrix< double > &gama, const void *data), int(*precond)(const Matrix< double > &r, Matrix< double > &p, const void *data), SHARK_DATA *dat, const void *activity_data, const void *residual_data, const void *precon_data, const void *other_data)

Function to setup the memory and pointers for the SHARK_DATA structure for the current simulation.

• int shark_add_customResidual (int i, double(*other_res)(const Matrix< double > &x, SHARK_DATA *shark dat, const void *other data), SHARK_DATA *shark dat)

Function to add user defined custom residual functions to the OtherList vector object in SHARK_DATA.

int shark_parameter_check (SHARK_DATA *shark_dat)

Function to check the Reaction and UnsteadyReaction objects for missing info.

int shark_energy_calculations (SHARK_DATA *shark_dat)

Function to calculate all Reaction and UnsteadyReaction energies.

• int shark_temperature_calculations (SHARK_DATA *shark_dat)

Function to calculate all Reaction and UnsteadyReaction parameters as a function of temperature.

int shark_pH_finder (SHARK_DATA *shark_dat)

Function will search MasterSpeciesList for existance of H + (aq) and OH - (aq) molecules.

int shark guess (SHARK DATA *shark dat)

Function provides a rough initial guess for the values of all non-linear variables.

int shark_initial_conditions (SHARK_DATA *shark_dat)

Function to establish the initial conditions of the shark simulation.

int shark executioner (SHARK DATA *shark dat)

Function to execute a shark simulation at a single time step or pH value.

int shark_timestep_const (SHARK_DATA *shark_dat)

Function to set up all time steps in the simulation to a specified constant.

int shark timestep adapt (SHARK DATA *shark dat)

Function to set up all time steps in the simulation based on success or failure to converge.

int shark_preprocesses (SHARK_DATA *shark_dat)

Function to call other functions for calculation of parameters and setting of time steps.

int shark solver (SHARK DATA *shark dat)

Function to call the PJFNK solver routine given the current SHARK DATA information.

int shark_postprocesses (SHARK_DATA *shark_dat)

Function to convert PJFNK solutions to concentration values and print to the output file.

int shark_reset (SHARK_DATA *shark_dat)

Function to reset the values of all stateful information in SHARK_DATA.

int shark_residual (const Matrix < double > &x, Matrix < double > &F, const void *data)

Default residual function for shark evaluations.

int SHARK (SHARK_DATA *shark_dat)

Function to call all above functions to perform a shark simulation.

int SHARK_SCENARIO (const char *yaml_input)

Function to perform a shark simulation based on the conditions in a yaml formatted input file.

int SHARK_TESTS ()

Function to perform a series of shark calculation tests.

int SHARK_TESTS_OLD ()

Function to perform a series of shark calculation tests (older version)

6.17.1 Detailed Description

Speciation-object Hierarchy for Adsorption Reactions and Kinetics.

shark.cpp

This file contains structures and functions associated with solving speciation and kinetic problems in aqueous systems. The primary aim for the development of these algorithms was to solve speciation and adsorption problems for the recovery of uranium resources from seawater. Seawater is an extradorinarily complex medium in which to work, which is why these algorithms are being constructed in a piece-wise, object-oriented fashion. This allows us to displace much of the complexity of the problem by breaking it down into smaller, more managable pieces.

Each piece of SHARK contributes to a residual function when solving the overall speciation, reaction, kinetic chemical problem. These residuals are then fed into the PJFNK solver function in lark.h. The variables of the system are the log(C) concentration values of each species in the system. We solve for log(C) concentrations, rather than just C, because the PJFNK method is an unbounded solution algorithm. So to prevent the algorithm from producing negative values for concentration, we reformulate all residuals in terms of the log(C) values. In this way, regardless of the value found for log(C), the concentration C will always be greater than 0.

Currenty, SHARK supports standard aqueous speciation problems with simple kinetic models based on an unsteady form of the standard reaction stoichiometry. As more methods and algorithms are completed, the SHARK simulations will be capable of doing much, much more.

Warning

Much of this is still underconstruction and many methods or interfaces may change. Use with caution.

Author

Austin Ladshaw

Date

05/27/2015

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This software was designed and built at the Georgia Institute of Technology by Austin Ladshaw for PhD research in the area of adsorption and surface science. Copyright (c) 2015, all rights reserved.

6.17.2 Macro Definition Documentation

6.17.2.1 #define Rstd 8.3144621

Gas Law Constant in J/K/mol (or) L*kPa/K/mol (Standard Units)

6.17.2.2 #define Na 6.0221413E+23

Avagadro's Number - Units: molecules/mol.

6.17.2.3 #define kB 1.3806488E-23

Boltzmann's Constant - Units: J/K or C*V/K.

6.17.2.4 #define e 1.6021766208E-19

Elementary Electric Charge - Units: C.

6.17.2.5 #define Faraday 96485.33289

Faraday's Constant - C/mol.

6.17.2.6 #define VolumeSTD 15.17

Standard Segment Volume - cm^3/mol.

6.17.2.7 #define AreaSTD 2.5E5

Standard Segment Area - m^2/mol.

6.17.2.8 #define CoordSTD 10

Standard Coordination Number.

6.17.2.9 #define LengthFactor(z, r, s) (((z/2.0)*(r-s)) - (r-1.0))

Calculation of the Length Factor Parameter in UNIQUAC.

6.17.2.10 #define VacuumPermittivity 8.8541878176E-12

Vacuum Permittivity Constant - F/m or C/V/m.

6.17.2.11 #define WaterRelPerm 80.1

Approximate Relative Permittivity for water - Unitless.

6.17.2.12 #define AbsPerm(Rel) (Rel*VacuumPermittivity)

Calculation of Absolute Permittivity of a medium - F/m or C/V/m.

6.17.3 Typedef Documentation

6.17.3.1 typedef struct SHARK_DATA SHARK_DATA

Data structure for SHARK simulations.

C-style object holding data and function pointers associated with solving aqueous speciation and reaction kinetics. This object couples all other objects available in shark.h in order to provide residual calculations for each individual function that makes up the overall system model. Those residuals are brought together inside the residual function and fed into the lark.h PJFNK solver routine. That solver then attempts to find a solution to all non-linear variables simultaneously. Any function or data pointers in this structure can be overriden to change how you interface with and solve the problem. Users may also provide a set of custom residual functions through the "OtherList" vector object. Those residual function must all have the same format.

6.17.4 Enumeration Type Documentation

6.17.4.1 enum valid mb

Enumeration for the list of valid activity models for non-ideal solutions.

Note

The SIT and PITZER models are not currently supported.

Enumerator

BATCH

CSTR

PFR

```
6.17.4.2 enum valid_act
```

Enumeration for the list of valid activity models for non-ideal solutions.

Note

The SIT and PITZER models are not currently supported.

Enumerator

IDEAL
DAVIES
DEBYE_HUCKEL
SIT
PITZER

6.17.4.3 enum valid surf act

Enumeration for the list of valid surface activity models for non-ideal adsorption.

Note

We had to create an IDEAL_ADS option to replace the IDEAL enum already in use for non-ideal solution or aqueous phases. (ADS => adsorption)

Enumerator

IDEAL_ADS FLORY_HUGGINS UNIQUAC_ACT

6.17.5 Function Documentation

```
6.17.5.1 void print2file_shark_info ( SHARK_DATA * shark_dat )
```

Function to print out simulation conditions and options to the output file.

```
6.17.5.2 void print2file_shark_header ( SHARK_DATA * shark_dat )
```

Function to print out the head of species and time stamps to the output file.

```
6.17.5.3 void print2file_shark_results_new ( SHARK_DATA * shark_dat )
```

Function to print out the simulation results for the current time step.

```
6.17.5.4 void print2file_shark_results_old ( SHARK_DATA * shark_dat )
```

Function to print out the simulation results for the previous time step.

6.17.5.5 double calculate ionic strength (const Matrix < double > & x, MasterSpeciesList & MasterList)

Function to calculate the ionic strength of the solution.

This function calculates the ionic strength of a system given the concentrations of the species present in solution, as well as any other relavent information from SHARK_DATA such as charge.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
MasterList	reference to the MasterSpeciesList object holding species information

6.17.5.6 int FloryHuggins (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for simple non-ideal adsorption (for adsorption reaction object)

This is a simple surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's only parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. NOTE: Only for AdsorptionReaction!

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the AdsorptionReaction object holding parameter information

6.17.5.7 int FloryHuggins_unsteady (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for simple non-ideal adsorption (for unsteady adsorption object)

This is a simple surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's only parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. NOTE: Only for UnsteadyAdsorption!

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the UnsteadyAdsorption object holding parameter information

6.17.5.8 int FloryHuggins_multiligand (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for simple non-ideal adsorption (for multiligand adsorption object)

This is a simple surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's only parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. NOTE: Only for MultiligandAdsorption!

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the MultiligandAdsorption object holding parameter information

6.17.5.9 int FloryHuggins_chemi (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for simple non-ideal adsorption (for chemisorption reaction object)

This is a simple surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's only parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. NOTE: Only for ChemisorptionReaction!

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the ChemisorptionReaction object holding parameter information

6.17.5.10 int FloryHuggins_multichemi (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for simple non-ideal adsorption (for multiligand chemisorption object)

This is a simple surface activity model to be used with the Chemisorption objects to evaluate the non-ideal behavoir of the surface phase. The model's only parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Chemisorption Object itself as the const void *data structure. NOTE: Only for MultiligandChemisorption!

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the MultiligandChemisorption object holding parameter information

6.17.5.11 int UNIQUAC (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for adsorption reaction object)

This is a more complex surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's primary parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. However, future development may require some additional parameters, which will be accessed later. NOTE: Only for AdsorptionReaction!

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the AdsorptionReaction object holding parameter information

6.17.5.12 int UNIQUAC_unsteady (const Matrix < double > & r, Matrix < double > & F, const void * data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for unsteady adsorption object)

This is a more complex surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's primary parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. However, future development may require some additional parameters, which will be accessed later. NOTE: Only for UnsteadyAdsorption!

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the UnsteadyAdsorption object holding parameter information

6.17.5.13 int UNIQUAC_multiligand (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for multiligand adsorption object)

This is a more complex surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's primary parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. However, future development may require some additional parameters, which will be accessed later. NOTE: Only for MultiligandAdsorption!

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the MultiligandAdsorption object holding parameter information

6.17.5.14 int UNIQUAC_chemi (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for chemisorption reaction object)

This is a more complex surface activity model to be used with the Adsorption objects to evaluate the non-ideal behavoir of the surface phase. The model's primary parameters are the shape factors in adsorption and the relative concentrations of each surface species. Therefore, we will pass the Adsorption Object itself as the const void *data structure. However, future development may require some additional parameters, which will be accessed later. NOTE: Only for ChemisorptionReaction!

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the ChemisorptionReaction object holding parameter information

6.17.5.15 int UNIQUAC_multichemi (const Matrix < double > & x, Matrix < double > & F, const void * data)

Surface Activity function for the UNIQUAC model for non-ideal adsorption (for multiligand chemisorption object)

This is a more complex surface activity model to be used with the Chemisorption objects to evaluate the non-ideal behavoir of the surface phase. The model's primary parameters are the shape factors in adsorption and the relative

concentrations of each surface species. Therefore, we will pass the Chemisorption Object itself as the const void *data structure. However, future development may require some additional parameters, which will be accessed later. NOTE: Only for MultiligandChemisorption!

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to the MultiligandChemisorption object holding parameter information

6.17.5.16 intideal_solution (const Matrix < double > & x, Matrix < double > & F, const void * data)

Activity function for Ideal Solution.

This is one of the default activity models available. It assumes the system behaves ideally and sets the activity coefficients to 1 for all species.

Parameters

X	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to a data structure needed to evaluate the activity model

6.17.5.17 int Davies_equation (const Matrix < double > & x, Matrix < double > & F, const void * data)

Activity function for Davies Equation.

This is one of the default activity models available. It uses the Davies semi-empirical model to calculate average activities of each species in solution. This model is typically valid for systems involving high ionic strengths upto 0.5 M (mol/L).

Parameters

Χ	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to a data structure needed to evaluate the activity model

6.17.5.18 int DebyeHuckel_equation (const Matrix < double > & x, Matrix < double > & F, const void * data)

Activity function for Debye-Huckel Equation.

This is one of the default activity models available. It uses the Debye-Huckel limiting model to calculate average activities of each species in solution. This model is typically valid for systems involving low ionic strengths and is only good for solutions between 0 and 0.01 M.

Parameters

Х	matrix of the log(C) concentration values at the current non-linear step
F	matrix of activity coefficients that are to be altered by this function
data	pointer to a data structure needed to evaluate the activity model

6.17.5.19 int surf_act_choice (const std::string & input)

First test of SIT Model.

Function takes a given string and returns a flag denoting which surface activity model was choosen This function returns an integer flag that will be one of the valid surface activity model flags from the valid_surf_act enum. If the input string was not recognized, then it defaults to returning the IDEAL_ADS flag.

Parameters

input	string for the name of the surface activity model
-------	---

6.17.5.20 int act_choice (const std::string & input)

Function takes a given string and returns a flag denoting which activity model was choosen.

This function returns an integer flag that will be one of the valid activity model flags from the valid_act enum. If the input string was not recognized, then it defaults to returning the IDEAL flag.

Parameters

|--|

6.17.5.21 int reactor_choice (const std::string & input)

Function takes a give string and returns a flag denoting which type of reactor was choosen for the system.

This function returns an integer flag that will be one of the valid reactor type flags from the valid_mb enum. If the input string was not recognized, then it defaults to returning the BATCH flag.

Parameters

input	string for the name of the activity model

6.17.5.22 bool linesearch_choice (const std::string & input)

Function returns a bool to determine the form of line search requested.

This function returns true if the user requests a bouncing line search algorithm and false if the user wants a standard line search. If the input string is unrecognized, then it returns false.

Parameters

input string for the line search method

6.17.5.23 int linearsolve_choice (const std::string & input)

Function returns the linear solver flag for the PJFNK method.

This function takes in a string argument and returns the integer flag for the appropriate linear solver in PJFNK. If the input string was unrecognized, then it returns the GMRESRP flag.

Parameters

input string for the linear solver method option
--

6.17.5.24 int Convert2LogConcentration (const Matrix < double > & x, Matrix < double > & logx)

Function to convert the given values of variables (x) to the log of those variables (logx)

This function returns an integer flag to denote success of failure. It takes a constant matrix argument x and replaces the elements of the matrix logx with the base 10 log of those x values. This is used mainly to convert a set of concentrations (x) to their respective log(C) values (logx).

Parameters

X	matrix of values to take the base 10 log of
logx	matrix whose entries are to be changed to base 10 log(x)

6.17.5.25 int Convert2Concentration (const Matrix < double > & logx, Matrix < double > & x)

Function to convert the given log values of variables (logx) to the values of those variables (x)

This function returns an integer flag to denote success of failure. It takes a constant matrix argument logx and replaces the elements of the matrix x with 10° logx. This is used mainly to convert a set of log(C) values (logx) to their respective concentration values (x).

Parameters

logx	matrix of values to apply as the power of 10 (i.e., 10^logx)
Χ	matrix whose entries are to be changed to the result of 10^logx

6.17.5.26 int read_scenario (SHARK DATA * shark_dat)

Function to go through the yaml object for the scenario document.

This function checks the yaml object for the expected keys and values of the scenario document to setup the shark simulation for the input given in the input file.

6.17.5.27 int read_multiligand_scenario (SHARK_DATA * shark_dat)

Function to go through the yaml object to setup memory space for multiligand objects.

This function checks the yaml object for the expected keys and values of the multiligand scenario documents to setup the shark simulation for the input given in the input file.

6.17.5.28 int read_multichemi_scenario (SHARK_DATA * shark_dat)

Function to go through the yaml object to setup memory space for multiligand chemisorption objects.

This function checks the yaml object for the expected keys and values of the multiligand chemisorption scenario documents to setup the shark simulation for the input given in the input file.

```
6.17.5.29 int read_options ( SHARK_DATA * shark_dat )
```

Function to go through the yaml object for the solver options document.

This function checks the yaml object for the expected keys and values of the solver options document to setup the shark simulation for the input given in the input file.

```
6.17.5.30 int read_species ( SHARK_DATA * shark_dat )
```

Function to go through the yaml object for the master species document.

This function checks the yaml object for the expected keys and values of the master species document to setup the shark simulation for the input given in the input file.

```
6.17.5.31 int read_massbalance ( SHARK_DATA * shark_dat )
```

Function to go through the yaml object for the mass balance document.

This function checks the yaml object for the expected keys and values of the mass balance document to setup the shark simulation for the input given in the input file.

```
6.17.5.32 int read_equilrxn ( SHARK_DATA * shark_dat )
```

Function to go through the yaml object for the equilibrium reaction document.

This function checks the yaml object for the expected keys and values of the equilibrium reaction document to setup the shark simulation for the input given in the input file.

```
6.17.5.33 int read_unsteadyrxn ( SHARK_DATA * shark_dat )
```

Function to go through the yaml object for the unsteady reaction document.

This function checks the yaml object for the expected keys and values of the unsteady reaction document to setup the shark simulation for the input given in the input file.

```
6.17.5.34 int read_adsorbobjects ( SHARK_DATA * shark_dat )
```

Function to go through the yaml object for each Adsorption Object.

This function checks the yaml object for the expected keys and values of the adsorption object documents to setup the shark simulation for the input given in the input file.

Note

Each adsorption object will have its own document header by the name of that object

```
6.17.5.35 int read_unsteadyadsorbobjects ( SHARK DATA * shark_dat )
```

Function to go through the yaml object for each Unsteady Adsorption Object.

This function checks the yaml object for the expected keys and values of the unsteady adsorption object documents to setup the shark simulation for the input given in the input file.

Note

Each unsteady adsorption object will have its own document header by the name of that object

6.17.5.36 int read_multiligandobjects (SHARK_DATA * shark_dat)

Function to go through the yaml object for each MultiligandAdsorption Object.

This function checks the yaml object for the expected keys and values of the multiligand object documents to setup the shark simulation for the input given in the input file.

Note

Each ligand object will have its own document header by the name of that object

6.17.5.37 int read_chemisorbobjects (SHARK_DATA * shark_dat)

Function to go through the yaml object for each Chemisorption Object.

This function checks the yaml object for the expected keys and values of the chemisorption object documents to setup the shark simulation for the input given in the input file.

Note

Each adsorption object will have its own document header by the name of that object

6.17.5.38 int read_multichemiobjects (SHARK DATA * shark_dat)

Function to go through the yaml object for each MultiligandChemisorption Object.

This function checks the yaml object for the expected keys and values of the multiligand chemisorption object documents to setup the shark simulation for the input given in the input file.

Note

Each ligand object will have its own document header by the name of that object

6.17.5.39 int setup_SHARK_DATA (FILE * file, int(*)(const Matrix< double > &x, Matrix< double > &res, const void *data) residual, int(*)(const Matrix< double > &x, Matrix< double > &gama, const void *data) activity, int(*)(const Matrix< double > &r, Matrix< double > &p, const void *data) precond, SHARK_DATA * dat, const void * activity_data, const void * residual_data, const void * precon_data, const void * other_data)

Function to setup the memory and pointers for the SHARK_DATA structure for the current simulation.

This function will be called after reading the scenario file and is used to setup the memory and other pointers for the user requested simulation. This function must be called before running a simulation or trying to read in the remander of the yaml formatted input file. Options may be overriden manually after calling this function.

Parameters

file	pointer for the output file where shark results will be stored
residual	pointer to the residual function that will be fed into the PJFNK solver
activity	pointer to the activity function that will determine the activity coefficients
precond	pointer to the linear preconditioning operation to be applied to the Jacobian
dat	pointer to the SHARK_DATA data structure
activity_data	optional pointer for data needed in activity functions
residual_data	optional pointer for data needed in residual functions
Generated by Doxyger precon_data	optional pointer for data needed in preconditioning functions
other_data	optional pointer for data needed in the evaluation of user defined residual functions

6.17.5.40 int shark_add_customResidual (int i, double(*)(const Matrix< double > &x, SHARK_DATA *shark_dat, const void *other_data) other_res, SHARK_DATA * shark_dat)

Function to add user defined custom residual functions to the OtherList vector object in SHARK_DATA.

This function will need to be used if the user wants to include custom residuals into the system via the OtherList object in SHARK_DATA. For each i residual you want to add, you must call this function passing your residual function and the SHARK_DATA structure pointer. The order that those functions are executed in are determined by the integer i.

Parameters

i	index that the other_res function will appear at in the OtherList object
other_res	function pointer for the user's custom residual function
shark_dat	pointer to the SHARK_DATA data structure

6.17.5.41 int shark_parameter_check (SHARK_DATA * shark_dat)

Function to check the Reaction and UnsteadyReaction objects for missing info.

This function checks the Reaction and UnsteadyReaction objects for missing information. If information is missing, this function will return an error that will cause the program to force quit.

6.17.5.42 int shark_energy_calculations (SHARK_DATA * shark_dat)

Function to calculate all Reaction and UnsteadyReaction energies.

This function will call the calculate energy functions for Reaction and UnsteadyReaction objects.

6.17.5.43 int shark_temperature_calculations (SHARK_DATA * shark_dat)

Function to calculate all Reaction and UnsteadyReaction parameters as a function of temperature.

This function will call all temperature dependent functions in Reaction and UnsteadyReaction to calculate equilibirium and reaction rate parameters as a function of system temperature.

6.17.5.44 int shark_pH_finder (SHARK_DATA * shark_dat)

Function will search MasterSpeciesList for existance of H + (aq) and OH - (aq) molecules.

This function searches all molecules in the MasterSpeciesList object for the H + (aq) and OH - (aq) molecules. If they are found, then it sets the pH_index and pOH_index of the SHARK_DATA structure and indicates that the system contains these variables.

6.17.5.45 int shark_guess (SHARK_DATA * shark_dat)

Function provides a rough initial guess for the values of all non-linear variables.

This function constructs an rough initial guess for the values of all non-linear variables in the system. The guess is based primarily off of trying to statisfy all mass balance constraints, initial conditions, and pH constraints if any apply.

```
6.17.5.46 int shark_initial_conditions ( SHARK_DATA * shark_dat )
```

Function to establish the initial conditions of the shark simulation.

This function will establish the initial conditions for a transient problem by solving the speciation of the system while holding the transient/unsteady variables constant at their respective initial values. However, if the system we are trying to solve is steady, then this function just calls the shark_guess function.

```
6.17.5.47 int shark_executioner ( SHARK_DATA * shark_dat )
```

Function to execute a shark simulation at a single time step or pH value.

This function calls the preprocess, solver, and postprocess functions in order. If a particular solve did not converge, then it will retry the solver routine until it runs out of tries or attains convergence.

```
6.17.5.48 int shark_timestep_const ( SHARK_DATA * shark_dat )
```

Function to set up all time steps in the simulation to a specified constant.

This function will set all time steps for the current simulation to a constant that is specified in the input file. The time step will not be changed unless the simulation fails, then it will be reduced in order to try to get the system to converge.

```
6.17.5.49 int shark_timestep_adapt ( SHARK_DATA * shark_dat )
```

Function to set up all time steps in the simulation based on success or failure to converge.

This function will set all time steps for the current simulation based on some factor multiple of the prior time step used and whether or not the previous solution step was successful. If the previous step converged, then the new time step will be 1.5x the old time step. If it failed, then the simulation will be retried with a new time step of 0.5x the old time step.

```
6.17.5.50 int shark_preprocesses ( SHARK_DATA * shark_dat )
```

Function to call other functions for calculation of parameters and setting of time steps.

This function will call the shark_temperature_calculations function and the appropriate time step function. If the user requests a constant time step, it will call the shark_timestep_const function. Otherwise, it calls the shark_timestep_adapt function.

```
6.17.5.51 int shark_solver ( SHARK_DATA * shark_dat )
```

Function to call the PJFNK solver routine given the current SHARK DATA information.

This function will perform the necessary steps before and after calling the PJFNK solver routine. Based on the simulation flags, the solver function will perform an intial guess for unsteady variables, call the PJFNK method, and the printout a console message about the performance. If a terminal failure occurs during the solver, it will print out the current state of residuals, variables, and the Jacobian matrix to the console. Analyzing this information could provide clues as to why failure occured.

```
6.17.5.52 int shark_postprocesses ( SHARK_DATA * shark_dat )
```

Function to convert PJFNK solutions to concentration values and print to the output file.

This function will convert the non-linear variables to their respective concentration values, then print the solve information out to the output file.

```
6.17.5.53 int shark_reset ( SHARK_DATA * shark_dat )
```

Function to reset the values of all stateful information in SHARK DATA.

This function will reset all stateful matrix data in the SHARK_DATA structure in preparation of the next time step simulation.

```
6.17.5.54 int shark_residual ( const Matrix < double > & x, Matrix < double > & F, const void * data )
```

Default residual function for shark evaluations.

This function calls each individual object's residual function to formulate the overall residual function used in the $P \leftarrow JFNK$ solver routine. It will also call the activity function. The order in which these function calls occurs is as follows: (i) activities, (ii) Reaction, (iii) UnsteadyReaction, (iv) MassBalance, (v) OtherList, and (vi) MasterSpeciesList. If a constant pH is specified, then the MasterSpeciesList residual call is replaced with a constraint on the H + (aq) variable (if one exists).

```
6.17.5.55 int SHARK ( SHARK_DATA * shark_dat )
```

Function to call all above functions to perform a shark simulation.

This function is called after reading in all inputs, setting all constants, and calling the setup function. It will call all the necessary functions and subroutines iteratively until the desired simulation is complete.

```
6.17.5.56 int SHARK_SCENARIO ( const char * yaml_input )
```

Function to perform a shark simulation based on the conditions in a yaml formatted input file.

This is the primary function used to run shark simulations from the UI. It requires that the user provide one input file that is formatted with yaml keys, symbols, and spacing so that it can be recognized by the parser. This style of input file is much easier to use and understand than the input files used for SCOPSOWL or SKUA. Below shows an example of a typical input file. Note that the # symbol is used in the input file to comment out lines of text that the parser does not need to read.

Example Yaml Input for SHARK

```
#This will serve as a test input file for shark to demo how to structure the document #In practice, this section should be listed first, but it doesn't really matter #DO NOT USE TABS IN THESE INPUT FILES #— Starts a document ... Ends a document #All keys must be proceeded by a : #All lists/header must be preceeded by a - #Spacing of the keys will indicate which list/header they belong to Scenario:
```

 vars_fun: numvar: 25 num_ssr: 15 num_mbe: 7 num_usr: 2

num_other: 0 #Not required or used in current version

```
sys_data:
      act fun: davies
      const pH: false
      pH: 7 #Only required if we are specifying a const_pH
      temp: 298.15 #Units must be in Kelvin
      dielec: 78.325 #Units must be in (1/Kelvin)
      rel perm: 80.1 #Unitless number
      res_alk: 0 #Units must be in mol/L (Residual Alkalinity)
      volume: 1.0 #Units must be in L
    · run time:
      steady: false #NOTE: All time must be represented in hours
      specs_curve: false #Only needed if steady = true, and will default to false
      dt: 0.001 #Only required if steady = false
      time_adapt: true #Only needed if steady = false, and will default to false
      sim_time: 96.0 #Only required if steady = false
      t_out: 0.01 #Only required if steady = false
#The following header is entirely optional, but is used to set solver options
SolverOptions:
line_search: true #Default = true, and is recommended to be true
search_type: standard
linear_solve: gmresrp #Note: FOM will be fastest for small problems
restart: 25 #Note: restart only used if using GMRES or GCR type solvers
nl_maxit: 50
nl_abstol: 1e-5
nl reltol: 1e-8
lin_reltol: 1e-10 #Min Tol = 1e-15
lin_abstol: 1e-10 #Min Tol = 1e-15
nl print: true
I print: true
#After the Scenario read, shark will call the setup_function, then read info below
MasterSpecies:
#Header names are specific
#Keys are chosen by user, but must span numbers 0 through numvar-1
#Keys will denote the ordering of the variables
#Note: Currently, the number of reg molecules is very limited
    reg:
      0: CI - (aq)
      1: NaHCO3 (aq)
      2: NaCO3 - (aq)
      3: Na + (aq)
      4: HNO3 (aq)
      5: NO3 - (aq)
      6: H2CO3 (aq)
      7: HCO3 - (aq)
      8: CO3 2- (aq)
      9: UO2 2+ (aq)
```

10: UO2NO3 + (aq)

```
11: UO2(NO3)2 (aq)
 12: UO2OH + (aq)
 13: UO2(OH)3 - (aq)
 14: (UO2)2(OH)2 2+ (aq)
 15: (UO2)3(OH)5 + (aq)
 16: UO2CO3 (aq)
 17: UO2(CO3)2 2- (aq)
 18: UO2(CO3)3 4- (aq)
 19: H2O (I)
 20: OH - (aq)
 21: H + (aq)
 #Keys for the sub-headers must follow same rules as keys from above
· unreg:
    - 22:
      formula: A(OH)2 (aq)
      charge: 0
      enthalpy: 0
       entropy: 0
      have_HS: false
      energy: 0
      have_G: false
      phase: Aqueous
      name: Amidoxime
      lin_form: none
    - 23:
      formula: UO2AO2 (aq)
      charge: 0
       enthalpy: 0
       entropy: 0
      have_HS: false
      energy: 0
      have G: false
      phase: Aqueous
      name: Uranyl-amidoximate
      lin_form: none
    - 24:
      formula: UO2CO3AO2 2- (aq)
      charge: -2
       enthalpy: 0
       entropy: 0
      have_HS: false
       energy: 0
      have_G: false
      phase: Aqueous
      name: Uranyl-carbonate-amidoximate
      lin form: none
      #NOTE: Total concentrations must be given in mol/L
      MassBalance:
       #Header names under MassBalance are choosen by the user
```

#All other keys will be checked

```
water:
 total conc: 1
     – delta:
       "H2O (I)": 1
· carbonate:
 total conc: 0.0004175
     - delta:
       "NaHCO3 (aq)": 1
       "NaCO3 - (aq)": 1
       "H2CO3 (aq)": 1
       "HCO3 - (aq)": 1
       "CO3 2- (aq)": 1
       "UO2CO3 (aq)": 1
       "UO2(CO3)2 2- (aq)": 2
       "UO2(CO3)3 4- (aq)": 3
       "UO2CO3AO2 2- (aq)": 1
       #Other mass balances skipped for demo purposes...
       #Document for equilibrium or steady reactions
       EquilRxn:
       #Headers under EquilRxn separate out each reaction object
       #Keys for these headers only factor into the order of the equations
       #Stoichiometry follows the convention that products are pos(+) and reactants are neg(-)
       #Note: logK is only required if any species in stoichiometry is unregistered
       #Example: below represents - \{H2O(I)\} -> \{H + (aq)\} + \{OH - (aq)\}
       #Note: a valid reaction statement requires at least 1 stoichiometry args
       #Note: You can also provide reaction energies: enthalpy, entropy, and energy
• rxn00:
  logK: -14
     - stoichiometry:
       "H2O (I)": -1
       "OH - (aq)": 1
       "H + (aq)": 1
• rxn01:
 logK: -6.35
     – stoichiometry:
       "H2CO3 (aq)": -1
       "HCO3 - (aq)": 1
       "H + (aq)": 1
       #Other reactions skipped for demo purposes...
       #Document for unsteady reactions
       UnsteadyRxn:
       #Same basic standards for this doc as the EquilRxn
```

```
#Main difference is the inclusion of rate information
       #You are required to give at least 1 rate
       #You are also required to denote which variable is unsteady
       #You must give the initial concentration for the variable in mol/L
       #Rate units are in (L/mol)^n/hr
       #Note: we also have keys for forward ref, reverse ref,
       #activation energy, and temp affinity.
       #These are optional if forward and/or reverse are given
       #Note: You can also provide reaction energies: enthalpy, entropy, and energy
rxn00:
  unsteady_var: UO2AO2 (aq)
  initial condition: 0
  logK: -1.35
  forward: 4.5e+6
  reverse: 1.00742e+8
     - stoichiometry:
       "UO2 2+ (aq)": -1
       "A(OH)2 (aq)": -1
       "UO2AO2 (aq)": 1
       "H + (aq)": 2
rxn01:
  unsteady_var: UO2CO3AO2 2- (aq)
  initial condition: 0
  logK: 3.45
  forward: 2.55e+15
  reverse: 9.04774e+11
    - stoichiometry:
       "UO2 2+ (aq)": -1
       "CO3 2- (aq)": -1
       "A(OH)2 (aq)": -1
       "UO2CO3AO2 2- (aq)": 1
       "H + (aq)": 2
       ...
       Note
```

It may be advantageous to look at some other shark input file examples. More input files are provided in the input_files/SHARK directory of the ecosystem project folder. Please refer to your own source file location for more input file examples for SHARK.

```
6.17.5.57 int SHARK_TESTS ( )
```

Function to perform a series of shark calculation tests.

This function sets up and solves a test problem for shark. It is callable from the UI.

```
6.17.5.58 int SHARK_TESTS_OLD ( )
```

Function to perform a series of shark calculation tests (older version)

This function sets up and solves a test problem for shark. It is NOT callable from the UI.

6.18 skua.h File Reference

Surface Kinetics for Uptake by Adsorption.

```
#include "finch.h"
#include "magpie.h"
#include "egret.h"
```

Classes

struct SKUA_PARAM

Data structure for species' parameters in SKUA.

struct SKUA_DATA

Data structure for all simulation information in SKUA.

Macros

```
    #define SKUA_HPP_
```

```
\bullet \ \ \text{\#define D\_inf}(\mathsf{Dref}, \mathsf{Tref}, \mathsf{B}, \mathsf{p}, \mathsf{T}) \ ( \ \mathsf{Dref} * \mathsf{pow}(\mathsf{p+sqrt}(\mathsf{DBL\_EPSILON}), (\mathsf{Tref/T}) - \mathsf{B}) \ ) \\
```

Empirical correction of diffusivity (um²/hr)

#define D_o(Diff, E, T) (Diff * exp(-E/(Rstd*T)))

Arrhenius Rate Expression for Diffusivity (um²/hr)

#define D_c(Diff, phi) (Diff * (1.0/((1.0+1.1E-6)-phi)))

Approximate Darken Diffusivity Equation (um²/hr)

Functions

void print2file_species_header (FILE *Output, SKUA_DATA *skua_dat, int i)

Function to print out the species' headers to output file.

• void print2file_SKUA_time_header (FILE *Output, SKUA_DATA *skua_dat, int i)

Function to print out time and space headers to output file.

void print2file_SKUA_header (SKUA_DATA *skua_dat)

Function calls the other header functions to establish output file structure.

void print2file_SKUA_results_old (SKUA_DATA *skua_dat)

Function to print out the old time step simulation results to the output file.

void print2file_SKUA_results_new (SKUA_DATA *skua_dat)

Function to print out the new time step simulation results to the output file.

double default_Dc (int i, int I, const void *data)

Default function for surface diffusivity.

double default_kf (int i, const void *data)

Default function for film mass transfer coefficent.

double const Dc (int i, int I, const void *data)

Constant surface diffusivity function.

double simple_darken_Dc (int i, int I, const void *data)

Simple Darken model for surface diffusivity.

double theoretical darken Dc (int i, int I, const void *data)

Theoretical Darken model for surface diffusivity.

double empirical_kf (int i, const void *data)

Empirical function for film mass transfer coefficent.

double const_kf (int i, const void *data)

Constant function for film mass transfer coefficent.

int molefractionCheck (SKUA_DATA *skua_dat)

Function to check mole fractions in gas and solid phases for errors.

• int setup_SKUA_DATA (FILE *file, double(*eval_Dc)(int i, int I, const void *user_data), double(*eval_Kf)(int i, const void *user_data), const void *user_data, MIXED_GAS *gas_data, SKUA_DATA *skua_dat)

Function to setup the function pointers and vector objects in memory to setup the SKUA simulation.

int SKUA_Executioner (SKUA_DATA *skua_dat)

Function to execute preprocesses, solvers, and postprocesses for a SKUA simulation.

int set SKUA ICs (SKUA DATA *skua dat)

Function to establish the initial conditions of adsorption in the adsorbent.

int set_SKUA_timestep (SKUA_DATA *skua_dat)

Function to establish the time step for the current simulation.

int SKUA_preprocesses (SKUA_DATA *skua_dat)

Function to perform the necessary preprocess operations before a solve.

int set_SKUA_params (const void *user_data)

Function to call the diffusivity function during the solve.

int SKUA_postprocesses (SKUA_DATA *skua_dat)

Function to perform the necessary postprocess operations after a solve.

• int SKUA reset (SKUA DATA *skua dat)

Function to reset the stateful information in SKUA after a simulation.

int SKUA (SKUA_DATA *skua_dat)

Function to iteratively call all execution steps to evolve a simulation through time.

• int SKUA_SCENARIOS (const char *scene, const char *sorbent, const char *comp, const char *sorbate)

Function callable from the UI to perform a SKUA simulation based on user supplied input files.

• int SKUA_TESTS ()

Function to perform a test of the SKUA functions and routines.

6.18.1 Detailed Description

Surface Kinetics for Uptake by Adsorption.

skua.cpp

This file contains structures and functions associated with solving the surface diffusion partial differential equations for adsorption kinetics in spherical and/or cylindrical adsorbents. For this system, it is assumed that the pore size is so small that all molecules are confined to movement exclusively on the surface area of the adsorbent. The total amount of adsorption for each species is drive by the MAGPIE model for non-ideal mixed gas adsorption. Spatial and temporal varience in adsorption is caused by a combination of different kinetics between adsorbing species and different adsorption affinities for the surface.

The function for surface diffusion involves four parameters, although not all of these parameters are required to be used. Surface diffusion theoretically varies with temperature according to the Arrhenius rate expression, but we also add in an empirical correction term to account for variations in diffusivity with the partial pressure of the species in the gas phase.

```
D\_surf = D\_ref * exp(-E / (R*T)) * pow(p, (T\_ref/T) - B)
```

D_ref is the Reference Diffusivity (um 2 2/hr), E is the activation energy for adsorption (J/mol), R is the gas law constant (J/K/mol), T is the system temperature (K), p is the partial pressure of the adsorbing species (kPa), T_ref is the Reference Temperature (K), and B is the Affinity constant.

Author

Austin Ladshaw

Date

01/26/2015

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This software was designed and built at the Georgia Institute of Technology by Austin Ladshaw for PhD research in the area of adsorption and surface science. Copyright (c) 2015, all rights reserved.

6.18.2 Macro Definition Documentation

6.18.2.1 #define SKUA_HPP_

6.18.2.2 #define D_inf(Dref, Tref, B, p, T) ($Dref * pow(p+sqrt(DBL_EPSILON),(Tref/T)-B)$)

Empirical correction of diffusivity (um²/hr)

6.18.2.3 #define D_o(Diff, E, T) (Diff $* \exp(-E/(Rstd*T))$)

Arrhenius Rate Expression for Diffusivity (um²/hr)

6.18.2.4 #define D_c(Diff, phi) (Diff * (1.0/((1.0+1.1E-6)-phi)))

Approximate Darken Diffusivity Equation (um²/hr)

6.18.3 Function Documentation

6.18.3.1 void print2file_species_header (FILE * Output, SKUA_DATA * skua_dat, int i)

Function to print out the species' headers to output file.

6.18.3.2 void print2file_SKUA_time_header (FILE * Output, SKUA_DATA * skua_dat, int i)

Function to print out time and space headers to output file.

6.18.3.3 void print2file_SKUA_header (SKUA_DATA * skua_dat)

Function calls the other header functions to establish output file structure.

6.18.3.4 void print2file_SKUA_results_old (SKUA_DATA * skua_dat)

Function to print out the old time step simulation results to the output file.

6.18.3.5 void print2file_SKUA_results_new (SKUA_DATA * skua_dat)

Function to print out the new time step simulation results to the output file.

6.18.3.6 double default_Dc (int i, int l, const void * data)

Default function for surface diffusivity.

This is the default function provided by SKUA for the calculation of the surface diffusivity parameter. The diffusivity is calculated based on the Arrhenius rate expression, then corrected for using the empirical correction term with the outside partial pressure of the gas species.

Parameters

i	index of the gas/adsorbed phase species that this function acts on
1	index of the node in the spatial discretization that this function acts on
data	pointer to the SKUA_DATA structure

6.18.3.7 double default_kf (int i, const void * data)

Default function for film mass transfer coefficent.

This is the default function provided by SKUA for the calculation of the film mass transfer parameter. By default, we are usually going to couple the SKUA model with a pore diffusion model (see scopsowl.h). Therefore, the film mass transfer coefficient would be zero, because we would only consider a Dirichlet boundary condition for this sub-problem.

Parameters

i	index of the gas/adsorbed phase species that this function acts on
data	pointer to the SKUA_DATA structure

6.18.3.8 double const_Dc (int i, int l, const void * data)

Constant surface diffusivity function.

This function allows the user to specify just a single constant value for surface diffusivity. The value of diffusivity applied at all nodes will be the ref_diffusion parameter in SKUA_PARAM.

Parameters

i	index of the gas/adsorbed phase species that this function acts on
1	index of the node in the spatial discretization that this function acts on
data	pointer to the SKUA_DATA structure

6.18.3.9 double simple_darken_Dc (int i, int I, const void * data)

Simple Darken model for surface diffusivity.

This function uses an approximation to Darken's model for surface diffusion. The approximation is exact if the isotherm for adsorption takes the form of the Langmuir model, but is only approximate if the isotherm is heterogeneous. Forming the approximation in this manner is significantly cheaper than forming the true Darken model expression for the GSTA isotherm.

Parameters

i	index of the gas/adsorbed phase species that this function acts on
1	index of the node in the spatial discretization that this function acts on
data	pointer to the SKUA_DATA structure

6.18.3.10 double theoretical_darken_Dc (int i, int I, const void * data)

Theoretical Darken model for surface diffusivity.

This function uses the full theoretical expression of the Darken's diffusion model to calculate the surface diffusivity. This calculation involves formulating the reference state pressures for the adsorbed amount at every node, then calculating derivatives of the adsorption isotherm for each species. It is more accurate than the simple Darken model function, but costs significantly more computational time.

Parameters

i	index of the gas/adsorbed phase species that this function acts on
1	index of the node in the spatial discretization that this function acts on
data	pointer to the SKUA_DATA structure

6.18.3.11 double empirical_kf (int i, const void * data)

Empirical function for film mass transfer coefficent.

This function provides an empirical estimate of the mass transfer coefficient using the gas velocity, molecular diffusivities, and dimensionless numbers (see egret.h). It is used as the default film mass transfer function IF the boundary condition is specified to be a Neumann type boundary by the user.

Parameters

i	index of the gas/adsorbed phase species that this function acts on
data	pointer to the SKUA_DATA structure

6.18.3.12 double const_kf (int i, const void * data)

Constant function for film mass transfer coefficent.

This function allows the user to specify a constant value for the film mass transfer coefficient. The value of the film mass transfer coefficient will be the value of film transfer given in the SKUA PARAM data structure.

Parameters

i	index of the gas/adsorbed phase species that this function acts on
data	pointer to the SKUA_DATA structure

6.18.3.13 int molefractionCheck (SKUA_DATA * skua_dat)

Function to check mole fractions in gas and solid phases for errors.

This function is called after reading input and before calling the primary solution routines. It will force and error and quit the program if their are inconsistencies in the mole fractions it was given. All mole fractions must sum to 1, otherwise there is missing information.

6.18.3.14 int setup_SKUA_DATA (FILE * file, double(*)(int i, int I, const void *user_data) eval_Dc, double(*)(int i, const void *user_data) eval_Kf, const void * user_data, MIXED_GAS * gas_data, SKUA_DATA * skua_dat)

Function to setup the function pointers and vector objects in memory to setup the SKUA simulation.

This function is called to setup the SKUA problem in memory and set function pointers to either defaults or user specified functions. It must be called prior to calling any other SKUA function and will report an error if the object was not setup properly.

Parameters

file	pointer to the output file for SKUA simulations	
eval_Dc	pointer to the function to evaluate the surface diffusivity	
eval_Kf	pointer to the function to evaluate the film mass transfer coefficient	
user_data	pointer to a user defined data structure used in the calculation the the parameters	
gas_data	pointer to the MIXED_GAS data structure for egret.h calculations	
skua_dat	pointer to the SKUA_DATA data structure	

6.18.3.15 int SKUA_Executioner (SKUA_DATA * skua_dat)

Function to execute preprocesses, solvers, and postprocesses for a SKUA simulation.

This function calls the preprocess, solver, and postprocess functions to complete a single time step in a SKUA simulation. User's will want to call this function whenever a time step simulation result is needed. This is used primarily when coupling with other models (see scopsowl.h).

6.18.3.16 int set_SKUA_ICs (SKUA_DATA * skua_dat)

Function to establish the initial conditions of adsorption in the adsorbent.

This function needs to be called before doing any simulation or execution of a time step, but only once per simulation. It sets the value of adsorption for each adsorbable species to the specified initial values given via qT and xIC in SKUA DATA.

6.18.3.17 int set_SKUA_timestep (SKUA_DATA * skua_dat)

Function to establish the time step for the current simulation.

This function is called to set a time step value for a particular simulation step. By default, the time step is set to (1/4)x space step size. If you need to change the step size, you must do so manually.

6.18.3.18 int SKUA_preprocesses (SKUA_DATA * skua_dat)

Function to perform the necessary preprocess operations before a solve.

This function performs preprocess operations prior to calling the solver routine. Those preprocesses include establishing boundary conditions and performing a MAGPIE simulation for the adsorption on the surface (see magpie.h).

6.18.3.19 int set_SKUA_params (const void * user_data)

Function to call the diffusivity function during the solve.

This is the function passed into FINCH to be called during the FINCH solver (see finch.h). It will call the diffusion functions set by the user in the setup function above. This is not overridable.

6.18.3.20 int SKUA_postprocesses (SKUA_DATA * skua_dat)

Function to perform the necessary postprocess operations after a solve.

This function performs postprocess operations after a solve was completed successfully. Those operations include estimating average total adsorption, average adsorbed mole fractions, and heat of adsorption for each species. Results are then printed to the output file.

```
6.18.3.21 int SKUA_reset ( SKUA_DATA * skua_dat )
```

Function to reset the stateful information in SKUA after a simulation.

This function sets all the old state data to the newly formed state data. It needs to be called after a successful execution of the simulation step and before calling for the next time step to be solved. Do not call out of turn, otherwise information will be lost.

```
6.18.3.22 int SKUA ( SKUA_DATA * skua_dat )
```

Function to iteratively call all execution steps to evolve a simulation through time.

This function is used in conjunction with the scenario call from the UI to numerically solve the adsorption kinetics problem in time. It will call the initial conditions function once, then iteratively call the reset, time step, and executioner functions for SKUA to push the simulation forward in time. This function will be called from the SKUA_SCENARIOS function.

```
6.18.3.23 int SKUA_SCENARIOS ( const char * scene, const char * sorbent, const char * comp, const char * sorbate )
```

Function callable from the UI to perform a SKUA simulation based on user supplied input files.

This is the primary function to be called when running a stand-alone SKUA simulation. Parameters and system information for the simulation are given in a series of input files that come in as character arrays. These inputs are all required to call this function.

Parameters

scene	Sceneario Input File
sorbent	Adsorbent Input File
comp	Component Input File
sorbate	Adsorbate Input File

Note

Each input file has a particular format that must be strictly adhered to in order for the simulation to be carried out correctly. The format for each input file, and an example, is provided below...

Scenario Input Format

System Temperature (K) [tab] Total Pressure (kPa) [tab] Gas Velocity (cm/s) Simulation Time (hrs) [tab] Print Out Time (hrs)
BC Type (0 = Neumann, 1 = Dirichlet)
Number of Gas Species
Initial Total Adsorption (mol/kg)

Name of ith Species [tab] Adsorbable? (0 = false, 1 = true) [tab] Gas Phase Molefraction [tab] Initial Sorbed Molefraction (repeat above for all species)

Example Scenario Input

353.15 101.35 0.36 4.0 0.05 0 5 0.0 N2 0 0.7634 0.0 O2 0 0.2081 0.0 Ar 0 0.009 0.0 CO2 0 0.0004 0.0

Above example is for a 5-component mixture of N2, O2, Ar, CO2, and H2O, but we are only considering the H2O as adsorbable.

Adsorbent Input File

H2O 1 0.0191 0.0

Domain Coord. (2 = spherical, 1 = cylindrical) { [tab] Char. Length (um) (i.e., cylinder length) } (NOTE: Char. Length is only needed if problem is not spherical)
Pellet Radius (um)

Example Adsorbent Input

1 6.0

Above example is for a cylindrical adsorbent with a length of 5 um and radius of 2 um.

Component Input File

Molar Weight of ith species (g/mol) [tab] Specific Heat of ith species (J/g/K) Sutherland Viscosity (g/cm/s) [tab] Sutherland Temperature (K) [tab] Sutherland Constant (K) of ith species (repeat above for all species in same order they appeared in the Scenario Input File)

Example Component Input

28.016 1.04 0.0001781 300.55 111.0 32.0 0.919 0.0002018 292.25 127.0 39.948 0.522 0.0002125 273.11 144.4 44.009 0.846 0.000148 293.15 240.0 18.0 1.97 0.0001043 298.16 784.72

Above example is a continuation of the Scenario Input example wherein each grouping represents parameters that are associated with N2, O2, Ar, CO2, and H2O, respectively. The order is VERY important!

Adsorbate Input File

Type of Surface Diffusion Function (0 = constant, 1 = simple Darken, 2 = theoretical Darken) Reference Diffusivity (um^2/hr) [tab] Activation Energy (J/mol) of ith adsorbable species Reference Temperature (K) [tab] Affinity Constant (-) of ith adsorbable species van der Waals Volume (cm^3/mol) of ith species GSTA adsorption capacity (mol/kg) of ith species Number of GSTA parameters of ith species Enthalpy (J/mol) of nth site [tab] Entropy of nth site (J/K/mol) of ith species (repeat enthalpy and entropy for all n sites in species i) (repeat above for all species i)

Example Adsorbate Input

```
0
0.8814 0.0
267.999 0.0
13.91
11.67
4
-46597.5 -53.6994
-125024 -221.073
-193619 -356.728
-272228 -567.459
1.28 540.1
374.99 0.01
3.01
1.27
2
-46597.5 -53.6994
-125024 -221.073
```

Above example would be for a simulation involving two adsorbable species using a constant surface diffusion function. Each adsorbable species has it's own set of kinetic and equilibrium parameters that must be given in the same order as the species appeared in the Scenario Input. Note: we do not need to supply this information for non-adsorbable species.

```
6.18.3.24 int SKUA_TESTS ( )
```

Function to perform a test of the SKUA functions and routines.

This function is callable from the UI and will perform a test simulation of the SKUA system of equations. Results from that test are output into a sub-directory called output and named SKUA_Test_Output.txt.

6.19 skua_opt.h File Reference

Optimization Routine for the SKUA Model.

```
#include "skua.h"
```

Classes

struct SKUA_OPT_DATA

Data structure for the SKUA Optimization Routine.

Functions

• int SKUA_OPT_set_y (SKUA_OPT_DATA *skua_opt)

Function to set the rest of the gas phase mole fractions based on current mole fraction of adsorbing gas.

int initial_guess_SKUA (SKUA_OPT_DATA *skua_opt)

Function to set up an initial guess for the surface diffusivity parameter in SKUA.

void eval_SKUA_Uptake (const double *par, int m_dat, const void *data, double *fvec, int *info)

Function that works in conjunction with the Imfit routine to minimize the euclidean norm between function and data.

• int SKUA_OPTIMIZE (const char *scene, const char *sorbent, const char *comp, const char *sorbate, const char *data)

Function called to perform the optimization routine given a specific set of information and data.

6.19.1 Detailed Description

Optimization Routine for the SKUA Model.

skua_opt.cpp

This file contains structures and functions associated with performing non-linear least-squares optimization of the SKUA simulation results against actual kinetic adsorption data. The optimization routine here allows you to run data comparisons and optimizations in three forms: (i) Rough optimizations - cheaper operations, but less accurate, (ii) Exact optmizations - much more expensive, but greater accuracy, and (iii) data/model comparisons - no optimization, just using system parameters to compare simulation results agains a set of data.

Depending on the level of optimization desired, this routine could take several minutes or several hours. The optimization/comparisons are printed out in two files: (i) a parameter file, which contains the simulation partial pressures and temperatures and the optimized diffusivities with the euclidean norm of the fitting and (ii) a comparison file that shows the model value and data value at each time step for each kinetic curve.

The optimized diffusion parameters are given for each individual kinetic data curve. Each data curve will have a different pairing of partial pressure and temperature. Because of this, you will get a list of different diffusivities for each data curve. To get the optimum kinetic parameters from this list of diffusivities, you must fit the diffusion parameter values to the following diffusion function model...

```
D opt = D ref * exp(-E / (R*T)) * pow(p, (T ref/T) - B)
```

where D_ref is the Reference Diffusivity (um^2/hr), E is the activation energy for adsorption (J/mol), R is the gas law constant (J/K/mol), T is the system temperature (K), p is the partial pressure of the adsorbing species (kPa), T_ref is the Reference Temperature (K), and B is the Affinity constant. This algorithm does not automatically produce these parameters for you, but gives you everything you need to produce them yourself.

This routine allows you to optimize multiple kinetic curves at one time. However, all data must be for the same adsorbent-adsorbate system. In other words, the adsorbent and adsorbate pair must be the same for each kinetic curve analyzed. Also, each experiment must have been done in a thin bed or continuous flow system where the adsorbents were exposed to a nearly constant outside partial pressure for all time steps and the gas velocity of that system is assumed constant for all experiments. This experimental setup is very typical for studying adsorption kinetics for gas-solid systems.

Author

Austin Ladshaw

Date

05/11/2015

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This software was designed and built at the Georgia Institute of Technology by Austin Ladshaw for PhD research in the area of adsorption and surface science. Copyright (c) 2015, all rights reserved.

6.19.2 Function Documentation

```
6.19.2.1 int SKUA_OPT_set_y ( SKUA_OPT_DATA * skua_opt )
```

Function to set the rest of the gas phase mole fractions based on current mole fraction of adsorbing gas.

This function takes the current mole fraction of the adsorbing gas and calculates the gas mole fractions of the other gases in the sytem based on the standard inlet gas composition given in the scenario file.

```
6.19.2.2 int initial_guess_SKUA ( SKUA_OPT_DATA * skua_opt )
```

Function to set up an initial guess for the surface diffusivity parameter in SKUA.

This function performs the Rough optimization on the surface diffusivity based on the idea of reducing or eliminating function bias between data and simulation. A positive function bias means that the simulation curve is "higher" than the data curve and a negative function bias means that the simulation curve is "lower" than the data curve. We use this information to incrementally adjust the rate of surface diffusion until this bias is near zero. When bias is near zero, the simulation is nearly optimized, but further refinement may be necessary to find the true minimum solution.

```
6.19.2.3 void eval_SKUA_Uptake ( const double * par, int m_dat, const void * data, double * fvec, int * info )
```

Function that works in conjunction with the Imfit routine to minimize the euclidean norm between function and data.

This function will run the SKUA simulation at a given value of surface diffusivity and produce residuals that feed into the Levenberg-Marquardt's algorithm for non-linear least-squares regression. The form of this function is specific to the format required by the Imfit routine.

Parameters

par	array of parameters that are to be optimized
m_dat	number of data points or functions to evaluate
data	user supplied data structure holding information necessary to form the residuals
fvec	array of residuals computed at the current parameter values
info	integer pointer denoting whether or not the user requests to end a particular simulation

6.19.2.4 int SKUA_OPTIMIZE (const char * scene, const char * sorbent, const char * comp, const char * sorbate, const char * data)

Function called to perform the optimization routine given a specific set of information and data.

This is the function that is callable by the UI. The user must provide 5 input files to the routine in order to establish simulation conditions, adsorbent properties, component properties, adsorbate equilibrium parameters, and the set of data that we are comparing the simulations to. Each input file has a very specific structure and order to the information that it contains. The structure here is DIFFERENT than the structure for just running standard SKUA simulations (see skua.h).

Parameters

scene	Sceneario Input File
sorbent	Adsorbent Input File
comp	Component Input File
sorbate	Adsorbate Input File
data Generated by	Kinetic Adsorption Data File

Note

Much of the structure of these input files are "similar" to that of the input files used in SKUA_SCENARIOS (see skua.h), but with some notable differences. Below gives the format for each input file with an example. Make sure your input files follow this format before calling this routine from the UI.

Scenario Input File

```
Optimization? (0 = false, 1 = true) [tab] Rough Optimization? (0 = false, 1 = true)

Surf. Diff. (0 = constant, 1 = simple Darken, 2 = theoretical Darken) [tab] BC Type (0 = Neumann, 1 = Dirichlet)

Total Pressure (kPa) [tab] Gas Velocity (cm/s)
```

Number of Gaseous Species

Initial Adsorption Total (mol/kg)

Name [tab] Adsorbable? (0 = false, 1 = true) [tab] Inlet Gas Mole Fraction [tab] Initial Adsorbed Mole Fraction (NOTE: The above line is repeated for all species in gas phase. Also, this algorithm only allows you to consider one adsorbable gas component. Inlet gas mole fractions must be non-zero for all non-adsorbing gases and must sum to 1.)

Example Scenario Input

1 0 0 0 101.35 0.36 5 0.0 N2 0 0.7825 0.0 O2 0 0.2081 0.0 Ar 0 0.009 0.0 CO2 0 0.0004 0.0 H2O 1 0.0 0.0

Above example is for running optimizations on data collected with a gas stream at 0.36 cm/s with 5 gas species in the mixture, only H2O of which is adsorbing. The "base line" or "inlet gas" without H2O has a composition of N2 at 0.7825, O2 at 0.2081, Ar at 0.009, and CO2 at 0.0004.

Adsorbent Input File

```
Domain Coord. (2 = spherical, 1 = cylindrical) { [tab] Char. Length (um) (i.e., cylinder length) } (NOTE: Char. Length is only needed if problem is not spherical)
Pellet Radius (um)
```

Example Adsorbent Input

1 6.0

2.0

Above example is for a cylindrical adsorbent with a length of 5 um and radius of 2 um.

Component Input File

Molar Weight of ith species (g/mol) [tab] Specific Heat of ith species (J/g/K) Sutherland Viscosity (g/cm/s) [tab] Sutherland Temperature (K) [tab] Sutherland Constant (K) of ith species (repeat above for all species in same order they appeared in the Scenario Input File)

Example Component Input

28.016 1.04 0.0001781 300.55 111.0 32.0 0.919 0.0002018 292.25 127.0 39.948 0.522 0.0002125 273.11 144.4 44.009 0.846 0.000148 293.15 240.0 18.0 1.97 0.0001043 298.16 784.72

Above example is exactly the same as in the SCOPSOWL_SCENARIO example (see scopsowl.h). There is no difference in the input file formats for this input. Keep in mind that the order is VERY important! All species information must be in the same order that the species appeared in the Scenario input file.

Adsorbate Input File

Reference Diffusivity (um^2/hr) [tab] Activation Energy (J/mol) of ith adsorbable species Reference Temperature (K) [tab] Affinity Constant (-) of ith adsorbable species van der Waals Volume (cm^3/mol) of ith species GSTA adsorption capacity (mol/kg) of ith species Number of GSTA parameters of ith species Enthalpy (J/mol) of nth site [tab] Entropy of nth site (J/K/mol) of ith species (repeat enthalpy and entropy for all n sites in species i) (repeat above for all species i)

Example Adsorbate Input

0 0 0 0 13.91 11.67 4 -46597.5 -53.6994 -125024 -221.073 -193619 -356.728 -272228 -567.459

Above example gives the equilibrium parameters associated with the H2O-MS3A single component adsorption system. Note that the kinetic parameters (Ref. Diff., Act. Energy, Ref. Temp., and Affinity) were all given a value of zero. These values are irrelavent if we are running an optimization because they will be replaced with a single estimate for the diffusivity that is being optimization for. However, if we wanted to run this routine with comparisons and not do any optimization, then you would need to provide non-zero values for these parameters (at least for Ref. Diff.).

Data Input File

Number of Kinetic Data Curves

Number of data points in the ith curve

Temperature (K) [tab] Partial Pressure (kPa) [tab] Equilibrium Adsorption (mol/kg) all of ith curve

Time point 1 (hrs) [tab] Adsorption 1 (mol/kg) of ith curve

Time point 1 (hrs) [tab] Adsorption 2 (mol/kg) of ith curve

... (Repeat for all time-adsorption data points)

(Repeat above for all curves i)

Example Data Input

```
40

2990

298.15 0.000310922 2.9

0 0

0.1666666667 0.001834419

0.333611111 0.004880247

0.5 0.008306803

...

2789

298.15 0.00055189 5

0 0

0.166944444 0.003350185

0.333611111 0.007418267

0.5 0.009930906

0.6666666667 0.014597236

0.833611111 0.021377373
```

Above is a partial example for a data set of 40 kinetic curves. The first curve contains 2990 data points and has temperature of 298.15 K, partial pressure of 0.000310922 kPa, and an equilibrium adsorption of 2.9. Each first time point should start from 0 hours and each initial adsorption should correspond to the value of initial adsorption indicated in the Scenario input file. Then, this structure is repeated for all adsorptio curves.

6.20 Trajectory.h File Reference

Single Particle Trajectory Analysis for Magnetic Filtration.

```
#include "macaw.h"
#include <random>
#include <chrono>
```

Classes

struct TRAJECTORY_DATA

Functions

- double Magnetic_R (const Matrix< double > &dX, const Matrix< double > &dY, int i, double b, double mu_0, double chi_p, double M, double H0, double a)
- double Magnetic_T (const Matrix< double > &dX, const Matrix< double > &dY, int i, double b, double mu_0, double chi_p, double M, double H0, double a)
- double Grav_R (const Matrix < double > &dX, int i, double b, double rho_p, double rho_f)
- double Grav_T (const Matrix< double > &dX, int i, double b, double rho_p, double rho_f)
- double Van_R (const Matrix< double > &dX, const Matrix< double > &dY, int i, double Hamaker, double b, double a)
- double V_RAD (const Matrix< double > &dX, const Matrix< double > &dY, int i, double V0, double rho_f, double a, double eta)
- double V_THETA (const Matrix< double > &dX, const Matrix< double > &dY, int i, double V0, double rho_f, double a, double eta)
- double Brown_RAD (double n_rand, double m_rand, double sigma_n, double sigma_m)

- double Brown_THETA (double s_rand, double t_rand, double sigma_n, double sigma_m)
- int POLAR (Matrix < double > &POL, const Matrix < double > &dX, const Matrix < double > &dY, const void *data, int i)
- double In_PVel_Rad (const Matrix< double > &POL)
- double In PVel Theta (const Matrix< double > &POL)
- int In_P_Velocity (const Matrix< double > &POL, Matrix< double > &Vr, Matrix< double > &Vt)
- double PVel_Rad (const Matrix< double > &POL, const Matrix< double > &Vr, int i, double mp, double beta, double t, double sigma_v, double rand_n)
- double PVel_Theta (const Matrix< double > &POL, const Matrix< double > &Vt, int i, double mp, double beta, double t, double sigma v, double rand s)
- int P_Velocity (const Matrix< double > &POL, Matrix< double > &Vr, Matrix< double > &Vt, int i, const void *data)
- double RADIAL_FORCE (const Matrix< double > &POL, const Matrix< double > &Vr, int i, double beta, double mp, double dt, double a)
- double TANGENTIAL_FORCE (const Matrix< double > &POL, const Matrix< double > &Vt, const Matrix
 double > &dY, int i, double beta, double mp, double dt, double a)
- double Capture_Force (const Matrix< double > &POL, const Matrix< double > &Vr, int i, double beta, double mp, double dt, double a)
- int CARTESIAN (const Matrix< double > &POL, const Matrix< double > &Vr, const Matrix< double > &Vt, Matrix< double > &H, const Matrix< double > &dY, int i, const void *data)
- int DISPLACEMENT (Matrix< double > &dX, Matrix< double > &dY, const Matrix< double > &H, int i)
- int LOCATION (const Matrix< double > &dY, const Matrix< double > &dX, Matrix< double > &X, Matrix< double > &Y, int i)
- double Removal_Efficiency (double Sum_Cap, const void *data)
- int Trajectory_SetupConstants (TRAJECTORY_DATA *dat)
- int Number_Generator (TRAJECTORY_DATA *dat)
- int Run_Trajectory ()

Run_Trajectory function.

6.20.1 Detailed Description

Single Particle Trajectory Analysis for Magnetic Filtration.

Trajectory.cpp

Alex, Please provide details here... and elsewhere in the file.

Author

Alex Wiechert

Date

08/25/2015

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This software was designed and built at the Georgia Institute of Technology by Alex Wiechert for PhD research in the area of environmental surface science. Copyright (c) 2015, all rights reserved.

- 6.20.2 Function Documentation
- 6.20.2.1 double Magnetic_R (const Matrix < double > & dX, const Matrix < double > & dY, int i, double b, double mu_0 , double chi_p , double d, double d, double d0, double d0,
- 6.20.2.2 double Magnetic_T (const Matrix < double > & dX, const Matrix < double > & dY, int i, double b, double mu_0, double chi p, double M, double H0, double a)
- 6.20.2.3 double Grav_R (const Matrix < double > & dX, int i, double b, double rho_p, double rho_f)
- 6.20.2.4 double Grav_T (const Matrix < double > & dX, int i, double b, double rho_p, double rho_f)
- 6.20.2.5 double Van_R (const Matrix < double > & dX, const Matrix < double > & dY, int i, double dX, int dX double dX doub
- 6.20.2.6 double V_RAD (const Matrix< double > & dX, const Matrix< double > & dY, int i, double V0, double rho_f, double a, double eta)
- 6.20.2.7 double V_THETA (const Matrix< double > & dX, const Matrix< double > & dY, int i, double VO, dou
- 6.20.2.8 double Brown_RAD (double n_rand, double m_rand, double sigma_n, double sigma_m)
- 6.20.2.9 double Brown_THETA (double s_rand, double t_rand, double sigma_n, double sigma_m)
- 6.20.2.10 int POLAR (Matrix< double > & POL, const Matrix< double > & dX, const Matrix< double > & dY, const void * data, int i)
- 6.20.2.11 double In_PVel_Rad (const Matrix< double > & POL)
- 6.20.2.12 double In_PVel_Theta (const Matrix < double > & POL)
- 6.20.2.13 int In_P_Velocity (const Matrix < double > & POL, Matrix < double > & Vr, Matrix < double > & Vt)
- 6.20.2.14 double PVel_Rad (const Matrix < double > & POL, const Matrix < double > & Vr, int i, double mp, double beta, double t, double $sigma_v$, double $rand_n$)
- 6.20.2.15 double PVel_Theta (const Matrix < double > & POL, const Matrix < double > & Vt, int i, double mp, double beta, double t, double sigma_v, double rand_s)
- 6.20.2.16 int P_Velocity (const Matrix < double > & POL, Matrix < double > & Vr, Matrix < double > & Vt, int i, const void * data)
- 6.20.2.17 double RADIAL_FORCE (const Matrix< double > & POL, const Matrix< double > & Vr, int i, double beta, double mp, double dt, double a)
- 6.20.2.18 double TANGENTIAL_FORCE (const Matrix< double > & POL, const Matrix< double > & Vt, const Matrix< double > & dY, int i, double beta, double mp, double dt, double a)
- 6.20.2.19 double Capture_Force (const Matrix < double > & POL, const Matrix < double > & Vr, int i, double beta, double mp, double dt, double a)

6.21 ui.h File Reference 383

```
6.20.2.20 int CARTESIAN ( const Matrix < double > & POL, const Matrix < double > & Vr, const Matrix < double > & Vt, Matrix < double > & Vt,
```

6.21 ui.h File Reference

User Interface for Ecosystem.

```
#include <fstream>
#include <string>
#include <iostream>
#include "error.h"
#include "yaml_wrapper.h"
#include "flock.h"
#include "school.h"
#include "sandbox.h"
#include "Trajectory.h"
```

Classes

struct UI_DATA

Data structure holding the UI arguments.

Macros

- #define UI_HPP_
- #define ECO_VERSION "1.0.0"

Macro expansion for executable current version number.

#define ECO_EXECUTABLE "eco"

Macro expansion for executable current name.

Enumerations

```
    enum valid_options {
        TEST, EXECUTE, EXIT, CONTINUE,
        HELP, dogfish, eel, egret,
        finch, lark, macaw, mola,
        monkfish, sandbox, scopsowl, shark,
        skua, gsta_opt, magpie, scops_opt,
        skua_opt, trajectory }
```

Valid options available upon execution of the code.

Functions

void aui_help ()

Function to display help for Advanced User Interface.

· void bui_help ()

Function to display help for Basic User Interface.

bool exit (const std::string &input)

Function returns true if user requests exit.

bool help (const std::string &input)

Function returns trun if the user requests help.

bool version (const std::string &input)

Function returns true if user requests to know the executable version.

bool test (const std::string &input)

Function returns true if user requests to run a test.

bool exec (const std::string &input)

Function returns true if the user requests to run a simulation/executable.

bool path (const std::string &input)

Function returns true if the user indicates that input files share a common path.

bool input (const std::string &input)

Function returns true if the user indicates that the next arguments are input files.

• bool valid_test_string (const std::string &input, UI_DATA *ui_dat)

Function returns true if the user gave a valid test option.

bool valid_exec_string (const std::string &input, UI_DATA *ui_dat)

Function returns true if the user gave a valid execution option.

int number_files (UI_DATA *ui_dat)

Function returns the number of expected input files for the user's run option.

bool valid_addon_options (UI_DATA *ui_dat)

Function returns true if the user has choosen a valid additional runtime option.

void display_help (UI_DATA *ui_dat)

Function to call the appropriate help menu based on type of interface.

void display_version (UI_DATA *ui_dat)

Function to display ecosystem version information to the console.

int invalid_input (int count, int max)

Function returns a CONTINUE or EXIT when invalid input is given.

• bool valid input main (UI DATA *ui dat)

Function returns true if user gave valid input in Basic UI.

bool valid_input_tests (UI_DATA *ui_dat)

Function returns true if user gave a valid test function to run.

bool valid_input_execute (UI_DATA *ui_dat)

Function returns true if user gave a valid executable function to run.

6.21 ui.h File Reference 385

int test_loop (UI_DATA *ui_dat)

Function that loops the Basic UI until a valid test option was selected.

int exec_loop (UI_DATA *ui_dat)

Function that loops the Basic UI until a valid executable option was selected.

int run_test (UI_DATA *ui_dat)

Function will call the user requested test function.

int run_exec (UI_DATA *ui_dat)

Function will call the user requested executable function.

• int run_executable (int argc, const char *argv[])

Function called by the main and runs both user interfaces for the program.

6.21.1 Detailed Description

User Interface for Ecosystem.

ui.cpp

These routines define how the user will interface with the software

Author

Austin Ladshaw

Date

08/25/2015

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This software was designed and built at the Georgia Institute of Technology by Austin Ladshaw for PhD research in the area of adsorption and surface science. Copyright (c) 2015, all rights reserved.

6.21.2 Macro Definition Documentation

6.21.2.1 #define UI_HPP_

6.21.2.2 #define ECO_VERSION "1.0.0"

Macro expansion for executable current version number.

6.21.2.3 #define ECO_EXECUTABLE "eco"

Macro expansion for executable current name.

6.21.3 Enumeration Type Documentation

6.21.3.1 enum valid_options

Valid options available upon execution of the code.

Enumeration of valid options for executing the ecosystem code. More options become available as the code updates. Some options that appear here may not be viewable in the "help" screen of the executable. Those options are hidden, but are still valid entries.

Enumerator

TEST EXECUTE EXIT CONTINUE HELP dogfish eel egret finch lark macaw mola monkfish sandbox scopsowl shark skua gsta_opt magpie

6.21.4 Function Documentation

```
6.21.4.1 void aui_help()
```

scops_opt skua_opt trajectory

Function to display help for Advanced User Interface.

The Advanved User Interface help screen is accessed by including run option -h or -help when executing the program from command line.

```
6.21.4.2 void bui_help()
```

Function to display help for Basic User Interface.

The Basic User Interface help screen is accessed by running the executable, then typing "help" at any point during the console prompts. Exception to this occurs when the console prompts you to provide input files for your choosen routine. In this circumstance, the executable always assumes that what the user types in will be an input file.

```
6.21.4.3 bool exit ( const std::string & input )
```

Function returns true if user requests exit.

This function will check the input string for "exit" or "quit" and terminate the executable. Only checked if using the Basic User Interface.

6.21 ui.h File Reference 387

Parameters

input input string user gives to the console
--

6.21.4.4 bool help (const std::string & input)

Function returns trun if the user requests help.

This function will check the input string for "help", "-h", or "--help" and will tell the executable to display the help menu. The help menu that gets displayed depends on how the executable was run to begin with.

Parameters

input	input string user gives to the console
-------	--

6.21.4.5 bool version (const std::string & input)

Function returns true if user requests to know the executable version.

This function will check the input string for "version", "-v", or "--version" and will tell the executable to display version information about the executable.

Parameters

input	input string user gives to the console
-------	--

6.21.4.6 bool test (const std::string & input)

Function returns true if user requests to run a test.

This function will check the input string for "-t" or "--test" and determine whether or not the user requests to run an ecosystem test function.

Parameters

input	input string user gives to the console

6.21.4.7 bool exec (const std::string & input)

Function returns true if the user requests to run a simulation/executable.

This function will check the input string for "-e" or "--execute" and determine whether or not the user requests to run an ecosystem executable function.

Parameters

input	input string the user gives to the console
-------	--

388 CONTENTS

6.21.4.8 bool path (const std::string & input)

Function returns true if the user indicates that input files share a common path.

This function will check the input string for "-p" or "--path" and determine whether or not the user will give a common path to all input files needed for the specified simulation. Only used in Advanced User Interface.

Parameters

input	input string the user gives to the console
-------	--

6.21.4.9 bool input (const std::string & input)

Function returns true if the user indicates that the next arguments are input files.

This function will check the input string for "-i" or "--input" and determine whether or not the user's next arguments are input files for a specific simulation. Only used in Advanced User Interface.

Parameters

<i>input</i> input string the user gives to the cons
--

6.21.4.10 bool valid_test_string (const std::string & input, UI_DATA * ui_dat)

Function returns true if the user gave a valid test option.

This function will check the input string given by the user and determine whether that string denotes a valid test. Then, it will mark the option variable in ui_dat with the appropriate option from the valid_options enum.

Parameters

iı	nput	input string the user gives to the console
u	ıi_dat	pointer to the data structure for the ui object

6.21.4.11 bool valid_exec_string (const std::string & input, UI_DATA * ui_dat)

Function returns true if the user gave a valid execution option.

This function will check the input string given by the user and determine whether that string denotes a valid execution option. Then, it will mark the option variable in ui_dat with the appropriate option from the valid_options enum.

Parameters

input	input string the user gives to the console
ui_dat	pointer to the data structure for the ui object

6.21.4.12 int number_files (UI_DATA * ui_dat)

Function returns the number of expected input files for the user's run option.

6.21 ui.h File Reference 389

This function will check the option variable in the ui_dat structure to determine the number of input files that is expected to be given. Running different executable functions in ecosystem may require various number of input files.

Parameters

	ui_dat	pointer to the data structure for the ui object
--	--------	---

6.21.4.13 bool valid_addon_options (UI_DATA * ui_dat)

Function returns true if the user has choosen a valid additional runtime option.

This function will check all additional input options in the user_input variable of ui_dat to determine if the user requests any additional options during runtime. Valid additional options are -p or -path and -i or -input.

Parameters

or the ui object

6.21.4.14 void display_help (UI_DATA * ui_dat)

Function to call the appropriate help menu based on type of interface.

This function looks at the ui_dat structure and the user's OS files to determine what help menu to display and how to display it. There are two different types of help menus that can be displayed: (i) Advanced Help and (ii) Basic Help. Additionally, this function checks the OS file system for the existence of installed help files. If it finds those files, then it instructs the command terminal to read the contents of those files with the "less" command. Otherwise, it will just print the appropriate help menu to the console window.

Parameters

ui_d	t	pointer to the data structure for the ui object	
------	---	---	--

6.21.4.15 void display_version (UI_DATA * ui_dat)

Function to display ecosystem version information to the console.

This function will check the ui_dat structure to see which type of interface the user is using, then print out the version information for the executable being run.

Parameters

ui dat	pointer to the data structure for the ui object

6.21.4.16 int invalid_input (int count, int max)

Function returns a CONTINUE or EXIT when invalid input is given.

This function looks at the current count and the max iterations and determines whether or not to force the executable to terminate. If the user provides too many incorrect options during the Basic User Interface, then the executable will force quit.

390 CONTENTS

Parameters

count	number of times the user has provided a bad option
max	maximum allowable bad options before force quit

6.21.4.17 bool valid_input_main (UI_DATA * ui_dat)

Function returns true if user gave valid input in Basic UI.

This function is only called if the user is running the Basic UI. It checks the given console argument stored in user_input of ui_dat for a valid option. If no valid option is given, then this function returns false.

Parameters

ı	ui_dat	pointer to the data structure for the ui object
---	--------	---

6.21.4.18 bool valid_input_tests (UI_DATA * ui_dat)

Function returns true if user gave a valid test function to run.

This function checks the user_input argument of ui_dat for a valid test option. If no valid test was given, then this function returns false.

Parameters

ui_dat	pointer to the data structure for the ui object
--------	---

6.21.4.19 bool valid_input_execute (UI_DATA * ui_dat)

Function returns true if user gave a valid executable function to run.

This function checks the user_input argument of ui_dat for a valid executable option. If no valid executable was given, then this function returns false.

Parameters

ui_dat	pointer to the data structure for the ui object
--------	---

6.21.4.20 int test_loop (UI_DATA * ui_dat)

Function that loops the Basic UI until a valid test option was selected.

This function loops the Basic UI menu for running a test until a valid test is selected by the user. If a valid test is not selected, and the maximum number of loops has been reached, then this function will cause the program to force quit.

Parameters

ui_dat	pointer to the data structure for the ui object
ui_aai	pointer to the data structure for the ui object

```
6.21.4.21 int exec_loop ( UI_DATA * ui_dat )
```

Function that loops the Basic UI until a valid executable option was selected.

This function loops the Basic UI menu for running an executable until a valid executable is selected by the user. If a valid executable is not selected, and the maximum number of loops has been reached, then this function will cause the program to force quit.

Parameters

ui_dat	pointer to the data structure for the ui object
--------	---

```
6.21.4.22 int run_test ( UI_DATA * ui_dat )
```

Function will call the user requested test function.

This function checks the option variable of the ui_dat structure and runs the corresponding test function.

Parameters

|--|

Function will call the user requested executable function.

This function checks the option variable of the ui_dat structure and runs the corresponding executable function.

Parameters

ui_dat pointer to	the data structure for the ui object
---------------------	--------------------------------------

6.21.4.24 int run_executable (int argc, const char * argv[])

Function called by the main and runs both user interfaces for the program.

This function is called in the main.cpp file and passes the console arguments given at run time.

Parameters

arg	0	number of arguments provided by the user at the time of execution	
arg	V	list of C-strings that was provided by the user at the time of execution	

6.22 yaml_wrapper.h File Reference

C++ Wrapper for the C-YAML Library.

392 CONTENTS

```
#include "yaml.h"
#include "error.h"
#include <map>
#include <string>
#include <iostream>
#include <utility>
#include <stdexcept>
```

Classes

class ValueTypePair

Value-Type Pair object to recognize data type of a string that was read.

class KeyValueMap

Key-Value-Type Map object creating a map of the KeyValuePair objects.

· class SubHeader

Object for the Lowest level of Header for the yaml_wrapper.

· class Header

Object for headers in a yaml document (inherits from SubHeader)

class Document

Object for the various documents in the yaml file.

class YamlWrapper

Object for the entire yaml file holding all documents, header, sub-headers, keys, and values.

· class yaml cpp class

Primary object used when reading and digitally storing yaml files.

Typedefs

• typedef enum data_type data_type

Enum for valid data types in ValueTypePair.

• typedef enum header_state header_state

Enum for state of the headers in the yaml_wrapper.

Enumerations

```
    enum data_type {
        STRING, BOOLEAN, DOUBLE, INT,
        UNKNOWN }
```

enum header_state { ANCHOR, ALIAS, NONE }

Functions

std::string allLower (const std::string &input)

Function to return an all lower case string based on the passed argument.

• bool isEven (int n)

Function to return true if the given argument is an even number.

int YAML_WRAPPER_TESTS ()

Function to run tests on all the objects that yaml_cpp_class.

• int YAML_CPP_TEST (const char *file)

Function to run a test read for the yaml_cpp_class on a given file.

6.22.1 Detailed Description

C++ Wrapper for the C-YAML Library.

yaml_wrapper.cpp

This file holds objects, structures, and functions associatied with using the C-YAML library. A C++ wrapper has been created for the Kirill Simonov (2006) LibYAML library to more easily store and query information in yaml style input files. The wrapper uses the C-YAML parser to identify the file structure and store the read in information from that document into an object using C++ maps. Those maps are hold information in a series of Key-Value pairs as well as lists of Key-Value pairs. This allows the user to create well organized input files to change the behavior of simulations.

The yaml_wrapper is restricted to the same limitations in the C-YAML source code in terms of how the documents are allowed to be structured for TOKEN based parsing. C-YAML only recognizes specific tokens and will only allow a certain level of Sub-Header mapping. Therefore, this wrapper has the same limitations. Below is an example of acceptable formatting for a C-YAML document.

#Test input file for YAML and SHARK

TestDoc1: &hat

· scenario: numvar: 25 act_fun: DAVIES steadystate: FALSE

t out: 1 pH: 0

· testblock: another: block

> – subblock: sub: block

> > TestDoc2: *hat

· masterspecies:

"CI - (aq)": 0 "Na + (aq)": 1 "H2O (I)": 2 3: NaCl (aq)

TestDoc3:

apple: red pear: green

· array: #Block banana: yellow #List 1 in array

394 CONTENTS

```
- list1: &a #also a block
           a: 1 #key : value
           b: 2
           c: 3
           #List 2 in array
         - list2: *a
           a: 4
           b: 5
           c: 6
           TestDoc4:
    · anchor: &anchor
      stuff: to do
    • alias: *anchor
      add: to stuff
    • list:
         - anchored: &list_anchor
           info: blah
           atta: boy
         - aliased: *list_anchor
           info: bruh
           atta: ber
           #WARNING: MAKE SURE FILE DOES NOT CONTAIN TABS!!!
TestDoc5:
    • grab: *anchor
      add2: more adds
         - listcopy: *a
    • block: {1: 2, 3: 4}
      still: in block
      Note
           You can view the actual yaml example file in the input_files/SHARK/test_input.yml sub-directory of the
           project folder.
      Author
           Austin Ladshaw
```

Date

07/29/2015

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6.22.2 Typedef Documentation

6.22.2.1 typedef enum data_type data_type

Enum for valid data types in ValueTypePair.

6.22.2.2 typedef enum header_state header_state

Enum for state of the headers in the yaml_wrapper.

6.22.3 Enumeration Type Documentation

6.22.3.1 enum data_type

Enumerator

STRING

BOOLEAN

DOUBLE

INT

UNKNOWN

396 CONTENTS

6.22.3.2 enum header_state

Enumerator

ANCHOR

ALIAS

NONE

6.22.4 Function Documentation

6.22.4.1 std::string allLower (const std::string & input)

Function to return an all lower case string based on the passed argument.

This function will copy the input paramter and convert that copy to all lower case. The copy is then returned and can be checked against valid or allowed strings.

Parameters

input string to copy and convert to lower case

6.22.4.2 bool isEven (int n)

Function to return true if the given argument is an even number.

6.22.4.3 int YAML_WRAPPER_TESTS ()

Function to run tests on all the objects that yaml_cpp_class.

This test function is currently NOT callable from the UI.

6.22.4.4 int YAML_CPP_TEST (const char * file)

Function to run a test read for the yaml_cpp_class on a given file.

This test/executable function is currently NOT callable from the UI.

Index

\sim AdsorptionReaction	AQUEOUS
AdsorptionReaction, 12	mola.h, 321
\sim Atom	ARNOLDI_DATA, 21
Atom, 26	beta, 22
\sim ChemisorptionReaction	e1, 23
ChemisorptionReaction, 41	Hkp1, 23
\sim Document	hp1, 22
Document, 50	iter, 22
\sim Header	k, 22
Header, 90	Output, 23
~KeyValueMap	sum, 23
KeyValueMap, 95	v, 23
~MassBalance	Vk, 23
MassBalance, 104	w, 23
~MasterSpeciesList	yk, 23
MasterSpeciesList, 109	abs_tol_bias
~Matrix	
	SCOPSOWL_OPT_DATA, 188
Matrix, 115	SKUA_OPT_DATA, 211
~Molecule	AbsPerm
Molecule, 126	shark.h, 349
~MultiligandAdsorption	act_choice
MultiligandAdsorption, 142	shark.h, 355
\sim MultiligandChemisorption	act_fun
MultiligandChemisorption, 152	AdsorptionReaction, 20
\sim PeriodicTable	MultiligandAdsorption, 148
PeriodicTable, 163	MultiligandChemisorption, 157
\sim Reaction	SHARK_DATA, 198
Reaction, 176	activation_energy
\sim SubHeader	SCOPSOWL_PARAM_DATA, 191
SubHeader, 214	SKUA PARAM, 212
~UnsteadyAdsorption	UnsteadyReaction, 246
UnsteadyAdsorption, 228	activities
~UnsteadyReaction	AdsorptionReaction, 20
UnsteadyReaction, 239	MultiligandAdsorption, 148
~ValueTypePair	MultiligandChemisorption, 157
ValueTypePair, 248	activities_old
~YamlWrapper	
	UnsteadyAdsorption, 235
YamlWrapper, 254	activity_data
~yaml_cpp_class	AdsorptionReaction, 20
yaml_cpp_class, 251	MultiligandAdsorption, 148
	MultiligandChemisorption, 157
A	SHARK_DATA, 203
magpie.h, 306	activity_new
a	SHARK_DATA, 202
TRAJECTORY_DATA, 221	activity_old
A_separator	SHARK_DATA, 202
TRAJECTORY_DATA, 221	addDocKey
A_wire	YamlWrapper, 255
TRAJECTORY_DATA, 221	addHeadKey
ADSORBED	Document, 52
mola.h, 321	addKey
ALIAS	KeyValueMap, 96
yaml_wrapper.h, 396	addPair
ANCHOR	Document, 51, 52
yaml_wrapper.h, 396	Header, 92
Jam_mappoint, 000	rioddoi, J L

KeyValueMap, 97	getlonicStrength, 18
SubHeader, 215	getMolarFactor, 17
addSubKey	getNumberRxns, 18
Header, 92	getReaction, 17
adjoint	getSpecificArea, 18
Matrix, 117	getSpecificMolality, 18
ads_rxn	getSurfaceCharge, 18
AdsorptionReaction, 21	getTotalMass, 18
ChemisorptionReaction, 47	getTotalVolume, 18
UnsteadyAdsorption, 235	getVolumeFactor, 18
adsorb_index	IncludeSurfCharge, 21
AdsorptionReaction, 20	includeSurfaceCharge, 19
SCOPSOWL_OPT_DATA, 186	Initialize_Object, 12
SKUA_OPT_DATA, 209	ionic_strength, 21
Adsorbable	isAreaBasis, 19
SCOPSOWL_PARAM_DATA, 192	List, 19
SKUA_PARAM, 212	modifyDeltas, 12
adsorbent_name	molar_factor, 20
AdsorptionReaction, 21	num_rxns, 21
MultiligandAdsorption, 147	setActivities, 14
MultiligandChemisorption, 157	setActivityEnum, 13
AdsorptionList	setActivityModelInfo, 13
SHARK_DATA, 196	setAdsorbIndices, 13
AdsorptionReaction, 8	setAdsorbentName, 14
\sim AdsorptionReaction, 12	setAqueousIndex, 13
act_fun, 20	setAqueousIndexAuto, 13
activities, 20	setAreaBasisBool, 14
activity_data, 20	setAreaFactor, 13
ads_rxn, 21	setBasis, 14
adsorb_index, 20	setChargeDensity, 15
adsorbent_name, 21	setChargeDensityValue, 14
AdsorptionReaction, 12	setIonicStrength, 15
aqueous_index, 20	setIonicStrengthValue, 14
area_factors, 20	setMolarFactor, 13
AreaBasis, 21	setSpecificArea, 14
calculateActiveFraction, 15	setSpecificMolality, 14
calculateAqueousChargeExchange, 16	setSurfaceCharge, 14
calculateAreaFactors, 15	setSurfaceChargeBool, 14
calculateEquilibria, 15	setTotalMass, 14
calculateEquilibriumCorrection, 17	setTotalVolume, 14
calculateLangmuirAdsorption, 16	setVolumeFactor, 13
calculateLangmuirEquParam, 16	specific_area, 20
calculateLangmuirMaxCapacity, 15	specific_molality, 20
calculatePsi, 16	surface_activity, 19
calculateSurfaceChargeDensity, 15	surface_charge, 20
callSurfaceActivity, 15	total_mass, 20
charge_density, 21	total_volume, 20
checkAqueousIndices, 13	volume_factors, 20
Display_Info, 12	affinity
Eval_Residual, 17	SCOPSOWL_PARAM_DATA, 191
getActivity, 18	SKUA_PARAM, 212
getActivityEnum, 19	Ai
getAdsorbIndex, 19	OPTRANS_DATA, 160
getAdsorbentName, 19	alias
getAqueousIndex, 19	SubHeader, 216
getAreaFactor, 18	alkalinity
getBulkDensity, 18	MasterSpeciesList, 111
getChargeDensity, 18	all_pars

GSTA_OPT_DATA, 87	editProtons, 26
allLower	editRadii, 27
yaml_wrapper.h, 396	editValence, 26
alpha	Electrons, 27
BACKTRACK_DATA, 30	electrons, 28
BiCGSTAB_DATA, 32	Name, 29
CGS DATA, 36	NaturalState, 29
GCR DATA, 73	Neutrons, 27
PCG DATA, 161	neutrons, 28
anchor_alias_dne	oxidation_state, 28
error.h, 268	OxidationState, 27
	Protons, 27
Ap	protons, 28
PCG_DATA, 162	Register, 26
aqueous_index	
AdsorptionReaction, 20	removeElectron, 27
area_factors	removeNeutron, 27
AdsorptionReaction, 20	removeProton, 27
AreaBasis	Symbol, 29
AdsorptionReaction, 21	valence_e, 28
AreaSTD	AtomCategory
shark.h, 348	Atom, 28
arg	AtomName
GMRESR DATA, 79	Atom, 27
<u> </u>	AtomState
arg_matrix_same	Atom, 28
error.h, 268	AtomSymbol
argc	-
UI_DATA, 224	Atom, 28
argv	atomic_number
UI_DATA, 224	Atom, 29
arnoldi	atomic_radii
lark.h, 289	Atom, 28
arnoldi dat	atomic_weight
GMRESLP DATA, 76	Atom, 28
- · · ·	AtomicNumber
As CVOTEM DATA 010	Atom, 28
SYSTEM_DATA, 218	AtomicRadii
assertType	Atom, 27
KeyValueMap, 97	AtomicWeight
ValueTypePair, 248	Atom, 27
Atom, 23	atoms
\sim Atom, 26	Molecule, 131
Atom, 26	
AtomCategory, 28	aui_help
AtomName, 27	ui.h, 386
AtomState, 28	avg_fiber_density
AtomSymbol, 28	MONKFISH_DATA, 135
atomic_number, 29	avg_norm
atomic_radii, 28	SYSTEM_DATA, 218
	avg_sorption
atomic_weight, 28	MONKFISH_PARAM, 138
AtomicNumber, 28	avg_sorption_old
AtomicRadii, 27	MONKFISH_PARAM, 138
AtomicWeight, 27	avgDp
BondingElectrons, 27	scopsowl.h, 329
Category, 29	avgPar
DisplayInfo, 28	gsta_opt.h, 280
editAtomicWeight, 26	avgValue
editElectrons, 26	
editNeutrons, 26	gsta_opt.h, 280
editOxidationState, 26	b
Sale Additional Control Control	~

TRAJECTORY_DATA, 221	beta, 32
B0	breakdown, 32
TRAJECTORY_DATA, 221	iter, 32
BACKTRACK_DATA, 29	maxit, 32
alpha, 30	omega, <mark>32</mark>
constRho, 30	omega_old, 33
Fk, 30	Output, 33
	p, 34
fun_call, 30	r, 33
lambdaMin, 30	
normFkp1, 30	r0, 33
rho, 30	relres, 33
xk, 30	relres_base, 33
BATCH	res, 33
shark.h, 349	rho, 32
BOOLEAN	rho_old, 32
yaml_wrapper.h, 395	s, 34
backtrack_dat	t, 34
PJFNK_DATA, 171	tol_abs, 33
backtrackLineSearch	tol_rel, 33
lark.h, 299	v, 34
BasicUI	x, 33
UI DATA, 224	y, 34
begin	z, <mark>34</mark>
Document, 51	BiCGSTAB
Header, 91	lark.h, 288
	bicgstab
KeyValueMap, 96	lark.h, 293
YamlWrapper, 254, 255	bicgstab_dat
best_par	PJFNK DATA, 170
GSTA_OPT_DATA, 87	binary_diffusion
bestres	MIXED_GAS, 123
BiCGSTAB_DATA, 33	binder_fraction
CGS_DATA, 37	
GCR_DATA, 73	SCOPSOWL_DATA, 183
GMRESLP_DATA, 76	binder_poresize
GMRESRP_DATA, 82	SCOPSOWL_DATA, 183
PCG_DATA, 162	binder_porosity
PICARD DATA, 166	SCOPSOWL_DATA, 183
bestx	BondingElectrons
BICGSTAB DATA, 33	Atom, 27
CGS DATA, 37	Bounce
GCR DATA, 73	PJFNK_DATA, 170
GMRESLP DATA, 76	breakdown
GMRESRP DATA, 82	BiCGSTAB_DATA, 32
PCG DATA, 162	CGS_DATA, 36
= '	GCR_DATA, 72
PICARD_DATA, 166	Brown_RAD
PJFNK_DATA, 170	Trajectory.h, 382
beta	Brown THETA
ARNOLDI_DATA, 22	Trajectory.h, 382
BiCGSTAB_DATA, 32	bui_help
CGS_DATA, 36	ui.h, 386
FINCH_DATA, 65	,
GCR_DATA, 73	С
PCG_DATA, 161	CGS_DATA, 37
TRAJECTORY DATA, 222	GCR DATA, 74
BICGSTAB_DATA, 31	c temp
alpha, 32	GCR DATA, 74
bestres, 33	CARTESIAN
bestx, 33	Trajectory.h, 382
2000, 00	114,00tor y.11, 002

CC_E	ChemisorptionReaction, 43
FINCH_DATA, 66	MultiligandAdsorption, 144
CC_I	MultiligandChemisorption, 154
FINCH DATA, 66	UnsteadyAdsorption, 230
CE3	calculateAvgOxiState
egret.h, 264	Molecule, 128
CGS_DATA, 34	calculateElecticPotential
alpha, 36	ChemisorptionReaction, 44
bestres, 37	MultiligandAdsorption, 145
bestx, 37	MultiligandChemisorption, 154
beta, 36	calculateEnergies
breakdown, 36	Reaction, 177
c, 37	UnsteadyReaction, 242
iter, 36	calculateEquilibria
maxit, 36	AdsorptionReaction, 15
Output, 37	ChemisorptionReaction, 43
p, 37	MultiligandAdsorption, 144
r, 37	MultiligandChemisorption, 154
r0, 37	UnsteadyAdsorption, 230
relres, 36	calculateEquilibrium
relres_base, 37	Reaction, 177
res, 36	UnsteadyReaction, 242
	calculateEquilibriumCorrection
rho, 36	•
sigma, 36	AdsorptionReaction, 17
tol_abs, 36	ChemisorptionReaction, 44
tol_rel, 36	MultiligandAdsorption, 145
u, 37	MultiligandChemisorption, 155
v, 3 7	UnsteadyAdsorption, 232
w, 37	calculateLangmuirAdsorption
x, 37	AdsorptionReaction, 16
z, 38	calculateLangmuirEquParam
CGS	AdsorptionReaction, 16
lark.h, 288	calculateLangmuirMaxCapacity
CL E	AdsorptionReaction, 15
FINCH_DATA, 66	calculateMolarArea
CL I	Molecule, 128
FINCH DATA, 66	calculateMolarVolume
CONTINUE	Molecule, 128
ui.h, 386	calculateMolarWeight
CR E	Molecule, 128
FINCH DATA, 66	calculatePsi
CR I	
-	AdsorptionReaction, 16
FINCH_DATA, 66	UnsteadyAdsorption, 231
CSTR	calculateRate
shark.h, 349	UnsteadyReaction, 242
calculate_ionic_strength	calculateRates
shark.h, 350	UnsteadyAdsorption, 230
calculate_properties	calculateSurfaceChargeDensity
egret.h, 266	AdsorptionReaction, 15
calculateActiveFraction	ChemisorptionReaction, 43
AdsorptionReaction, 15	UnsteadyAdsorption, 231
UnsteadyAdsorption, 231	callSurfaceActivity
calculateAqueousChargeExchange	AdsorptionReaction, 15
AdsorptionReaction, 16	ChemisorptionReaction, 43
ChemisorptionReaction, 44	MultiligandAdsorption, 145
UnsteadyAdsorption, 231	MultiligandChemisorption, 154
calculateAreaFactors	UnsteadyAdsorption, 231
AdsorptionReaction, 15	callroutine
Ausorphornicachon, 10	Cambuline

FINCH_DATA, 69	calculateAqueousChargeExchange, 44
CanCalcHS	calculateAreaFactors, 43
Reaction, 178	calculateElecticPotential, 44
CanCalcG	calculateEquilibria, 43
Reaction, 179	calculateEquilibriumCorrection, 44
Cap	calculateSurfaceChargeDensity, 43
TRAJECTORY_DATA, 222	callSurfaceActivity, 43
Capture_Force	ChemisorptionReaction, 41
Trajectory.h, 382	Delta, 47
Carrier	Display Info, 41
SYSTEM_DATA, 219	Eval RxnResidual, 44
Cartesian	Eval_SiteBalanceResidual, 45
finch.h, 272	getActivity, 45
Category	getActivityEnum, 46
Atom, 29	getAdsorbIndex, 46
cgs	getAdsorbentName, 46
lark.h, 294	getAreaFactor, 45
cgs_dat	getBulkDensity, 46
PJFNK DATA, 170	getChargeDensity, 46
changeKey	getDelta, 45
Document, 51	getlonicStrength, 46
Header, 92	getLigandIndex, 46
YamlWrapper, 255	getNumberRxns, 46
char_length	getReaction, 45
MIXED_GAS, 123	getSpecificArea, 45
	getSpecificMolality, 46
char_macro	
SCOPSOWL_DATA, 182	getTotalMass, 46 getTotalVolume, 46
char_measure SKUA_DATA, 206	getVolumeFactor, 45
	includeSurfaceCharge, 46
char_micro	
SCOPSOWL_DATA, 183	Initialize_Object, 41
Charge Malagula 120	ligand_index, 47
Molecule, 129	modifyMBEdeltas, 41
charge MasterSpecial ist 111	setActivities, 43 setActivityEnum, 42
MasterSpeciesList, 111 Molecule, 130	· · · · · · · · · · · · · · · · · · ·
•	setAdaarbladiaaa 41
charge_density	setAdsorbIndices, 41
AdsorptionReaction, 21	setAdsorbentName, 43
MultiligandAdsorption, 148	setAreaFactor, 42
MultiligandChemisorption, 158	setChargeDensity, 43
check_Mass	setChargeDensityValue, 43
finch.h, 272	setDelta, 42
checkAqueousIndices	setDeltas, 42
AdsorptionReaction, 13	setIonicStrength, 43
MultiligandAdsorption, 143	setlonicStrengthValue, 43
UnsteadyAdsorption, 228	setLigandIndex, 42
CheckMass	setSpecificArea, 42
FINCH_DATA, 65	setSpecificMolality, 42
CheckMolefractions	setSurfaceChargeBool, 43
MIXED_GAS, 122	setTotalMass, 42
checkSpeciesEnergies	setTotalVolume, 43
Reaction, 177	setVolumeFactor, 42
UnsteadyReaction, 242	chi_p
ChemisorptionList	TRAJECTORY_DATA, 221
SHARK_DATA, 196	cleanup
ChemisorptionReaction, 38	yaml_cpp_class, 251
~ChemisorptionReaction, 41	clear
ads_rxn, 47	Document, 51

Header, 91	SCOPSOWL_DATA, 181
KeyValueMap, 96	coord_micro
SubHeader, 215	SCOPSOWL_DATA, 181
YamlWrapper, 255	CoordSTD
CN	shark.h, 348
FINCH_DATA, 65	copyAnchor2Alias
cofactor	Document, 52
Matrix, 116	Header, 92
columnExtend	YamlWrapper, 255
Matrix, 121	count
columnExtract	UI_DATA, 224
Matrix, 120	crystal_radius
columnProjection	SCOPSOWL_DATA, 182 Cstd
Matrix, 119	egret.h, 264
columnReplace	current equil
Matrix, 120	SCOPSOWL_OPT_DATA, 187
columnShrink	SKUA_OPT_DATA, 210
Matrix, 120	current points
columnVectorFill	SCOPSOWL OPT DATA, 186
Matrix, 119	SKUA OPT DATA, 209
columns	current_press
Matrix, 116	SCOPSOWL OPT DATA, 187
CompareFile	SKUA OPT DATA, 210
SCOPSOWL_OPT_DATA, 188	current_temp
SKUA_OPT_DATA, 211	SCOPSOWL OPT DATA, 187
Conc_new	SKUA_OPT_DATA, 210
SHARK_DATA, 202	current token
Conc_old	yaml_cpp_class, 252
SHARK_DATA, 202	Cylindrical
Console_Output	finch.h, 272
SHARK_DATA, 201	,
const_Dc	d
skua.h, 370	FINCH_DATA, 62
const_filmMassTransfer	D_c
scopsowl.h, 332	skua.h, <mark>369</mark>
const_kf	D_ii
skua.h, 371	egret.h, 265
const_pH	D_ij
SHARK_DATA, 201	egret.h, 265
const_pore_diffusion	D_inf
scopsowl.h, 332 constRho	skua.h, <mark>369</mark>
	D_0
BACKTRACK_DATA, 30	skua.h, 369
Constant CFill	DAVIES
Matrix, 118	shark.h, 350
Contains_pOH	DBL_EPSILON
SHARK_DATA, 201	magpie.h, 305
Contains_pH	DEBYE_HUCKEL
SHARK_DATA, 201 Converged	shark.h, 350
	dHo
SHARK_DATA, 201	GSTA_DATA, 85
Convert2Concentration	DISPLACEMENT
shark.h, 357	Trajectory.h, 383
Convert2LogConcentration	DIC EINCH DATA 64
shark.h, 357	FINCH_DATA, 64
coord	DOGFISH_DATA, 53
SKUA_DATA, 205	DirichletBC, 55 end time, 55
coord_macro	GHU_HHE, 55

eval_DI, 56	dogfish.h, 258
eval_kf, 56	default_IntraDiffusion
eval_qs, 56	dogfish.h, 258
eval_R, 56	default_Retardation
fiber_diameter, 56	dogfish.h, 258
fiber_length, 56	default_SurfaceConcentration
fiber_specific_area, 56	dogfish.h, 259
finch_dat, 56	default_adsorption
NonLinear, 55	scopsowl.h, 330
NumComp, 55	default_bcs
OutputFile, 56	finch.h, 275
param_dat, 57	default_density
Print2Console, 55	monkfish.h, 323
Print2File, 55	default_effective_diffusion
t_counter, 55	scopsowl.h, 331
t_print, 55	default_execution
time, 55	finch.h, 274
time_old, 55	default_exterior_concentration
total_sorption, 56	monkfish.h, 324
total_sorption_old, 56	default_film_transfer
total_steps, 55	monkfish.h, 324
user_data, 56	default_filmMassTransfer
DOGFISH_Executioner	scopsowl.h, 332
dogfish.h, 259 DOGFISH PARAM, 57	default_ic
film_transfer_coeff, 57	finch.h, 274
initial_sorption, 58	default_interparticle_diffusion monkfish.h, 323
intraparticle_diffusion, 57	default kf
sorbed_molefraction, 58	skua.h, 370
species, 58	default_monk_adsorption
surface_concentration, 58	monkfish.h, 323
DOGFISH TESTS	default monk equilibrium
dogfish.h, 260	monkfish.h, 323
DOGFISH_postprocesses	default_monkfish_retardation
dogfish.h, 260	monkfish.h, 324
DOGFISH_preprocesses	default_params
dogfish.h, 260	finch.h, 275
DOGFISH reset	default_pore_diffusion
dogfish.h, 260	scopsowl.h, 330
DOGFISH	default_porosity
dogfish.h, 260	monkfish.h, 322
DOUBLE	default postprocess
yaml_wrapper.h, 395	finch.h, 276
dSo	default_precon
GSTA_DATA, 85	finch.h, 275
Data	default_preprocess
Matrix, 121	finch.h, 274
Data_Map	default_res
SubHeader, 216	finch.h, 275
data_type	default_reset
yaml_wrapper.h, 395	finch.h, 276
Davies_equation	default_retardation
shark.h, 354	scopsowl.h, 330
DebyeHuckel_equation	<u>-</u>
	default_solve
shark.h, 354	default_solve finch.h, 274
shark.h, 354 default_Dc	default_solve finch.h, 274 default_surf_diffusion
shark.h, 354 default_Dc skua.h, 369	default_solve finch.h, 274 default_surf_diffusion scopsowl.h, 331
shark.h, 354 default_Dc	default_solve finch.h, 274 default_surf_diffusion

	5 5 .
finch.h, 274	DisplayPair
Delta	ValueTypePair, 249
ChemisorptionReaction, 47	DisplayTable
MassBalance, 107	PeriodicTable, 164
density	Dk
PURE GAS, 173	scopsowl.h, 329
determinate	Dn
Matrix, 116	FINCH DATA, 68
diagonalSolve	Dnp1
Matrix, 119	FINCH DATA, 68
	<u> </u>
dielectric_const	Do SINOU DATA 64
SHARK_DATA, 200	FINCH_DATA, 64
diffusion_type	Doc_Map
SCOPSOWL_OPT_DATA, 186	YamlWrapper, 256
SKUA_OPT_DATA, 209	Document, 47
dim_mis_match	\sim Document, 50
error.h, 268	addHeadKey, 52
Dirichlet	addPair, 51, 52
FINCH DATA, 65	begin, 51
dirichletBCFill	changeKey, 51
Matrix, 119	clear, 51
DirichletBC	copyAnchor2Alias, 52
DOGFISH DATA, 55	DisplayContents, 52
— · · · · · · · · · · · · · · · · · · ·	Document, 50
MONKFISH_DATA, 134	
SCOPSOWL_DATA, 183	end, 51
SKUA_DATA, 206	getAlias, 52
discretize	getAnchoredHeader, 53
FINCH_DATA, 70	getDataMap, 51
Display	getHeadFromSubAlias, 53
Matrix, 117	getHeadMap, 51
Display_Info	getHeader, 51
AdsorptionReaction, 12	getName, 52
ChemisorptionReaction, 41	getState, 52
MassBalance, 104	Head_Map, 53
MultiligandChemisorption, 152	isAlias, 53
Reaction, 176	isAnchor, 53
UnsteadyAdsorption, 228	operator(), 50
UnsteadyReaction, 239	operator=, 50
display_help	operator[], 50
ui.h, 389	resetKeys, 51
	-
display_version	revalidateAllKeys, 51
ui.h, 389	setAlias, 52
DisplayAll	setName, 52
MasterSpeciesList, 110	setNameAliasPair, 52
DisplayConcentrations	setState, 52
MasterSpeciesList, 110	size, 52
DisplayContents	dog_dat
Document, 52	MONKFISH_DATA, 136
Header, 92	dogfish
SubHeader, 216	ui.h, 386
yaml_cpp_class, 251	dogfish.h, 256
YamlWrapper, 255	DOGFISH_Executioner, 259
DisplayInfo	DOGFISH_TESTS, 260
Atom, 28	DOGFISH_postprocesses, 260
MasterSpeciesList, 110	DOGFISH_preprocesses, 260
Molecule, 130	 ·
	DOGFISH 260
DisplayMap	DOGFISH, 260
KeyValueMap, 97	default_FilmMTCoeff, 258

default_IntraDiffusion, 258	SCOPSOWL_OPT_DATA, 187
default Retardation, 258	SKUA_OPT_DATA, 210
default_SurfaceConcentration, 259	ECO EXECUTABLE
print2file DOGFISH header, 258	ui.h, 385
print2file DOGFISH result new, 258	ECO VERSION
print2file_DOGFISH_result_old, 258	ui.h, 385
print2file_species_header, 258	EEL TESTS
set DOGFISH ICs, 259	eel.h, 262
set_DOGFISH_params, 260	EGRET TESTS
set DOGFISH timestep, 260	egret.h, 266
setup_DOGFISH_DATA, 259	eMax
domain_diameter	
MONKFISH_DATA, 135	mSPD_DATA, 139
Dp	magpie.h, 308
scopsowl.h, 329	EXECUTE
Dp_ij	ui.h, 386 FXIT
egret.h, 265	= /
dq_dc	ui.h, 386
SCOPSOWL PARAM DATA, 190	edit
dq_dco	Matrix, 116
SCOPSOWL PARAM DATA, 191	editAllOxidationStates
dq_dp	Molecule, 128
magpie.h, 307	editAtomicWeight
dt	Atom, 26
FINCH DATA, 62	editCharge
MassBalance, 107	Molecule, 128
SHARK DATA, 199	editElectrons
TRAJECTORY_DATA, 222	Atom, 26
dt max	editEnergy
SHARK_DATA, 199	Molecule, 129
dt min	editEnthalpy
SHARK_DATA, 199	Molecule, 128
dt_old	editEntropy
FINCH DATA, 63	Molecule, 128
duplicate variable	editHS
error.h, 268	Molecule, 128
dX	editNeutrons
TRAJECTORY_DATA, 222	Atom, 26
dxj	editOneOxidationState
NUM JAC DATA, 159	Molecule, 128
dY	editOxidationState
TRAJECTORY_DATA, 222	Atom, 26
dynamic_viscosity	editPair
PURE GAS, 173	ValueTypePair, 248
dz	editProtons
FINCH DATA, 63	Atom, 26
TINOTI_DATA, 00	editRadii
е	Atom, 27
shark.h, 348	editValence
e0	Atom, 26
GMRESRP_DATA, 82	editValue
e0 bar	ValueTypePair, 248
GMRESRP_DATA, 82	editValue4Key
e1	KeyValueMap, 96
ARNOLDI_DATA, 23	eduGuess
e_norm	gsta_opt.h, 281
SCOPSOWL_OPT_DATA, 187	eel
SKUA_OPT_DATA, 210	ui.h, <mark>386</mark>
e_norm_old	eel.h, 261
_	

EEL_TESTS, 262	Reaction, 178
egret	eps
ui.h, 386	NUM_JAC_DATA, 159
egret.h, 262	PJFNK_DATA, 169
CE3, 264	Equilibrium
calculate_properties, 266	Reaction, 178
Cstd, 264	error
D_ii, 265	error.h, 269
D_ij, 265	error.h, 266
Dp_ij, 265	anchor_alias_dne, 268
EGRET_TESTS, 266	arg_matrix_same, 268
FilmMTCoeff, 265	dim_mis_match, 268
initialize_data, 265	duplicate_variable, 268
Mu, 265	empty_matrix, 268
Nu, 265	error, 269
PE3, 264	error_type, 267
PSI, 265	file_dne, 267
Po, 264	generic_error, 267
Pstd, 264	indexing_error, 267
RE3, 264	initial_error, 268
ReNum, 265	invalid_atom, 268
Rstd, 264	invalid_boolean, 268
ScNum, 265	invalid_components, 267
set_variables, 265 ek	invalid_console_input, 268 invalid_electron, 268
QR DATA, 174	invalid_fraction, 268
electric_potential	invalid_gas_sum, 268
MultiligandAdsorption, 148	invalid_gas_sum, 268
MultiligandChemisorption, 158	invalid_neutron, 268
Electrons	invalid_norm, 268
Atom, 27	invalid_proton, 268
electrons	invalid_size, 268
Atom, 28	invalid_solid_sum, 268
empirical_kf	invalid_species, 268
skua.h, 371	invalid_species, 268
empty_matrix	invalid_valence, 268
error.h, 268	key_not_found, 268
end	mError, 267
Document, 51	magpie reverse error, 267
Header, 91	matrix_too_small, 268
KeyValueMap, 96	matvec mis match, 268
YamlWrapper, 254	missing information, 268
end_temp	negative_mass, 268
SHARK DATA, 200	negative_time, 268
end time	no_diffusion, 268
DOGFISH DATA, 55	non_real_edge, 268
MONKFISH DATA, 134	non_square_matrix, 268
Energy	not_a_token, 268
Molecule, 130	nullptr_error, 268
energy	nullptr_func, 268
Reaction, 178	opt_no_support, 268
Enthalpy	ortho_check_fail, 268
Molecule, 129	out_of_bounds, 268
enthalpy	read_error, 268
Reaction, 178	rxn_rate_error, 268
Entropy	scenario_fail, 268
Molecule, 129	simulation_fail, 267
entropy	singular_matrix, 268

string_parse_error, 268	magpie.h, 310
tensor_out_of_bounds, 268	eval_kf
unregistered_name, 268	DOGFISH_DATA, 56
unstable_matrix, 268	MONKFISH_DATA, 136
vector_out_of_bounds, 268	SCOPSOWL_DATA, 184
zero_vector, 268	SKUA_DATA, 206
error_type	eval_po
error.h, 267	magpie.h, 310
eta	eval_po_PI
mSPD_DATA, 139	magpie.h, 309
TRAJECTORY_DATA, 221	eval_po_qo
eval_Cex	magpie.h, 309
MONKFISH_DATA, 136	eval_qs
Eval_ChargeResidual	DOGFISH_DATA, 56
MasterSpeciesList, 111	eval_R
eval_Dex	DOGFISH_DATA, 56
MONKFISH_DATA, 135	eval_retard
eval_DI	SCOPSOWL_DATA, 183
DOGFISH_DATA, 56	eval_rho
eval_GPAST	MONKFISH_DATA, 135
magpie.h, 310	eval_surfDiff
eval_GSTA	SCOPSOWL_DATA, 184
gsta_opt.h, 283	EvalActivity
Eval_IC_Residual	SHARK_DATA, 202
MassBalance, 107	evalprecon
UnsteadyAdsorption, 233	FINCH_DATA, 70
UnsteadyReaction, 245	evalres
Eval_ReactionRate	FINCH_DATA, 70
UnsteadyAdsorption, 233	evaluation
UnsteadyReaction, 244	SCOPSOWL_OPT_DATA, 186
Eval Residual	SKUA_OPT_DATA, 209
AdsorptionReaction, 17	exec
MassBalance, 106	ui.h, 387
MultiligandAdsorption, 145	exec_loop
Reaction, 178	ui.h, 391
UnsteadyAdsorption, 232	executeYamlRead
UnsteadyReaction, 244, 245	yaml_cpp_class, 251
eval_Ret	exit
MONKFISH_DATA, 135	ui.h, 386
Eval_RxnResidual	Explicit_Eval
ChemisorptionReaction, 44	UnsteadyAdsorption, 233
MultiligandChemisorption, 155	UnsteadyReaction, 245
eval_SCOPSOWL_Uptake	ExplicitFlux
scopsowl_opt.h, 340	FINCH_DATA, 65
eval_SKUA_Uptake	exterior_concentration
 skua_opt.h, 377	MONKFISH_PARAM, 137
Eval_SiteBalanceResidual	exterior_transfer_coeff
ChemisorptionReaction, 45	MONKFISH_PARAM, 137
MultiligandChemisorption, 155	F
eval_ads	PJFNK DATA, 170
MONKFISH_DATA, 135	f_bias
SCOPSOWL_DATA, 183	SCOPSOWL_OPT_DATA, 187
eval_diff	SKUA_OPT_DATA, 210
SCOPSOWL_DATA, 184	f_bias_old
SKUA_DATA, 206	SCOPSOWL_OPT_DATA, 188
eval_eps	SKUA OPT DATA, 210
MONKFISH_DATA, 135	fC_E
eval_eta	FINCH_DATA, 67
ightharpoonup	<u> </u>

fC_I	pjfnk_dat, 70
FINCH_DATA, 67	pres, 69
FINCH_DATA, 58	RIC, 64
beta, 65	res, 69
CC_E, 66	resettime, 70
CC_I, 66	Rn, 69
CL_E, 66	Rnp1, 69
CL_I, 66	Ro, 64
CR_E, 66	s, <mark>63</mark>
CR I, 66	setbcs, 70
callroutine, 69	setic, 69
CheckMass, 65	setparams, 70
CN, 65	setpostprocess, 70
d, 62	setpreprocess, 70
DIC, 64	settime, 70
Dirichlet, 65	Sn, 69
discretize, 70	Snp1, 69
Dn, 68	solve, 70
Dnp1, 68	SteadyState, 65
Do, 64	T, 63
dt, 62	t, 63
dt_old, 63	t_old, 63
dz, 63	tol_abs, 66
evalprecon, 70	tol_rel, 65
evalres, 70	total_iter, 66
ExplicitFlux, 65	u_star, 68
fC_E, 67	uAvg, 63
fC_I, 67	uAvg_old, 63
fL_E, 66	uIC, 63
fL_I, 66	uT_old, 63
fR_E, 67	ubest, 68
fR_I, 67	un, 68
Fn, 69	unm1, 68
Fnp1, 69	unp1, 68
gE, 69	uo, 64
gl, 69	Update, 65
Iterative, 65	uT, 63
kIC, 64	uz_l_E, 67
kfn, 64	uz_l_l, 67
kfnp1, 64	uz_lm1_E, 68
kn, 68	uz_lm1_l, 67
knp1, 68	uz_lp1_E, 68
ko, 64	uz_lp1_l, <mark>67</mark>
L, 63	vIC, 64
lambda_E, 65	vn, 68
lambda_l, 64	vnp1, <mark>68</mark>
LN, 65	vo, 64
max_iter, 66	FINCH_Picard
ME, 67	finch.h, 272
MI, 67	FINCH_TESTS
NE, 67	finch.h, 276
NI, 67	fL_E
nl_method, 66	FINCH_DATA, 66
NormTrack, 65	fL_I
OE, 67	FINCH_DATA, 66
OI, 67	FLORY_HUGGINS
param_data, 71	shark.h, 350
picard_dat, 70	FOM

lark.h, 288	nl_picard, 273
fR_E	ospre_discretization, 275
FINCH_DATA, 67	print2file_dim_header, 274
fR_I	print2file_newline, 274
FINCH_DATA, 67	print2file_result_new, 274
Faraday	print2file_result_old, 274
shark.h, 348	print2file_tab, 274
fiber diameter	print2file time header, 274
DOGFISH_DATA, 56	setup FINCH DATA, 273
fiber_length	Spherical, 272
DOGFISH_DATA, 56	uAverage, 272
fiber_specific_area	uTotal, 272
DOGFISH_DATA, 56	vanAlbada_discretization, 275
File_Output	finch_coord_type
SHARK DATA, 201	finch.h, 272
file dne	finch dat
error.h, 267	DOGFISH_DATA, 56
	MONKFISH_DATA, 136
file_name	
yaml_cpp_class, 251	SCOPSOWL_DATA, 184
Files	SKUA_DATA, 207
UI_DATA, 224	finch_solve_type
film_transfer	finch.h, 272
SCOPSOWL_PARAM_DATA, 191	findAllTypes
SKUA_PARAM, 212	KeyValueMap, 97
film_transfer_coeff	findType
DOGFISH_PARAM, 57	KeyValueMap, 97
MONKFISH_PARAM, 138	ValueTypePair, 248
FilmMTCoeff	Fk
egret.h, 265	BACKTRACK_DATA, 30
finch	flock.h, 276
ui.h, 386	FloryHuggins
finch.h, 269	shark.h, 351
Cartesian, 272	FloryHuggins_chemi
check_Mass, 272	shark.h, 352
Cylindrical, 272	FloryHuggins_multichemi
default_bcs, 275	shark.h, 352
default_execution, 274	FloryHuggins_multiligand
default_ic, 274	shark.h, 351
default params, 275	FloryHuggins_unsteady
default_postprocess, 276	shark.h, 351
default precon, 275	flow rate
default_preprocess, 274	MassBalance, 107
default_res, 275	SHARK DATA, 200
default reset, 276	Fn
default_solve, 274	FINCH DATA, 69
default timestep, 274	Fnp1
FINCH Picard, 272	FINCH DATA, 69
FINCH_TESTS, 276	Fobj
	•
finch_coord_type, 272 finch_solve_type, 272	GSTA_OPT_DATA, 87 fom
I_direct, 272	lark.h, 291
LARK_PJFNK, 272	formation_energy
LARK_Picard, 272	Molecule, 131
lark_picard_step, 273	formation_enthalpy
max, 272	Molecule, 130
min, 272	formation_entropy
minmod, 272	Molecule, 130
minmod_discretization, 275	Formula

Molecule, 131	res, 76
forward_rate	restart, 75
UnsteadyReaction, 245	steps, 75
forward_ref_rate	tol_abs, 75
UnsteadyReaction, 246	tol_rel, 75
fun_call	x, 76
BACKTRACK_DATA, 30	GMRESLP
PJFNK_DATA, 168	lark.h, 288
funeval	GMRESR_DATA, 76
PJFNK_DATA, 171	arg, 79
FV DIFAIR DATA 470	GCR_Output, 78
PJFNK_DATA, 170	GMRES_Output, 78
FX	gcr_abs_tol, 78
NUM_JAC_DATA, 159	gcr_dat, 79
FXP	gcr_maxit, 78
NUM_JAC_DATA, 159	gcr_rel_tol, 78
GAS	gcr_restart, 78
mola.h, 321	gmres_dat, 79
GCR DATA, 71	gmres_maxit, 78
alpha, 73	gmres_restart, 78
bestres, 73	gmres_tol, 78
bestx, 73	iter_inner, 78
beta, 73	iter_outer, 78
breakdown, 72	matvec, 79
c, 74	matvec_data, 79
c_temp, 74	N, 78
iter_inner, 72	term_precon, 79
iter_outer, 72	terminal_precon, 79
maxit, 72	total_iter, 78
Output, 73	GMRESRP_DATA, 79
r, 73	bestres, 82
relres, 73	bestx, 82
relres_base, 73	e0, <mark>82</mark>
res, 73	e0_bar, <mark>82</mark>
restart, 72	H, 82
tol_abs, 73	H_bar, 82
tol_rel, 73	iter_inner, 81
total_iter, 72	iter_outer, 81
transpose_dat, 74	iter_total, 81
u, 74	maxit, 81
u temp, 74	Output, 82
x, 73	r, 82
GCR_Output	relres, 81
GMRESR_DATA, 78	relres_base, 81
GCR	res, 81
lark.h, 288	restart, 81
GMRES_Output	sum, 83
GMRESR_DATA, 78	tol_abs, 81
GMRESLP_DATA, 74	tol_rel, 81
arnoldi_dat, 76	v, 83
bestres, 76	Vk, 82
bestx, 76	w, 83
iter, 75	x, 82
maxit, 75	y, <mark>82</mark>
Output, 76	Zk, 82
r, 76	GMRESRP
relres, 76	lark.h, 288
relres_base, 76	GMRESR

lark.h, 288	generic_error
GPAST_DATA, 83	error.h, 267
gama_inf, 84	Get_ActivationEnergy
He, 84	UnsteadyReaction, 244
Plo, 84	Get_Affinity
po, 84	UnsteadyReaction, 244
poi, 84	Get_Area
present, 84	MassBalance, 106
q, 84	Get_Delta
qo, 84	MassBalance, 105
x, 84	Get_Energy
y, 84	Reaction, 178
GSTA_DATA, 85	UnsteadyReaction, 243
dHo, 85	Get_Enthalpy
	Reaction, 177
dSo, 85	
m, 85	UnsteadyReaction, 243
qmax, 85	Get_Entropy
GSTA_OPT_DATA, 86	Reaction, 178
all_pars, 87	UnsteadyReaction, 243
best_par, 87	Get_Equilibrium
Fobj, 87	Reaction, 177
iso, 87	UnsteadyReaction, 243
Kno, 87	Get_FlowRate
n_par, 87	MassBalance, 106
norms, 87	Get_Forward
opt_qmax, 87	UnsteadyReaction, 243
P, 87	Get_ForwardRef
q, 87	UnsteadyReaction, 244
qmax, 87	Get_InitialConcentration
total_eval, 87	MassBalance, 106
gama	Get InitialValue
mSPD_DATA, 139	UnsteadyReaction, 243
gama_inf	Get_InletConcentration
GPAST_DATA, 84	MassBalance, 106
	Get MaximumValue
gas_dat	_
SCOPSOWL_DATA, 184	UnsteadyReaction, 243
SKUA_DATA, 207	Get_Name
gas_temperature	MassBalance, 106
MIXED_GAS, 123	Get_Reverse
SCOPSOWL_DATA, 182	UnsteadyReaction, 244
gas_velocity	Get_ReverseRef
SCOPSOWL_DATA, 182	UnsteadyReaction, 244
SKUA_DATA, 206	Get_Species_Index
gcr	UnsteadyReaction, 243
lark.h, 295	Get_Stoichiometric
gcr_abs_tol	Reaction, 177
GMRESR_DATA, 78	UnsteadyReaction, 243
gcr_dat	Get_TimeStep
GMRESR_DATA, 79	MassBalance, 106
PJFNK DATA, 171	UnsteadyReaction, 244
gcr_maxit	Get TotalConcentration
GMRESR_DATA, 78	MassBalance, 106
gcr_rel_tol	Get_Type
GMRESR_DATA, 78	MassBalance, 106
gcr_restart	Get_Volume
GMRESR_DATA, 78	MassBalance, 106
gE	get_index MasterSpeciasList 111
FINCH_DATA, 69	MasterSpeciesList, 111

get species	getChemisorptionObject
MasterSpeciesList, 111	MultiligandChemisorption, 155
getActivity	getDataMap
AdsorptionReaction, 18	Document, 51
ChemisorptionReaction, 45	Header, 91
MultiligandAdsorption, 146	getDelta
MultiligandChemisorption, 156	ChemisorptionReaction, 45
UnsteadyAdsorption, 234	getDocFromHeadAlias
getActivityEnum	YamlWrapper, 255
-	getDocFromSubAlias
AdsorptionReaction, 19	
ChemisorptionReaction, 46	YamlWrapper, 255
MultiligandAdsorption, 146	getDocMap
MultiligandChemisorption, 156	YamlWrapper, 254
UnsteadyAdsorption, 235	getDocument
getAdsorbIndex	YamlWrapper, 254
AdsorptionReaction, 19	getDouble
ChemisorptionReaction, 46	KeyValueMap, 97
UnsteadyAdsorption, 235	ValueTypePair, 249
getAdsorbentName	getElectricPotential
AdsorptionReaction, 19	MultiligandAdsorption, 147
ChemisorptionReaction, 46	MultiligandChemisorption, 156
MultiligandAdsorption, 147	getHeadFromSubAlias
MultiligandChemisorption, 157	Document, 53
UnsteadyAdsorption, 235	getHeadMap
getAdsorptionObject	Document, 51
MultiligandAdsorption, 146	getHeader
getAlias	Document, 51
Document, 52	getInt
Header, 93	KeyValueMap, 97
SubHeader, 216	ValueTypePair, 249
getAnchoredDoc	getlonicStrength
YamlWrapper, 255	AdsorptionReaction, 18
getAnchoredHeader	ChemisorptionReaction, 46
Document, 53	MultiligandAdsorption, 147
getAnchoredSub	MultiligandChemisorption, 156
Header, 93	UnsteadyAdsorption, 235
getAqueousIndex	getLigandIndex
AdsorptionReaction, 19	ChemisorptionReaction, 46
UnsteadyAdsorption, 235	getLigandName
getAreaFactor	MultiligandAdsorption, 147
_	
AdsorptionReaction, 18	MultiligandChemisorption, 156
ChemisorptionReaction, 45	getMap
UnsteadyAdsorption, 234	KeyValueMap, 96
getBool	SubHeader, 215
KeyValueMap, 97	getMolarFactor
ValueTypePair, 249	AdsorptionReaction, 17
getBulkDensity	UnsteadyAdsorption, 234
AdsorptionReaction, 18	getName
ChemisorptionReaction, 46	Document, 52
MultiligandAdsorption, 146	Header, 92
MultiligandChemisorption, 156	SubHeader, 216
UnsteadyAdsorption, 234	getNumberLigands
getChargeDensity	MultiligandAdsorption, 146
AdsorptionReaction, 18	MultiligandChemisorption, 156
ChemisorptionReaction, 46	getNumberRxns
MultiligandAdsorption, 146	AdsorptionReaction, 18
MultiligandChemisorption, 156	ChemisorptionReaction, 46
UnsteadyAdsorption, 234	UnsteadyAdsorption, 235

getOldActivity	FINCH_DATA, 69
UnsteadyAdsorption, 234	gmres_dat
getPair	GMRESR_DATA, 79
KeyValueMap, 98	gmres_in
ValueTypePair, 249	KMS_DATA, 100
getReaction	gmres_maxit
AdsorptionReaction, 17	GMRESR_DATA, 78
ChemisorptionReaction, 45	gmres_out
UnsteadyAdsorption, 233	KMS_DATA, 100
getSpecificArea	gmres_restart
AdsorptionReaction, 18	GMRESR_DATA, 78
ChemisorptionReaction, 45	gmres_tol
MultiligandAdsorption, 146	GMRESR_DATA, 78
MultiligandChemisorption, 156	gmresLeftPreconditioned
UnsteadyAdsorption, 234	lark.h, 289
getSpecificMolality	gmresRightPreconditioned
AdsorptionReaction, 18	lark.h, 292
ChemisorptionReaction, 46	gmreslp_dat
UnsteadyAdsorption, 234	PJFNK_DATA, 170
getState	gmresr
Document, 52	lark.h, 296
Header, 93	gmresr_dat
SubHeader, 216	PJFNK_DATA, 171
getString KovVolueMop, 97	gmresrPreconditioner
KeyValueMap, 97 ValueTypePair, 249	lark.h, 296
getSubHeader	gmresrp_dat PJFNK_DATA, 170
Header, 91	gpast_dat
getSubMap	MAGPIE_DATA, 101
Header, 91	grad_mSPD
getSurfaceCharge	magpie.h, 308
AdsorptionReaction, 18	Grav R
UnsteadyAdsorption, 234	Trajectory.h, 382
getTotalMass	Grav_T
AdsorptionReaction, 18	Trajectory.h, 382
ChemisorptionReaction, 46	gsta_dat
MultiligandAdsorption, 146	MAGPIE_DATA, 101
MultiligandChemisorption, 156	gsta_opt
UnsteadyAdsorption, 234	ui.h, 386
getTotalVolume	gsta_opt.h, 277
AdsorptionReaction, 18	avgPar, 280
ChemisorptionReaction, 46	avgValue, 280
MultiligandAdsorption, 146	eduGuess, 281
MultiligandChemisorption, 156	eval_GSTA, 283
UnsteadyAdsorption, 234	gsta_optimize, 283
getType	gstaFunc, 281
KeyValueMap, 98	gstaObjFunc, 283
ValueTypePair, 249	isSmooth, 281
getValue	minIndex, 280
KeyValueMap, 98	minValue, 279
ValueTypePair, 249	Na, 279
getVolumeFactor	orderMag, 279
AdsorptionReaction, 18	orthoLinReg, 281
ChemisorptionReaction, 45	Po, 279
UnsteadyAdsorption, 234	R, 279
getYamlWrapper	rSq, <mark>280</mark>
yaml_cpp_class, 251 gl	roundlt, 279 twoFifths, 279

weightedAvg, 280	Document, 53
gsta_optimize	Header, 88
gsta_opt.h, 283	\sim Header, 90
gstaFunc	addPair, 92
gsta_opt.h, 281	addSubKey, 92
gstaObjFunc	begin, 91
gsta_opt.h, 283	changeKey, 92
Н	clear, 91
GMRESRP_DATA, 82	copyAnchor2Alias, 92
TRAJECTORY DATA, 222	DisplayContents, 92
H0	end, 91
TRAJECTORY DATA, 221	getAlias, 93
H bar	getAnchoredSub, 93
GMRESRP DATA, 82	getDataMap, 91
HELP	getName, 92
ui.h, 386	getState, 93
Hamaker	getSubHeader, 91 getSubMap, 91
TRAJECTORY DATA, 221	Header, 90
HaveEnergy	isAlias, 93
Molecule, 129	isAnchor, 93
HaveEquil	operator(), 91
Reaction, 179	operator=, 90
haveEquilibrium	operator[], 90, 91
Reaction, 177	resetKeys, 91
UnsteadyReaction, 242	setAlias, 92
HaveForRef	setName, 92
UnsteadyReaction, 246	setNameAliasPair, 92
HaveForward	setState, 92
UnsteadyReaction, 246	
haveForward	size, 92 Sub_Map, 93
UnsteadyReaction, 243	header state
haveForwardRef	yaml wrapper.h, 395
UnsteadyReaction, 242	help
HaveHS	ui.h, 387
Molecule, 129	Heterogeneous
Reaction, 179	SCOPSOWL_DATA, 182
haveHS	Hkp1
Molecule, 131	ARNOLDI DATA, 23
haveMinMax	hp1
MONKFISH_DATA, 134	ARNOLDI_DATA, 22
haveRate	,, <u></u> _, ,, , <u></u>
UnsteadyReaction, 242	1
HaveRevRef	SYSTEM_DATA, 218
UnsteadyReaction, 246	IDEAL_ADS
HaveReverse	shark.h, 350
UnsteadyReaction, 246	IDEAL
haveReverse	shark.h, 350
UnsteadyReaction, 243	INT
haveReverseRef	yaml_wrapper.h, 395
UnsteadyReaction, 243	ldeal
HaveG	SYSTEM_DATA, 219
Reaction, 179	ideal_solution
haveG	shark.h, 354
Molecule, 131	li
He	OPTRANS_DATA, 160
GPAST_DATA, 84	In_P_Velocity
magpie.h, 306	Trajectory.h, 382
Head_Map	In_PVel_Rad

Trajectory.h, 382	Matrix, 118
In_PVel_Theta	interparticle_diffusion
Trajectory.h, 382	MONKFISH_PARAM, 137
IncludeSurfCharge	intraparticle_diffusion
AdsorptionReaction, 21	DOGFISH_PARAM, 57
MultiligandAdsorption, 148	MONKFISH_PARAM, 138
MultiligandChemisorption, 158	invalid_atom
includeSurfaceCharge	error.h, 268
AdsorptionReaction, 19	invalid_boolean
ChemisorptionReaction, 46	error.h, 268
MultiligandAdsorption, 147	invalid_components
MultiligandChemisorption, 156	error.h, 267
UnsteadyAdsorption, 235	invalid_console_input
indexing_error	error.h, 268
error.h, 267	invalid_electron
initial_error	error.h, 268
error.h, 268	invalid_fraction
initial_guess_SCOPSOWL	error.h, 268
scopsowl_opt.h, 340	invalid_gas_sum
initial_guess_SKUA	error.h, 268
skua_opt.h, 377	invalid_input
initial_sorption	ui.h, 389
DOGFISH_PARAM, 58	invalid_molefraction
MONKFISH_PARAM, 137	error.h, 268
initial_value	invalid_neutron
UnsteadyReaction, 245	error.h, 268
InitialConcentration	invalid_norm
MassBalance, 108	error.h, 268
initialGuess_mSPD	invalid_proton
magpie.h, 309	error.h, 268
Initialize_Object	invalid_size
AdsorptionReaction, 12	error.h, 268
ChemisorptionReaction, 41	invalid_solid_sum
MassBalance, 104	error.h, 268
MultiligandAdsorption, 142	invalid_species
MultiligandChemisorption, 152	error.h, 268
Reaction, 176	invalid_type
UnsteadyAdsorption, 228	error.h, 268
UnsteadyReaction, 239	invalid_valence
initialize_data	error.h, 268
egret.h, 265	inverse
InletConcentration	Matrix, 117
MassBalance, 108	ionic_strength
inner_iter	AdsorptionReaction, 21
KMS_DATA, 100	MultiligandAdsorption, 148
inner_product	MultiligandChemisorption, 158
Matrix, 116	SHARK_DATA, 200
inner_reltol	isAlias
KMS_DATA, 100	Document, 53
input	Header, 93
ui.h, 388	SubHeader, 216
input_file	isAnchor
yaml_cpp_class, 251	Document, 53
input_files	Header, 93
UI_DATA, 224	SubHeader, 216
IntegralAvg	isAreaBasis
Matrix, 118	AdsorptionReaction, 19
IntegralTotal	UnsteadyAdsorption, 235

isEven	restart, 99
yaml_wrapper.h, 396	term_precon, 101
isRegistered	terminal_precon, 100
Molecule, 129	total iter, 100
isSmooth	KMS
gsta_opt.h, 281	lark.h, 288
isSteadyState	kB
MassBalance, 106	magpie.h, 306
isZeroInitialSolids	shark.h, 348
MassBalance, 106	Key_Value
iso	KeyValueMap, 98
GSTA_OPT_DATA, 87	key_not_found
iter	error.h, 268
ARNOLDI DATA, 22	KeyValueMap, 93
BiCGSTAB DATA, 32	•
CGS DATA, 36	~KeyValueMap, 95
GMRESLP DATA, 75	addKey, 96
PCG DATA, 161	addPair, 97
PICARD DATA, 165	assertType, 97
iter inner	begin, 96
-	clear, 96
GCR_DATA, 72	DisplayMap, 97
GMRESR_DATA, 78	editValue4Key, 96
GMRESRP_DATA, 81	end, <mark>96</mark>
iter_outer	findAllTypes, 97
GCR_DATA, 72	findType, 97
GMRESR_DATA, 78	getBool, 97
GMRESRP_DATA, 81	getDouble, 97
iter_total	getInt, 97
GMRESRP_DATA, 81	getMap, 96
Iterative	getPair, 98
FINCH_DATA, 65	getString, 97
	getType, 98
J	getValue, 98
SYSTEM_DATA, 218	Key_Value, 98
jacvec	KeyValueMap, 95
lark.h, 299	operator=, 96
	operator[], 96
K	size, 97
SYSTEM_DATA, 218	kfn
k	FINCH DATA, 64
ARNOLDI_DATA, 22	<u> </u>
TRAJECTORY_DATA, 221	kfnp1
kIC	FINCH_DATA, 64
FINCH_DATA, 64	kinematic_viscosity
KMS_DATA, 98	MIXED_GAS, 123
gmres_in, 100	kms_dat
gmres_out, 100	PJFNK_DATA, 171
inner_iter, 100	kmsPreconditioner
inner_reltol, 100	lark.h, 297
level, 99	kn
matvec, 100	FINCH_DATA, 68
matvec_data, 101	Kno
max_level, 99	GSTA_OPT_DATA, 87
maxit, 99	knp1
outer_abstol, 100	FINCH_DATA, 68
outer_iter, 100	ko
outer_reltol, 100	FINCH_DATA, 64
Output_inner, 100	krylov_method
Output_outer, 100	lark.h, 288
 	,

krylovMultiSpace	NumericalJacobian, 301
lark.h, 297	operatorTranspose, 294
	PCG, 288
L	pcg, 292
FINCH_DATA, 63	picard, 298
TRAJECTORY_DATA, 221	pjfnk, 300
L_Output	QRsolve, 298
PJFNK DATA, 169	
I direct	QR, 288
finch.h, 272	update_arnoldi_solution, 288
l iter	lark_picard_step
PJFNK_DATA, 168	finch.h, 273
	LengthFactor
L_wire	shark.h, 348
TRAJECTORY_DATA, 221	level
LARK_PJFNK	KMS_DATA, 99
finch.h, 272	MONKFISH_DATA, 134
LARK_Picard	SCOPSOWL_DATA, 181
finch.h, 272	ligand index
LARK_TESTS	ChemisorptionReaction, 47
lark.h, 301	•
LIQUID	ligand_obj
mola.h, 320	MultiligandAdsorption, 148
LOCATION	MultiligandChemisorption, 158
Trajectory.h, 383	lin_precon
ladshawSolve	SHARK_DATA, 203
Matrix, 117	lin_tol_abs
lambda_E	PJFNK_DATA, 169
FINCH DATA, 65	lin_tol_rel
-	PJFNK DATA, 169
lambda_l	LineSearch
FINCH_DATA, 64	PJFNK DATA, 170
lambdaMin	linear solver
BACKTRACK_DATA, 30	PJFNK DATA, 169
lark	linearsolve_choice
ui.h, 386	shark.h, 355
lark.h, 284	linesearch choice
arnoldi, 289	-
backtrackLineSearch, 299	shark.h, 355
BiCGSTAB, 288	List
bicgstab, 293	AdsorptionReaction, 19
CGS, 288	MassBalance, 107
cgs, 294	MultiligandAdsorption, 147
FOM, 288	MultiligandChemisorption, 157
fom, 291	Reaction, 178
GCR, 288	list_size
GMRESLP, 288	MasterSpeciesList, 111
GMRESRP, 288	LN
GMRESR, 288	FINCH_DATA, 65
	InKo
gcr, 295	magpie.h, 306
gmresLeftPreconditioned, 289	Inact mSPD
gmresRightPreconditioned, 292	-
gmresr, 296	magpie.h, 308
gmresrPreconditioner, 296	LocalMin
jacvec, 299	SHARK_DATA, 201
KMS, 288	lowerHessenberg2Triangular
kmsPreconditioner, 297	Matrix, 120
krylov_method, 288	lowerHessenbergSolve
krylovMultiSpace, 297	Matrix, 120
LARK_TESTS, 301	lowerTriangularSolve
MIN_TOL, 288	Matrix, 119
_ ,	,

M	max_fiber_density, 135
TRAJECTORY_DATA, 222	max_porosity, 135
m	min_fiber_density, 135
GSTA_DATA, 85	min_porosity, 135
M_PI	MultiScale, 134
macaw.h, 303	NonLinear, 134
mola.h, 320	NumComp, 134
m_rand	Output, 135
TRAJECTORY_DATA, 222	param_dat, 136
MACAW_TESTS	Print2Console, 133
 macaw.h, 303	Print2File, 133
MAGPIE_DATA, 101	single_fiber_density, 134
gpast_dat, 101	t_counter, 134
gsta_dat, 101	t_print, 134
mspd_dat, 101	time, 133
sys_dat, 101	time_old, 133
MAGPIE SCENARIOS	total sorption, 134
magpie.h, 311	total_sorption, 104
MAGPIE	total_sorption_old, 104
magpie.h, 311	user data, 136
mError	MONKFISH PARAM, 136
-	<u> </u>
error.h, 267	avg_sorption, 138
MIN_TOL	avg_sorption_old, 138
lark.h, 288	exterior_concentration, 137
MIXED_GAS, 121	exterior_transfer_coeff, 137
binary_diffusion, 123	film_transfer_coeff, 138
char_length, 123	initial_sorption, 137
CheckMolefractions, 122	interparticle_diffusion, 137
gas_temperature, 123	intraparticle_diffusion, 138
kinematic_viscosity, 123	sorbed_molefraction, 137
molefraction, 123	sorption_bc, 138
N, 122	species, 138
Reynolds, 123	MONKFISH_TESTS
species_dat, 123	monkfish.h, 325
total_density, 123	mSPD_DATA, 138
total_dyn_vis, 123	eMax, 139
total_molecular_weight, 123	eta, 139
total_pressure, 122	gama, 139
total_specific_heat, 123	s, 139
velocity, 123	v, 139
MOLA_TESTS	macaw
mola.h, 321	ui.h, 386
MONKFISH_DATA, 131	macaw.h, 301
avg_fiber_density, 135	M_PI, 303
DirichletBC, 134	MACAW_TESTS, 303
dog_dat, 136	Magnetic_R
domain_diameter, 135	Trajectory.h, 382
end_time, 134	Magnetic_T
eval_Cex, 136	Trajectory.h, 382
eval_Dex, 135	magpie
eval_Ret, 135	ui.h, 386
eval_ads, 135	magpie.h, 303
eval_eps, 135	A, 306
eval_kf, 136	DBL_EPSILON, 305
eval_rho, 135	dq_dp, 307
finch_dat, 136	eMax, 308
haveMinMax, 134	eval_GPAST, 310
level, 134	eval_eta, 310
	_ ·

eval_po, 310	Set_Name, 105
eval_po_PI, 309	Set_SteadyState, 105
eval_po_qo, 309	Set_TimeStep, 105
grad_mSPD, 308	Set_TotalConcentration, 104
He, 306	Set_Type, 105
initialGuess_mSPD, 309	Set_Volume, 105
kB, 306	Set_ZeroInitialSolids, 105
InKo, 306	SteadyState, 108
Inact_mSPD, 308	Sum_Delta, 105
MAGPIE_SCENARIOS, 311	TotalConcentration, 107
MAGPIE, 311	Type, 107
Na, 306	volume, 107
PI, 307	xsec_area, 107
	ZeroInitialSolids, 108
Po, 306	
q_p, 307	MassBalanceList
qo, 306	SHARK_DATA, 196
Qst, 307	MasterList
qT, 308	SHARK_DATA, 196
R, 306	MasterSpeciesList, 108
shapeFactor, 306	\sim MasterSpeciesList, 109
V, 306	alkalinity, 111
Z, 305	charge, 111
magpie_dat	DisplayAll, 110
SCOPSOWL_DATA, 184	DisplayConcentrations, 110
SKUA_DATA, 207	DisplayInfo, 110
magpie_reverse_error	Eval_ChargeResidual, 111
error.h, 267	get_index, 111
MassBalance, 102	get_species, 111
∼MassBalance, 104	list_size, 111
Delta, 107	MasterSpeciesList, 109, 110
Display Info, 104	operator=, 110
dt, 107	residual alkalinity, 112
Eval_IC_Residual, 107	set_alkalinity, 110
Eval Residual, 106	set list size, 110
flow_rate, 107	set species, 110
Get_Area, 106	size, 112
Get_Delta, 105 Get_FlowRate, 106	species, 112 speciesName, 111
Get InitialConcentration, 106	Matrix
Get_InletConcentration, 106	~Matrix, 115
Get_Name, 106	adjoint, 117
Get_TimeStep, 106	cofactor, 116
Get_TotalConcentration, 106	columnExtend, 121
Get_Type, 106	columnExtract, 120
Get_Volume, 106	columnProjection, 119
InitialConcentration, 108	columnReplace, 120
Initialize_Object, 104	columnShrink, 120
InletConcentration, 108	columnVectorFill, 119
isSteadyState, 106	columns, 116
isZeroInitialSolids, 106	ConstantICFill, 118
List, 107	Data, 121
MassBalance, 104	determinate, 116
Name, 108	diagonalSolve, 119
Set_Area, 105	dirichletBCFill, 119
Set_Delta, 104	Display, 117
Set FlowRate, 105	edit, 116
Set InitialConcentration, 105	inner_product, 116
Set_InletConcentration, 105	IntegralAvg, 118

IntegralTotal, 118	SCOPSOWL_OPT_DATA, 187
inverse, 117	SKUA_OPT_DATA, 209
ladshawSolve, 117	max_iter
lowerHessenberg2Triangular, 120	FINCH_DATA, 66
lowerHessenbergSolve, 120	max_level
lowerTriangularSolve, 119	KMS_DATA, 99
Matrix, 115	max_norm
naturalLaplacian3D, 117	SYSTEM DATA, 219
norm, 116	max_porosity
num_cols, 121	MONKFISH DATA, 135
num_rows, 121	max_value
operator*, 116, 117	UnsteadyReaction, 245
operator(), 115	maxit
operator+, 116	BiCGSTAB DATA, 32
operator-, 116	CGS DATA, 36
operator/, 116	GCR DATA, 72
operator=, 115	GMRESLP_DATA, 75
•	
outer_product, 117	GMRESRP_DATA, 81
qrSolve, 120	KMS_DATA, 99
rowExtend, 121	PCG_DATA, 161
rowExtract, 120	PICARD_DATA, 165
rowReplace, 120	ME
rowShrink, 120	FINCH_DATA, 67
rows, 116	MI
set_size, 115	FINCH_DATA, 67
SolnTransform, 118	min
sphericalAvg, 118	finch.h, 272
sphericalBCFill, 118	min_bias
sum, 116	SCOPSOWL_OPT_DATA, 187
transpose, 117	SKUA_OPT_DATA, 210
transpose_multiply, 117	min_fiber_density
tridiagonalFill, 117	MONKFISH_DATA, 135
tridiagonalSolve, 117	min porosity
tridiagonalVectorFill, 119	MONKFISH_DATA, 135
upperHessenberg2Triangular, 119	minIndex
upperHessenbergSolve, 120	gsta_opt.h, 280
upperTriangularSolve, 119	minValue
zeros, 115	gsta_opt.h, 279
Matrix < T >, 112	minmod
matrix_too_small	finch.h, 272
error.h, 268	minmod discretization
matvec	finch.h, 275
GMRESR_DATA, 79	missing_information
KMS DATA, 100	error.h, 268
matvec data	
GMRESR DATA, 79	MissingArg UI DATA, 224
<u> </u>	- · · · ·
KMS_DATA, 101	modifyDeltas
matvec_mis_match	AdsorptionReaction, 12
error.h, 268	MultiligandAdsorption, 143
max	UnsteadyAdsorption, 228
finch.h, 272	modifyMBEdeltas
UI_DATA, 224	ChemisorptionReaction, 41
max_bias	MultiligandChemisorption, 152
SCOPSOWL_OPT_DATA, 187	mola
SKUA_OPT_DATA, 210	ui.h, 386
max_fiber_density	mola.h, 313
MONKFISH_DATA, 135	ADSORBED, 321
max_guess_iter	AQUEOUS, 321

GAS, 321	isRegistered, 129
LIQUID, 320	molar_area, 130
M_PI, 320	molar_volume, 130
MOLA_TESTS, 321	molar_weight, 130
OTHER, 321	MolarArea, 129
PLASMA, 321	MolarVolume, 129
SOLID, 320	MolarWeight, 129
SphereArea, 320	MolecularFormula, 130
SphereVolume, 320	Molecule, 126
valid_phase, 320	MoleculeName, 130
molar_area	MoleculePhase, 130
Molecule, 130	MoleculePhaseID, 130
molar_factor	Name, 131
AdsorptionReaction, 20	Phase, 131
molar_volume	PhaseID, 131
Molecule, 130	Register, 127
molar_weight	registered, 131
Molecule, 130	removeAllAtoms, 129
MolarArea	removeOneAtom, 129
Molecule, 129	setFormula, 127
MolarVolume	setMolarArea, 128
Molecule, 129	setMolarVolume, 128
MolarWeight	setMolarWeigth, 128
Molecule, 129	MoleculeName
molecular_diffusion	Molecule, 130
PURE GAS, 173	Molecule Phase
- · · ·	
molecular_weight PURE GAS, 172	Molecule, 130 MoleculePhaseID
MolecularFormula	
	Molecule, 130
Molecule, 130	molefraction
Molecule, 124	MIXED_GAS, 123
~Molecule, 126	molefractionCheck
atoms, 131	skua.h, 371
calculateAvgOxiState, 128	monkfish
calculateMolarArea, 128	ui.h, 386
calculateMolarVolume, 128	monkfish.h, 321
calculateMolarWeight, 128	default_density, 323
Charge, 129	default_exterior_concentration, 324
charge, 130	default_film_transfer, 324
DisplayInfo, 130	default_interparticle_diffusion, 323
editAllOxidationStates, 128	default_monk_adsorption, 323
editCharge, 128	default_monk_equilibrium, 323
editEnergy, 129	default_monkfish_retardation, 324
editEnthalpy, 128	default_porosity, 322
editEntropy, 128	MONKFISH_TESTS, 325
editHS, 128	setup_MONKFISH_DATA, 324
editOneOxidationState, 128	mp
Energy, 130	TRAJECTORY_DATA, 222
Enthalpy, 129	Ms
Entropy, 129	TRAJECTORY_DATA, 221
formation_energy, 131	mspd_dat
formation_enthalpy, 130	MAGPIE_DATA, 101
formation_entropy, 130	Mu
Formula, 131	egret.h, 265
HaveEnergy, 129	mu_0
HaveHS, 129	TRAJECTORY_DATA, 221
haveHS, 131	MultiAdsList
haveG, 131	SHARK_DATA, 196

MultiChemList	setVolumeFactor, 144
SHARK_DATA, 196	specific_area, 148
MultiScale	surface_activity, 147
MONKFISH_DATA, 134	total_mass, 148
MultiligandAdsorption, 139	total_volume, 148
~MultiligandAdsorption, 142	MultiligandChemisorption, 149
act_fun, 148	~MultiligandChemisorption, 152
activities, 148	act_fun, 157
activity_data, 148	activities, 157
adsorbent_name, 147	activity_data, 157
calculateAreaFactors, 144	adsorbent_name, 157
calculateElecticPotential, 145	calculateAreaFactors, 154
calculateEquilibria, 144	calculateElecticPotential, 154
calculateEquilibriumCorrection, 145	calculateEquilibria, 154
callSurfaceActivity, 145	calculateEquilibriumCorrection, 155
charge_density, 148	callSurfaceActivity, 154
checkAqueousIndices, 143	charge_density, 158
electric_potential, 148	Display_Info, 152
Eval_Residual, 145	electric_potential, 158
getActivity, 146	Eval_RxnResidual, 155
getActivityEnum, 146	Eval_SiteBalanceResidual, 155
getAdsorbentName, 147	getActivity, 156
getAdsorptionObject, 146	getActivityEnum, 156
getBulkDensity, 146	getAdsorbentName, 157
getChargeDensity, 146	getBulkDensity, 156
getElectricPotential, 147	getChargeDensity, 156
getIonicStrength, 147	getChemisorptionObject, 155
getLigandName, 147	getElectricPotential, 156
getNumberLigands, 146	getIonicStrength, 156
getSpecificArea, 146	getLigandName, 156
getTotalMass, 146	getNumberLigands, 156
getTotalVolume, 146	getSpecificArea, 156
IncludeSurfCharge, 148	getTotalMass, 156
includeSurfaceCharge, 147	getTotalVolume, 156
Initialize_Object, 142	IncludeSurfCharge, 158
ionic_strength, 148	includeSurfaceCharge, 156
ligand_obj, 148	Initialize_Object, 152
List, 147	ionic_strength, 158
modifyDeltas, 143	ligand_obj, 158
MultiligandAdsorption, 142	List, 157
num_ligands, 147	modifyMBEdeltas, 152
setActivityEnum, 143	MultiligandChemisorption, 152
setActivityModelInfo, 143	num_ligands, 157
setAdsorbIndices, 143	setActivityEnum, 153
setAdsorbentName, 144	setActivityModelInfo, 153
setAqueousIndexAuto, 143	setAdsorbIndices, 152
setAreaFactor, 144	setAdsorbentName, 153
setChargeDensity, 145	setAreaFactor, 153
setElectricPotential, 144	setChargeDensity, 154
setIonicStrength, 145	setDeltas, 153
setLigandName, 144	setElectricPotential, 154
setMolarFactor, 143	setIonicStrength, 154
setSpecificArea, 144	setLigandIndices, 152
setSpecificMolality, 144	setLigandName, 153
setSurfaceCharge, 144	setSpecificArea, 153
setSurfaceChargeBool, 144	setSpecificMolality, 153
setTotalMass, 144	setSurfaceChargeBool, 154
setTotalVolume, 144	setTotalMass, 153
•	•

setTotalVolume, 154	FINCH_DATA, 66
setVolumeFactor, 153	nl_picard
specific_area, 157	finch.h, 273
surface_activity, 157	nl_relres
total_mass, 158	PJFNK_DATA, 169
total_volume, 158	nl_res
N.	PJFNK_DATA, 169
N CAUDEOD DATA TO	nl_res_base
GMRESR_DATA, 78	PJFNK_DATA, 169
MIXED_GAS, 122	nl_tol_abs
SYSTEM_DATA, 218	PJFNK_DATA, 169
n_par	nl_tol_rel
GSTA_OPT_DATA, 87	PJFNK DATA, 169
n_rand	no_diffusion
TRAJECTORY_DATA, 222	error.h, 268
NL_Output	non_real_edge
PJFNK_DATA, 169	error.h, 268
NONE	non_square_matrix
yaml_wrapper.h, 396	error.h, 268
NUM_JAC_DATA, 158	NonLinear
dxj, 159	DOGFISH DATA, 55
eps, 159	MONKFISH_DATA, 134
Fx, 159	SCOPSOWL DATA, 183
Fxp, 159	
Na	SKUA_DATA, 206 Norm
gsta_opt.h, 279	
magpie.h, 306	SHARK_DATA, 200
shark.h, 348	norm
Name	Matrix, 116
Atom, 29	normFkp1
MassBalance, 108	BACKTRACK_DATA, 30
Molecule, 131	NormTrack
name	FINCH_DATA, 65
SubHeader, 216	norms
naturalLaplacian3D	GSTA_OPT_DATA, 87
Matrix, 117	not_a_token
NaturalState	error.h, 268
Atom, 29	Nu
NE	egret.h, 265
FINCH_DATA, 67	nullptr_error
negative_mass	error.h, 268
error.h, 268	nullptr_func
negative time	error.h, 268
error.h, 268	num_cols
Neutrons	Matrix, 121
Atom, 27	num_curves
neutrons	SCOPSOWL_OPT_DATA, 186
Atom, 28	SKUA_OPT_DATA, 209
Newton_data	num_ligands
SHARK_DATA, 203	MultiligandAdsorption, 147
NI – ,	MultiligandChemisorption, 157
FINCH_DATA, 67	num_mbe
nl bestres	SHARK_DATA, 197
PJFNK_DATA, 169	num_multi_ssao
nl iter	SHARK DATA, 197
PJFNK_DATA, 168	num_multi_ssar
nl maxit	SHARK DATA, 198
PJFNK_DATA, 168	num_multi_sschem
nl_method	SHARK DATA, 197

num_multichem_rxns	Matrix, 115
SHARK_DATA, 198	YamlWrapper, 254
num_other	operator+
SHARK_DATA, 198	Matrix, 116
num_params	operator-
SCOPSOWL_OPT_DATA, 186	Matrix, 116
SKUA_OPT_DATA, 209	operator/
num_rows	Matrix, 116
Matrix, 121	operator=
num_rxns	Document, 50
AdsorptionReaction, 21	Header, 90
num_ssao	KeyValueMap, 96
SHARK_DATA, 197	MasterSpeciesList, 110
num_ssar	Matrix, 115
SHARK_DATA, 197	SubHeader, 215
num_sschem	ValueTypePair, 248
SHARK_DATA, 197	YamlWrapper, 254
num_sschem_rxns	operatorTranspose
SHARK_DATA, 197	lark.h, 294
NUM_SST	operator[]
SHARK_DATA, 197	Document, 50
NUM_USAO	Header, 90, 91
SHARK_DATA, 197	KeyValueMap, 96
num_usar SHARK_DATA, 197	SubHeader, 215
	opt_no_support
num_usr SHARK DATA, 197	error.h, 268
NumComp	opt_qmax
DOGFISH_DATA, 55	GSTA_OPT_DATA, 87
MONKFISH DATA, 134	Optimize
Number_Generator	SCOPSOWL_OPT_DATA, 187
Trajectory.h, 383	SKUA_OPT_DATA, 209
number_elements	option
PeriodicTable, 164	UI_DATA, 224
number_files	orderMag
ui.h, 388	gsta_opt.h, 279
NumericalJacobian	ortho_check_fail
lark.h, 301	error.h, 268
numvar	orthoLinReg
SHARK_DATA, 197	gsta_opt.h, 281
5. m. n. <u>-</u> 5. m., 107	ospre_discretization
OPTRANS_DATA, 159	finch.h, 275
Ai, 160	other_data
li, 160	SHARK_DATA, 203
OTHER	OtherList
mola.h, 321	SHARK_DATA, 196
OE	out_of_bounds
FINCH_DATA, 67	error.h, 268
OI	outer_abstol
FINCH_DATA, 67	KMS_DATA, 100
omega	outer_iter
BiCGSTAB_DATA, 32	KMS_DATA, 100
omega_old	outer_product
BiCGSTAB_DATA, 33	Matrix, 117
operator*	outer_reltol
Matrix, 116, 117	KMS_DATA, 100
operator()	Output
Document, 50	ARNOLDI_DATA, 23
Header, 91	BiCGSTAB_DATA, 33

CGS_DATA, 37	SHARK_DATA, 199
GCR_DATA, 73	pH_step
GMRESLP_DATA, 76	SHARK DATA, 200
GMRESRP_DATA, 82	PICARD DATA, 164
MONKFISH DATA, 135	bestres, 166
PCG DATA, 162	bestx, 166
PICARD DATA, 166	iter, 165
SYSTEM DATA, 219	
Output_inner	maxit, 165
KMS DATA, 100	Output, 166
-	r, 166
Output_outer	relres, 166
KMS_DATA, 100	relres_base, 166
OutputFile	res, 166
DOGFISH_DATA, 56	tol_abs, 165
SCOPSOWL_DATA, 183	tol_rel, 165
SHARK_DATA, 203	x0, 166
SKUA_DATA, 206	PITZER
owl_dat	shark.h, 350
SCOPSOWL_OPT_DATA, 188	Plo
oxidation_state	GPAST_DATA, 84
Atom, 28	PJFNK DATA, 166
OxidationState	backtrack dat, 171
Atom, 27	bestx, 170
	bicgstab_dat, 170
P	Bounce, 170
GSTA_OPT_DATA, 87	cgs_dat, 170
p	- —
BiCGSTAB_DATA, 34	eps, 169
CGS_DATA, 37	F, 170
PCG_DATA, 162	fun_call, 168
P_Velocity	funeval, 171
Trajectory.h, 382	Fv, 170
PCG_DATA, 160	gcr_dat, 171
alpha, 161	gmreslp_dat, 170
Ap, 162	gmresr_dat, 171
bestres, 162	gmresrp_dat, 170
bestx, 162	kms_dat, 171
beta, 161	L_Output, 169
iter, 161	I_iter, 168
maxit, 161	lin_tol_abs, 169
Output, 162	lin_tol_rel, 169
p, 162	LineSearch, 170
1.7	linear_solver, 169
r, 162	NL Output, 169
r_old, 162	nl bestres, 169
relres, 161	nl iter, 168
relres_base, 162	nl_maxit, 168
res, 161	
tol_abs, 161	nl_relres, 169
tol_rel, 161	nl_res, 169
x, 162	nl_res_base, 169
z, 162	nl_tol_abs, 169
z_old, 162	nl_tol_rel, 169
PCG	pcg_dat, 170
lark.h, 288	precon, 171
PE3	precon_data, 171
egret.h, 264	qr_dat, 171
PFR	res_data, 171
shark.h, 349	v, 170
pH_index	x, 170
· -	•

PLASMA	DisplayTable, 164
mola.h, 321	number_elements, 164
pOH_index	PeriodicTable, 163, 164
SHARK_DATA, 199	Table, 164
POLAR	рH
Trajectory.h, 382	SHARK_DATA, 199
POL	Phase - '
TRAJECTORY_DATA, 222	Molecule, 131
PSI	PhaseID
egret.h, 265	Molecule, 131
PURE_GAS, 171	PI
density, 173	
	magpie.h, 307
dynamic_viscosity, 173	SYSTEM_DATA, 218
molecular_diffusion, 173	pi
molecular_weight, 172	SYSTEM_DATA, 218
Schmidt, 173	picard
specific_heat, 172	lark.h, 298
Sutherland_Const, 172	picard_dat
Sutherland_Temp, 172	FINCH_DATA, 70
Sutherland_Viscosity, 172	pjfnk
PVel_Rad	lark.h, 300
Trajectory.h, 382	pjfnk_dat
PVel_Theta	FINCH_DATA, 70
Trajectory.h, 382	Po
Par	egret.h, 264
SYSTEM_DATA, 219	gsta_opt.h, 279
param_dat	magpie.h, 306
DOGFISH DATA, 57	po
MONKFISH_DATA, 136	GPAST DATA, 84
SCOPSOWL_DATA, 184	
	POI CRACT DATA 04
SKUA_DATA, 207	GPAST_DATA, 84
param_data	pore_diffusion
FINCH_DATA, 71	SCOPSOWL_PARAM_DATA, 191
param_guess	porosity
SCOPSOWL_OPT_DATA, 188	TRAJECTORY_DATA, 221
SKUA_OPT_DATA, 210	precon
param_guess_old	PJFNK_DATA, 171
SCOPSOWL_OPT_DATA, 188	precon_data
SKUA_OPT_DATA, 210	PJFNK_DATA, 171
ParamFile	SHARK_DATA, 203
SCOPSOWL_OPT_DATA, 188	pres
SKUA_OPT_DATA, 211	FINCH_DATA, 69
Path	present
UI_DATA, 224	GPAST DATA, 84
path	previous token
UI DATA, 224	yaml_cpp_class, 252
ui.h, 387	Print2Console
pcg	DOGFISH DATA, 55
lark.h, 292	MONKFISH DATA, 133
pcg_dat	SCOPSOWL DATA, 182
PJFNK_DATA, 170	SKUA_DATA, 206
pellet_density	Print2File
SCOPSOWL_DATA, 183	DOGFISH_DATA, 55
pellet_radius	MONKFISH_DATA, 133
SCOPSOWL_DATA, 182	SCOPSOWL_DATA, 182
SKUA_DATA, 206	SKUA_DATA, 206
PeriodicTable, 163	print2file_DOGFISH_header
\sim PeriodicTable, 163	dogfish.h, 258

print2file_DOGFISH_result_new dogfish.h, 258	SCOPSOWL_OPT_DATA, 188 SKUA_OPT_DATA, 211
print2file_DOGFISH_result_old	Q_in
dogfish.h, 258	TRAJECTORY_DATA, 221
print2file_SCOPSOWL_header	q_p
scopsowl.h, 330	magpie.h, 307
print2file_SCOPSOWL_result_new	q_sim
scopsowl.h, 330	SCOPSOWL_OPT_DATA, 188
print2file_SCOPSOWL_result_old	SKUA_OPT_DATA, 211
scopsowl.h, 330	qAvg
print2file_SCOPSOWL_time_header	SCOPSOWL_PARAM_DATA, 190
scopsowl.h, 330	qAvg_old
print2file_SKUA_header	SCOPSOWL_PARAM_DATA, 190
skua.h, 369	qIntegralAvg
print2file_SKUA_results_new	SCOPSOWL_PARAM_DATA, 190
skua.h, 369	qIntegralAvg_old
print2file_SKUA_results_old	SCOPSOWL_PARAM_DATA, 190
skua.h, 369	QR_DATA, 173
print2file_SKUA_time_header	ek, 174
skua.h, 369	Ro, 174
print2file_dim_header	x, 174
finch.h, 274	QRsolve
print2file_newline	lark.h, 298
finch.h, 274	qTn
print2file_result_new	SKUA DATA, 205
finch.h, 274	qTnp1
print2file_result_old	SKUA DATA, 206
finch.h, 274	qmax
print2file_shark_header	GSTA DATA, 85
shark.h, 350	GSTA_OPT_DATA, 87
print2file_shark_info	qo
shark.h, 350	GPAST_DATA, 84
print2file_shark_results_new	magpie.h, 306
shark.h, 350	SCOPSOWL_PARAM_DATA, 191
print2file_shark_results_old	QR
shark.h, 350	lark.h, 288
print2file_species_header	qr_dat
dogfish.h, 258	PJFNK DATA, 171
scopsowl.h, 330	grSolve
skua.h, 369	Matrix, 120
print2file_tab	_
finch.h, 274	Qst
print2file_time_header	magpie.h, 307 SCOPSOWL_PARAM_DATA, 190
finch.h, 274	
Protons	Qst_old
Atom, 27	SCOPSOWL_PARAM_DATA, 190
protons	QstAvg
Atom, 28	SCOPSOWL_PARAM_DATA, 191
Pstd	QstAvg_old
egret.h, 264	SCOPSOWL_PARAM_DATA, 191
PT	Qstn
SYSTEM_DATA, 218	SKUA_PARAM, 212
	Qstnp1
q	SKUA_PARAM, 212
GPAST_DATA, 84	Qsto
GSTA_OPT_DATA, 87	SCOPSOWL_PARAM_DATA, 191
q_bar	qT
TRAJECTORY_DATA, 222	magpie.h, 308
q_data	SYSTEM_DATA, 218

R	Set_Stoichiometric, 176
gsta_opt.h, 279	Stoichiometric, 178
magpie.h, 306	ReactionList
r	SHARK DATA, 196
BICGSTAB DATA, 33	reactor choice
CGS DATA, 37	shark.h, 355
GCR DATA, 73	reactor_type
GMRESLP DATA, 76	SHARK DATA, 198
— · · · · · · · · · · · · · · · · · · ·	- · · · ·
GMRESRP_DATA, 82	read_adsorbobjects shark.h, 358
PCG_DATA, 162	,
PICARD_DATA, 166	read_chemisorbobjects
r0	shark.h, 359
BiCGSTAB_DATA, 33	read_equilrxn
CGS_DATA, 37	shark.h, 358
r_old	read_error
PCG_DATA, 162	error.h, 268
RADIAL_FORCE	read_massbalance
Trajectory.h, 382	shark.h, 358
RE3	read_multichemi_scenario
egret.h, 264	 shark.h, 357
RIC	read_multichemiobjects
FINCH DATA, 64	shark.h, 359
rSq	read_multiligand_scenario
•	
gsta_opt.h, 280	shark.h, 357
RUN_SANDBOX	read_multiligandobjects
sandbox.h, 326	shark.h, 358
ReNum	read_options
egret.h, 265	shark.h, 357
Reaction, 174	read_scenario
\sim Reaction, 176	shark.h, 357
calculateEnergies, 177	read_species
calculateEquilibrium, 177	shark.h, 358
CanCalcHS, 178	read_unsteadyadsorbobjects
CanCalcG, 179	shark.h, 358
checkSpeciesEnergies, 177	read_unsteadyrxn
Display_Info, 176	shark.h, 358
energy, 178	readInputFile
	•
enthalpy, 178	yaml_cpp_class, 251
entropy, 178	Recover
Equilibrium, 178	SYSTEM_DATA, 219
Eval_Residual, 178	ref_diffusion
Get_Energy, 178	SCOPSOWL_PARAM_DATA, 191
Get_Enthalpy, 177	SKUA_PARAM, 212
Get_Entropy, 178	ref_pressure
Get_Equilibrium, 177	SCOPSOWL PARAM DATA, 191
Get_Stoichiometric, 177	SKUA PARAM, 212
HaveEquil, 179	ref temperature
haveEquilibrium, 177	SCOPSOWL_PARAM_DATA, 191
HaveHS, 179	SKUA_PARAM, 212
HaveG, 179	Register
Initialize_Object, 176	Atom, 26
List, 178	Molecule, 127
Reaction, 176	registered
Set_Energy, 177	Molecule, 131
Set_Enthalpy, 177	rel_tol_norm
Set_EnthalpyANDEntropy, 177	SCOPSOWL_OPT_DATA, 188
Set_Entropy, 177	SKUA_OPT_DATA, 211
Set_Equilibrium, 176	relative_permittivity
·	_ ·

SHARK_DATA, 200	Document, 51
relres	YamlWrapper, 255
BiCGSTAB_DATA, 33	reverse_rate
CGS_DATA, 36	UnsteadyReaction, 246
GCR_DATA, 73	reverse_ref_rate
GMRESLP_DATA, 76	UnsteadyReaction, 246
GMRESRP DATA, 81	Reynolds
PCG DATA, 161	MIXED GAS, 123
PICARD DATA, 166	rho
relres_base	BACKTRACK DATA, 30
BICGSTAB DATA, 33	BICGSTAB DATA, 32
CGS_DATA, 37	CGS DATA, 36
GCR DATA, 73	rho_f
GMRESLP DATA, 76	TRAJECTORY DATA, 221
GMRESRP DATA, 81	rho_old
— · · · · · · · · · · · · · · · · · · ·	BICGSTAB DATA, 32
PCG_DATA, 162	rho_p
PICARD_DATA, 166	TRAJECTORY DATA, 221
Removal_Efficiency	Rn
Trajectory.h, 383	FINCH DATA, 69
removeAllAtoms	<u> </u>
Molecule, 129	Rnp1
removeElectron	FINCH_DATA, 69
Atom, 27	Ro
removeNeutron	FINCH_DATA, 64
Atom, 27	QR_DATA, 174
removeOneAtom	Rough
Molecule, 129	SCOPSOWL_OPT_DATA, 187
removeProton	SKUA_OPT_DATA, 209
Atom, 27	roundIt
res	gsta_opt.h, 279
BiCGSTAB_DATA, 33	rowExtend
CGS DATA, 36	Matrix, 121
FINCH_DATA, 69	rowExtract
GCR DATA, 73	Matrix, 120
GMRESLP_DATA, 76	rowReplace
GMRESRP_DATA, 81	Matrix, 120
	rowShrink
PCG_DATA, 161	Matrix, 120
PICARD_DATA, 166	rows
res_data	Matrix, 116
PJFNK_DATA, 171	Rs
resetKeys	TRAJECTORY DATA, 221
Document, 51	Rstd
Header, 91	egret.h, 264
YamlWrapper, 255	shark.h, 348
resettime	Run_Trajectory
FINCH_DATA, 70	Trajectory.h, 383
Residual	run exec
SHARK_DATA, 202	_
SHARK_DATA, 202 residual_alkalinity	ui.h, 391
- · · ·	ui.h, 391 run_executable
residual_alkalinity MasterSpeciesList, 112	ui.h, 391 run_executable ui.h, 391
residual_alkalinity MasterSpeciesList, 112 residual_data	ui.h, 391 run_executable ui.h, 391 run_test
residual_alkalinity MasterSpeciesList, 112	ui.h, 391 run_executable ui.h, 391 run_test ui.h, 391
residual_alkalinity MasterSpeciesList, 112 residual_data SHARK_DATA, 203 restart	ui.h, 391 run_executable ui.h, 391 run_test ui.h, 391 rxn_rate_error
residual_alkalinity MasterSpeciesList, 112 residual_data SHARK_DATA, 203 restart GCR_DATA, 72	ui.h, 391 run_executable ui.h, 391 run_test ui.h, 391
residual_alkalinity MasterSpeciesList, 112 residual_data SHARK_DATA, 203 restart GCR_DATA, 72 GMRESLP_DATA, 75	ui.h, 391 run_executable ui.h, 391 run_test ui.h, 391 rxn_rate_error error.h, 268
residual_alkalinity MasterSpeciesList, 112 residual_data SHARK_DATA, 203 restart GCR_DATA, 72 GMRESLP_DATA, 75 GMRESRP_DATA, 81	ui.h, 391 run_executable ui.h, 391 run_test ui.h, 391 rxn_rate_error error.h, 268
residual_alkalinity MasterSpeciesList, 112 residual_data SHARK_DATA, 203 restart GCR_DATA, 72 GMRESLP_DATA, 75	ui.h, 391 run_executable ui.h, 391 run_test ui.h, 391 rxn_rate_error error.h, 268

mSPD_DATA, 139	e_norm_old, 187
s_rand	evaluation, 186
TRAJECTORY_DATA, 222	f_bias, 187
SCOPSOWL_DATA, 179	f_bias_old, 188
binder_fraction, 183	max_bias, 187
binder_poresize, 183	max_guess_iter, 187
binder_porosity, 183	min_bias, 187
char_macro, 182	num_curves, 186
char_micro, 183	num_params, 186
coord_macro, 181	Optimize, 187
coord_micro, 181	owl_dat, 188
crystal_radius, 182	param_guess, 188
DirichletBC, 183	param_guess_old, 188
eval ads, 183	ParamFile, 188
eval diff, 184	q_data, 188
eval_kf, 184	q_sim, 188
eval_retard, 183	rel_tol_norm, 188
eval surfDiff, 184	Rough, 187
finch_dat, 184	simulation equil, 187
gas_dat, 184	t, 188
gas temperature, 182	total eval, 186
gas_velocity, 182	y base, 188
Heterogeneous, 182	SCOPSOWL_OPT_set_y
level, 181	scopsowl_opt.h, 340
magpie_dat, 184	SCOPSOWL_OPTIMIZE
NonLinear, 183	scopsowl_opt.h, 340
OutputFile, 183	SCOPSOWL PARAM DATA, 189
param dat, 184	activation energy, 191
pellet_density, 183	Adsorbable, 192
pellet_radius, 182	affinity, 191
Print2Console, 182	dq dc, 190
Print2File, 182	dq dco, 191
sim_time, 181	film transfer, 191
skua dat, 184	pore diffusion, 191
SurfDiff, 182	qAvg, 190
t, 181	qAvg_old, 190
t counter, 182	qIntegralAvg, 190
t_old, 181	qIntegralAvg_old, 190
t_print, 182	qo, 191
tempy, 183	Qst, 190
total_pressure, 182	Qst_old, 190
total_steps, 181	QstAvg, 191
user_data, 184	QstAvg_old, 191
y, 183	Qsto, 191
SCOPSOWL_Executioner	ref_diffusion, 191
scopsowl.h, 333	ref_pressure, 191
SCOPSOWL_HPP_	ref_temperature, 191
scopsowl.h, 329	speciesName, 192
SCOPSOWL_OPT_DATA, 184	xIC, 190
abs_tol_bias, 188	SCOPSOWL_SCENARIOS
adsorb_index, 186	scopsowl.h, 336
CompareFile, 188	SCOPSOWL_TESTS
current_equil, 187	scopsowl.h, 338
current_points, 186	SCOPSOWL_postprocesses
current_press, 187	scopsowl.h, 334
current_temp, 187	SCOPSOWL_preprocesses
diffusion_type, 186	scopsowl.h, 334
e_norm, 187	SCOPSOWL_reset

scopsowl.h, 334	reactor_type, 198
SCOPSOWL	relative_permittivity, 200
scopsowl.h, 336	Residual, 202
SHARK_DATA, 192	residual_data, 203
act_fun, 198	shark.h, 349
activity_data, 203	simulationtime, 199
activity_new, 202	SpeciationCurve, 201
activity_old, 202	ss_ads_names, 198
AdsorptionList, 196	ss_chem_names, 198
ChemisorptionList, 196	ssmulti_names, 198
Conc_new, 202	ssmultichem_names, 198
Conc_old, 202	start_temp, 200
Console_Output, 201	steadystate, 201
const_pH, 201 Contains_pOH, 201	t_count, 199
Contains pH, 201	t_out, 199
Converged, 201	temp_step, 200 temperature, 200
dielectric_const, 200	Temperature Curve, 201
dt, 199	time, 199
dt, 199 dt_max, 199	time_old, 199
dt_max, 100 dt_min, 199	Time_old, 193
end_temp, 200	timesteps, 199
EvalActivity, 202	totalcalls, 198
File_Output, 201	totalsteps, 198
flow_rate, 200	UnsteadyAdsList, 196
ionic_strength, 200	UnsteadyList, 196
lin precon, 203	us ads names, 198
LocalMin, 201	volume, 200
MassBalanceList, 196	X_new, 202
MasterList, 196	X_old, 202
MultiAdsList, 196	xsec area, 200
MultiChemList, 196	yaml_object, 203
Newton_data, 203	ZeroInitialSolids, 201
Norm, 200	SHARK SCENARIO
num mbe, 197	shark.h, 362
num_multi_ssao, 197	SHARK TESTS OLD
num_multi_ssar, 198	shark.h, 366
num_multi_sschem, 197	SHARK_TESTS
num_multichem_rxns, 198	shark.h, 366
num_other, 198	SHARK
num_ssao, 197	shark.h, 362
num_ssar, 197	SIT
num_sschem, 197	shark.h, 350
num_sschem_rxns, 197	SKUA_DATA, 204
num_ssr, 197	char_measure, 206
num_usao, 197	coord, 205
num_usar, 197	DirichletBC, 206
num_usr, 197	eval_diff, 206
numvar, 197	eval_kf, 206
other_data, 203	finch_dat, 207
OtherList, 196	gas_dat, 207
OutputFile, 203	gas_velocity, 206
pH_index, 199	magpie_dat, 207
pH_step, 200	NonLinear, 206
pOH_index, 199	OutputFile, 206
pH, 199	param_dat, 207
precon_data, 203	pellet_radius, 206
ReactionList, 196	Print2Console, 206

Print2File, 206	ref_diffusion, 212
qTn, 205	ref_pressure, 212
qTnp1, 206	ref_temperature, 212
sim_time, 205	speciesName, 212
t, 205	xIC, 212
t_counter, 205	xn, 212
t_old, 205	xnp1, 212
t_print, 205	y_eff, 212
total_steps, 205	SKUA_SCENARIOS
user_data, 207	skua.h, <mark>373</mark>
y, 206	SKUA_TESTS
SKUA_Executioner	skua.h, <mark>375</mark>
skua.h, 372	SKUA_postprocesses
SKUA_HPP_	skua.h, 372
skua.h, 369	SKUA_preprocesses
SKUA_OPT_DATA, 207	skua.h, 372
abs_tol_bias, 211	SKUA_reset
adsorb_index, 209	skua.h, 373
CompareFile, 211	SKUA
current_equil, 210	skua.h, 373
current_points, 209	SOLID
current_press, 210	mola.h, 320 STRING
current_temp, 210	-
diffusion_type, 209 e_norm, 210	yaml_wrapper.h, 395 SYSTEM_DATA, 217
e_norm_old, 210	As, 218
evaluation, 209	avg_norm, 218
f_bias, 210	Carrier, 219
f_bias_old, 210	I, 218
max_bias, 210	Ideal, 219
max_guess_iter, 209	J, 218
min_bias, 210	K, 218
num curves, 209	max norm, 219
num_params, 209	N, 218
Optimize, 209	Output, 219
param_guess, 210	Par, 219
param_guess_old, 210	PI, 218
ParamFile, 211	pi, 218
q_data, 211	PT, 218
q_sim, 211	qT, 218
rel_tol_norm, 211	Recover, 219
Rough, 209	Sys, 219
simulation_equil, 210	T, 218
skua_dat, 211	total_eval, 218
t, 211	sandbox
total_eval, 209	ui.h, 386
y_base, 211	sandbox.h, 325
SKUA_OPT_set_y	RUN_SANDBOX, 326
skua_opt.h, 377	ScNum
SKUA_OPTIMIZE	egret.h, 265
skua_opt.h, 377	scenario_fail
SKUA_PARAM, 211	error.h, 268
activation_energy, 212	Schmidt CAS 179
Adsorbable, 212	PURE_GAS, 173
affinity, 212	school.h, 326
film_transfer, 212 Qstn, 212	scops_opt ui.h, 386
Qstrp1, 212	scopsowl
σοιιρ1, 212	Scopsowi

ui.h, 386	UnsteadyReaction, 240
scopsowl.h, 327	Set_Entropy
avgDp, 329	Reaction, 177
const_filmMassTransfer, 332	UnsteadyReaction, 240
const_pore_diffusion, 332	Set_Equilibrium
default_adsorption, 330	Reaction, 176
default_effective_diffusion, 331	UnsteadyReaction, 240
default_filmMassTransfer, 332	Set_FlowRate
default_pore_diffusion, 330	MassBalance, 105
default_retardation, 330	Set_Forward
default_surf_diffusion, 331	UnsteadyReaction, 241
Dk, 329	Set_ForwardRef
Dp, 329	UnsteadyReaction, 241
print2file_SCOPSOWL_header, 330	Set_InitialConcentration
print2file_SCOPSOWL_result_new, 330	MassBalance, 105
print2file_SCOPSOWL_result_old, 330	Set InitialValue
print2file_SCOPSOWL_time_header, 330	UnsteadyReaction, 240
print2file_species_header, 330	Set InletConcentration
SCOPSOWL Executioner, 333	MassBalance, 105
SCOPSOWL_HPP_, 329	Set_MaximumValue
SCOPSOWL_SCENARIOS, 336	UnsteadyReaction, 240
SCOPSOWL_TESTS, 338	Set_Name
SCOPSOWL_postprocesses, 334	MassBalance, 105
SCOPSOWL_preprocesses, 334	Set Reverse
SCOPSOWL reset, 334	UnsteadyReaction, 241
SCOPSOWL, 336	Set_ReverseRef
set_SCOPSOWL_ICs, 333	UnsteadyReaction, 241
set_SCOPSOWL_params, 334	set_SCOPSOWL_ICs
set_SCOPSOWL_timestep, 333	scopsowl.h, 333
setup_SCOPSOWL_DATA, 332	set_SCOPSOWL_params
zero_surf_diffusion, 331	scopsowl.h, 334
scopsowl_opt.h, 338	set_SCOPSOWL_timestep
eval_SCOPSOWL_Uptake, 340	scopsowl.h, 333
initial_guess_SCOPSOWL, 340	set_SKUA_ICs
SCOPSOWL_OPT_set_y, 340	skua.h, 372
SCOPSOWL_OPTIMIZE, 340	set_SKUA_params
Set_ActivationEnergy	skua.h, 372
UnsteadyReaction, 241	set_SKUA_timestep
Set_Affinity	skua.h, 372
UnsteadyReaction, 241	Set_Species_Index
Set Area	UnsteadyReaction, 239
MassBalance, 105	Set_SteadyState
set_DOGFISH_ICs	MassBalance, 105
dogfish.h, 259	Set_Stoichiometric
set_DOGFISH_params	Reaction, 176
dogfish.h, 260	UnsteadyReaction, 240
set_DOGFISH_timestep	Set TimeStep
dogfish.h, 260	MassBalance, 105
Set Delta	UnsteadyReaction, 242
MassBalance, 104	Set_TotalConcentration
Set_Energy	MassBalance, 104
Reaction, 177	Set_Type
UnsteadyReaction, 240	MassBalance, 105
Set_Enthalpy Reaction 177	Set_Volume MassBalance 105
Reaction, 177	MassBalance, 105
UnsteadyReaction, 240	Set_ZeroInitialSolids
Set_EnthalpyANDEntropy	MassBalance, 105
Reaction, 177	set_alkalinity

MasterSpeciesList, 110	UnsteadyAdsorption, 230
set_list_size	setChargeDensity
MasterSpeciesList, 110	AdsorptionReaction, 15
set_size	ChemisorptionReaction, 43
Matrix, 115	MultiligandAdsorption, 145
set_species	MultiligandChemisorption, 154
MasterSpeciesList, 110	UnsteadyAdsorption, 230
set_variables	setChargeDensityValue
egret.h, 265	AdsorptionReaction, 14
setActivities	ChemisorptionReaction, 43
AdsorptionReaction, 14	setDelta
ChemisorptionReaction, 43	ChemisorptionReaction, 42
setActivityEnum	setDeltas
AdsorptionReaction, 13	ChemisorptionReaction, 42
ChemisorptionReaction, 42	MultiligandChemisorption, 153
MultiligandAdsorption, 143	setElectricPotential
MultiligandChemisorption, 153	MultiligandAdsorption, 144
UnsteadyAdsorption, 229	MultiligandChemisorption, 154
setActivityModelInfo	setFormula
AdsorptionReaction, 13	Molecule, 127
ChemisorptionReaction, 42	setInputFile
MultiligandAdsorption, 143	yaml_cpp_class, 251
MultiligandChemisorption, 153	setlonicStrength
UnsteadyAdsorption, 229	AdsorptionReaction, 15
setAdsorbIndices	ChemisorptionReaction, 43
AdsorptionReaction, 13	MultiligandAdsorption, 145
ChemisorptionReaction, 41	MultiligandChemisorption, 154
MultiligandAdsorption, 143	UnsteadyAdsorption, 230
MultiligandChemisorption, 152	setIonicStrengthValue
UnsteadyAdsorption, 228	AdsorptionReaction, 14
setAdsorbentName	ChemisorptionReaction, 43
AdsorptionReaction, 14	setLigandIndex
ChemisorptionReaction, 43	ChemisorptionReaction, 42
MultiligandAdsorption, 144	setLigandIndices
MultiligandChemisorption, 153	MultiligandChemisorption, 152
UnsteadyAdsorption, 230	setLigandName
setAlias	MultiligandAdsorption, 144
Document, 52	MultiligandChemisorption, 153
Header, 92	setMolarArea
SubHeader, 215, 216	Molecule, 128
setAqueousIndex	setMolarFactor
AdsorptionReaction, 13	AdsorptionReaction, 13
UnsteadyAdsorption, 229	MultiligandAdsorption, 143
	-
setAqueousIndexAuto	UnsteadyAdsorption, 229
AdsorptionReaction, 13	setMolarVolume
MultiligandAdsorption, 143	Molecule, 128
UnsteadyAdsorption, 229	setMolarWeigth
setAreaBasisBool	Molecule, 128
AdsorptionReaction, 14	setName
UnsteadyAdsorption, 230	Document, 52
setAreaFactor	Header, 92
AdsorptionReaction, 13	SubHeader, 215
ChemisorptionReaction, 42	setNameAliasPair
MultiligandAdsorption, 144	Document, 52
MultiligandChemisorption, 153	Header, 92
UnsteadyAdsorption, 229	SubHeader, 216
setBasis	setSpecificArea
AdsorptionReaction, 14	AdsorptionReaction, 14
· · · · · · · · · · · · · · · · · · ·	

ChemisorptionReaction, 42	setup_MONKFISH_DATA
MultiligandAdsorption, 144	monkfish.h, 324
MultiligandChemisorption, 153	setup_SCOPSOWL_DATA
UnsteadyAdsorption, 229	scopsowl.h, 332
setSpecificMolality	setup SHARK DATA
AdsorptionReaction, 14	shark.h, 359
ChemisorptionReaction, 42	setup_SKUA_DATA
MultiligandAdsorption, 144	skua.h, 371
MultiligandChemisorption, 153	shapeFactor
UnsteadyAdsorption, 229	magpie.h, 306
setState	shark
Document, 52	ui.h, 386
Header, 92	shark.h, 343
SubHeader, 216	AbsPerm, 349
setSurfaceCharge	act_choice, 355
AdsorptionReaction, 14	AreaSTD, 348
MultiligandAdsorption, 144	BATCH, 349
UnsteadyAdsorption, 229	CSTR, 349
setSurfaceChargeBool	calculate_ionic_strength, 350
AdsorptionReaction, 14	Convert2Concentration, 357
ChemisorptionReaction, 43	Convert2LogConcentration, 357
MultiligandAdsorption, 144	CoordSTD, 348
MultiligandChemisorption, 154	DAVIES, 350
UnsteadyAdsorption, 230	DEBYE_HUCKEL, 350
setTotalMass	Davies_equation, 354
AdsorptionReaction, 14	DebyeHuckel_equation, 354
ChemisorptionReaction, 42	e, 348
MultiligandAdsorption, 144	FLORY_HUGGINS, 350
MultiligandChemisorption, 153	Faraday, 348
UnsteadyAdsorption, 230	FloryHuggins, 351
setTotalVolume	FloryHuggins_chemi, 352
AdsorptionReaction, 14	FloryHuggins_multichemi, 352
ChemisorptionReaction, 43	FloryHuggins_multiligand, 351
MultiligandAdsorption, 144	FloryHuggins_unsteady, 351
MultiligandChemisorption, 154	IDEAL_ADS, 350
UnsteadyAdsorption, 230	IDEAL, 350
setVolumeFactor	ideal_solution, 354
AdsorptionReaction, 13	kB, 348
ChemisorptionReaction, 42	LengthFactor, 348
MultiligandAdsorption, 144	linearsolve_choice, 355
MultiligandChemisorption, 153	linesearch_choice, 355
UnsteadyAdsorption, 229	Na, 348
setbcs	PFR, 349
FINCH DATA, 70	PITZER, 350
setic	print2file_shark_header, 350
FINCH DATA, 69	print2file_shark_info, 350
setparams	print2file shark results new, 350
FINCH_DATA, 70	print2file_shark_results_old, 350
setpostprocess	reactor_choice, 355
FINCH DATA, 70	read adsorbobjects, 358
setpreprocess	read_chemisorbobjects, 359
FINCH DATA, 70	read_equilrxn, 358
settime	read_massbalance, 358
FINCH DATA, 70	read multichemi scenario, 357
setup DOGFISH DATA	read_multichemiobjects, 359
dogfish.h, 259	read_multiligand_scenario, 357
setup_FINCH_DATA	read_multiligandobjects, 358
finch.h, 273	read_nutiligandobjects, 356
IIIIGII.II, 273	reau_options, 557

read_scenario, 357	shark_reset
read_species, 358	shark.h, <mark>361</mark>
read_unsteadyadsorbobjects, 358	shark_residual
read_unsteadyrxn, 358	shark.h, 362
Rstd, 348	shark_solver
SHARK_DATA, 349	shark.h, 361
SHARK_SCENARIO, 362	shark_temperature_calculations
SHARK TESTS OLD, 366	shark.h, 360
SHARK TESTS, 366	shark timestep adapt
SHARK, 362	shark.h, 361
SIT, 350	shark_timestep_const
setup_SHARK_DATA, 359	shark.h, 361
shark_add_customResidual, 360	sigma
shark_energy_calculations, 360	CGS DATA, 36
shark_executioner, 361	sigma_m
shark guess, 360	TRAJECTORY_DATA, 222
shark_initial_conditions, 360	sigma_n
shark pH finder, 360	TRAJECTORY_DATA, 222
shark_parameter_check, 360	
	sigma_v
shark_postprocesses, 361	TRAJECTORY_DATA, 222
shark_preprocesses, 361	sigma_vz
shark_reset, 361	TRAJECTORY_DATA, 222
shark_residual, 362	sigma_z
shark_solver, 361	TRAJECTORY_DATA, 222
shark_temperature_calculations, 360	sim_time
shark_timestep_adapt, 361	SCOPSOWL_DATA, 181
shark_timestep_const, 361	SKUA_DATA, 205
surf_act_choice, 355	simple_darken_Dc
UNIQUAC_ACT, 350	skua.h, 370
UNIQUAC_chemi, 353	simulation_equil
UNIQUAC_multichemi, 353	SCOPSOWL_OPT_DATA, 187
UNIQUAC_multiligand, 353	SKUA_OPT_DATA, 210
UNIQUAC_unsteady, 352	simulation_fail
UNIQUAC, 352	error.h, 267
VacuumPermittivity, 349	simulationtime
valid_act, 349	SHARK_DATA, 199
valid_mb, 349	single_fiber_density
valid_surf_act, 350	MONKFISH_DATA, 134
VolumeSTD, 348	singular_matrix
WaterRelPerm, 349	error.h, 268
shark_add_customResidual	size
shark.h, 360	Document, 52
shark_energy_calculations	Header, 92
shark.h, 360	KeyValueMap, 97
shark executioner	MasterSpeciesList, 112
shark.h, 361	YamlWrapper, 255
shark_guess	skua
shark.h, 360	ui.h, 386
shark_initial_conditions	skua.h, 367
shark.h, 360	const Dc, 370
shark_pH_finder	const kf, 371
shark.h, 360	D_c, 369
shark_parameter_check	D_inf, 369
shark, nestrocosses	D_o, 369
shark_postprocesses	default_Dc, 369
shark.h, 361	default_kf, 370
shark_preprocesses	empirical_kf, 371
shark.h, 361	molefractionCheck, 371

print2file_SKUA_header, 369	MultiligandAdsorption, 148
print2file_SKUA_results_new, 369	MultiligandChemisorption, 157
print2file_SKUA_results_old, 369	specific_heat
print2file_SKUA_time_header, 369	PURE_GAS, 172
print2file species header, 369	specific_molality
SKUA Executioner, 372	AdsorptionReaction, 20
SKUA_HPP_, 369	SphereArea
SKUA SCENARIOS, 373	mola.h, 320
SKUA_TESTS, 375	SphereVolume
SKUA postprocesses, 372	mola.h, 320
SKUA_preprocesses, 372	Spherical
SKUA reset, 373	finch.h, 272
_ :	
SKUA, 373	sphericalAvg
set_SKUA_ICs, 372	Matrix, 118
set_SKUA_params, 372	sphericalBCFill
set_SKUA_timestep, 372	Matrix, 118
setup_SKUA_DATA, 371	ss_ads_names
simple_darken_Dc, 370	SHARK_DATA, 198
theoretical_darken_Dc, 370	ss_chem_names
skua_dat	SHARK_DATA, 198
SCOPSOWL_DATA, 184	ssmulti_names
SKUA_OPT_DATA, 211	SHARK_DATA, 198
skua_opt	ssmultichem_names
ui.h, 386	SHARK DATA, 198
skua_opt.h, 375	start temp
eval_SKUA_Uptake, 377	SHARK DATA, 200
initial_guess_SKUA, 377	state
SKUA OPT set y, 377	SubHeader, 217
SKUA OPTIMIZE, 377	SteadyState
Sn	FINCH DATA, 65
FINCH DATA, 69	MassBalance, 108
Snp1	steadystate
FINCH_DATA, 69	SHARK DATA, 201
	- · · ·
SolnTransform	steps
Matrix, 118	GMRESLP_DATA, 75
solve	Stoichiometric
FINCH_DATA, 70	Reaction, 178
sorbed_molefraction	string_parse_error
DOGFISH_PARAM, 58	error.h, 268
MONKFISH_PARAM, 137	Sub_Map
sorption_bc	Header, 93
MONKFISH_PARAM, 138	SubHeader, 213
SpeciationCurve	\sim SubHeader, 214
SHARK_DATA, 201	addPair, 215
species	alias, 216
DOGFISH_PARAM, 58	clear, 215
MONKFISH PARAM, 138	Data_Map, 216
MasterSpeciesList, 112	DisplayContents, 216
species_dat	getAlias, 216
MIXED_GAS, 123	getMap, 215
species_index	getName, 216
UnsteadyReaction, 246	getState, 216
speciesName	isAlias, 216
MasterSpeciesList, 111	isAnchor, 216
SCOPSOWL PARAM DATA, 192	
:	name, 216
SKUA_PARAM, 212	operator=, 215
specific_area	operator[], 215
AdsorptionReaction, 20	setAlias, 215, 216

setName, 215	t_print
setNameAliasPair, 216	DOGFISH_DATA, 55
setState, 216	MONKFISH_DATA, 134
state, 217	SCOPSOWL DATA, 182
SubHeader, 214, 215	SKUA DATA, 205
sum	t_rand
ARNOLDI_DATA, 23	
GMRESRP_DATA, 83	TRAJECTORY_DATA, 222
	TANGENTIAL_FORCE
Matrix, 116	Trajectory.h, 382
Sum_Delta	TEST
MassBalance, 105	ui.h, 386
surf_act_choice	TRAJECTORY_DATA, 219
shark.h, 355	a, <mark>22</mark> 1
SurfDiff	A_separator, 221
SCOPSOWL_DATA, 182	A_wire, 221
surface_activity	b, 221
AdsorptionReaction, 19	B0, 221
MultiligandAdsorption, 147	beta, 222
MultiligandChemisorption, 157	Cap, 222
surface_charge	chi_p, 221
AdsorptionReaction, 20	dt, 222
surface_concentration	dX, 222
DOGFISH_PARAM, 58	
Sutherland_Const	dY, 222
PURE GAS, 172	eta, 221
Sutherland_Temp	H, 222
PURE GAS, 172	H0, 221
Sutherland_Viscosity	Hamaker, 221
_ •	k, 221
PURE_GAS, 172	L, 221
Symbol	L_wire, 221
Atom, 29	M, 222
Sys	m_rand, 222
SYSTEM_DATA, 219	mp, 222
sys_dat	Ms, 221
MAGPIE_DATA, 101	mu_0, 221
т	n_rand, 222
T FINCH DATA 62	POL, 222
FINCH_DATA, 63	porosity, 221
SYSTEM_DATA, 218	q_bar, 222
t	—
BiCGSTAB_DATA, 34	Q_in, 221
FINCH_DATA, 63	rho_f, 221
SCOPSOWL_DATA, 181	rho_p, 221
SCOPSOWL_OPT_DATA, 188	Rs, 221
SKUA_DATA, 205	s_rand, <mark>222</mark>
SKUA_OPT_DATA, 211	sigma_m, 222
t_count	sigma_n, <mark>222</mark>
SHARK_DATA, 199	sigma_v, <mark>222</mark>
t counter	sigma_vz, 222
DOGFISH_DATA, 55	sigma_z, <mark>222</mark>
MONKFISH_DATA, 134	t rand, 222
SCOPSOWL_DATA, 182	Temp, 221
SKUA_DATA, 205	V0, <mark>22</mark> 1
t old	V_separator, 221
FINCH DATA, 63	V_soparator, 221
SCOPSOWL_DATA, 181	V_wire, 221 Vr, 222
SKUA_DATA, 181	Vt, 222
t_out	X, 222
SHARK_DATA, 199	Y, 222

Y_initial, 222	GCR_DATA, 73
Table	GMRESLP_DATA, 75
PeriodicTable, 164	GMRESRP_DATA, 81
Temp	PCG_DATA, 161
TRAJECTORY_DATA, 221	PICARD_DATA, 165
temp_step	total_density
SHARK_DATA, 200	MIXED_GAS, 123
temperature	total_dyn_vis
SHARK_DATA, 200	MIXED_GAS, 123
temperature_affinity	total_eval
UnsteadyReaction, 246	GSTA_OPT_DATA, 87
TemperatureCurve	SCOPSOWL_OPT_DATA, 186
SHARK_DATA, 201	SKUA_OPT_DATA, 209
tempy	SYSTEM_DATA, 218
SCOPSOWL_DATA, 183	total_iter
tensor_out_of_bounds	FINCH_DATA, 66
error.h, 268	GCR_DATA, 72
term_precon	GMRESR_DATA, 78
GMRESR_DATA, 79	KMS_DATA, 100
KMS_DATA, 101	total_mass
terminal_precon	AdsorptionReaction, 20
GMRESR_DATA, 79	MultiligandAdsorption, 148
KMS_DATA, 100	MultiligandChemisorption, 158
test	total_molecular_weight
ui.h, 387	MIXED_GAS, 123
test_loop	total_pressure
ui.h, 390	MIXED_GAS, 122
theoretical_darken_Dc	SCOPSOWL_DATA, 182
skua.h, 370	total_sorption
time	DOGFISH_DATA, 56
DOGFISH_DATA, 55	MONKFISH_DATA, 134
MONKFISH_DATA, 133	total_sorption_old
SHARK_DATA, 199	DOGFISH_DATA, 56
time_old	MONKFISH_DATA, 134
DOGFISH_DATA, 55	total_specific_heat
MONKFISH_DATA, 133	MIXED_GAS, 123
SHARK_DATA, 199	total_steps
time_step	DOGFISH_DATA, 55
UnsteadyReaction, 246	MONKFISH_DATA, 133
TimeAdaptivity	SCOPSOWL_DATA, 181
SHARK_DATA, 201	SKUA_DATA, 205
timesteps	total_volume
SHARK_DATA, 199	AdsorptionReaction, 20
token_parser	MultiligandAdsorption, 148
yaml_cpp_class, 252	MultiligandChemisorption, 158
tol_abs	TotalConcentration
BiCGSTAB_DATA, 33	MassBalance, 107
CGS_DATA, 36	totalcalls
FINCH_DATA, 66	SHARK_DATA, 198
GCR_DATA, 73	totalsteps
GMRESLP_DATA, 75	SHARK_DATA, 198
GMRESRP_DATA, 81	trajectory
PCG_DATA, 161	ui.h, 386
PICARD_DATA, 165	Trajectory.h, 380
tol_rel	Brown_RAD, 382
BiCGSTAB_DATA, 33	Brown_THETA, 382
CGS DATA, 36	CARTESIAN, 382
FINCH_DATA, 65	Capture_Force, 382
<i>-</i>	• = /

DISPLACEMENT, 383	BasicUI, 224
Grav_R, 382	count, 224
Grav_T, 382	Files, 224
In_P_Velocity, 382	input_files, 224
In_PVel_Rad, 382	max, 224
In_PVel_Theta, 382	MissingArg, 224
LOCATION, 383	option, 224
Magnetic R, 382	Path, 224
Magnetic T, 382	path, 224
Number Generator, 383	user_input, 223
P Velocity, 382	value_type, 223
POLAR, 382	UI HPP
PVel_Rad, 382	ui.h, 385
PVel Theta, 382	ulC
RADIAL FORCE, 382	
Removal_Efficiency, 383	FINCH_DATA, 63
Run Trajectory, 383	UNIQUAC_ACT
TANGENTIAL FORCE, 382	shark.h, 350
Trajectory_SetupConstants, 383	UNIQUAC_chemi
V RAD, 382	shark.h, 353
V_HAD, 302 V_THETA, 382	UNIQUAC_multichemi
Van_R, 382	shark.h, 353
Trajectory_SetupConstants	UNIQUAC_multiligand
	shark.h, 353
Trajectory.h, 383	UNIQUAC_unsteady
transpose	shark.h, <mark>352</mark>
Matrix, 117	UNIQUAC
transpose_dat	shark.h, 352
GCR_DATA, 74	UNKNOWN
transpose_multiply	yaml_wrapper.h, 395
Matrix, 117	uT_old
tridiagonalFill	FINCH DATA, 63
Matrix, 117	uTotal
tridiagonalSolve	finch.h, 272
Matrix, 117	ubest
tridiagonalVectorFill	FINCH DATA, 68
Matrix, 119	ui.h, 383
twoFifths	aui_help, 386
gsta_opt.h, 279	bui_help, 386
Туре	CONTINUE, 386
MassBalance, 107	display_help, 389
type	display_reip, 303 display_version, 389
ValueTypePair, 249	dogfish, 386
	ECO EXECUTABLE, 385
U	ECO VERSION, 385
CGS_DATA, 37	-
GCR_DATA, 74	EXECUTE, 386
u_star	EXIT, 386
FINCH_DATA, 68	eel, 386
u_temp	egret, 386
GCR_DATA, 74	exec, 387
uAverage	exec_loop, 391
finch.h, 272	exit, 386
uAvg	finch, 386
FINCH_DATA, 63	gsta_opt, 386
uAvg_old	HELP, 386
FINCH_DATA, 63	help, 387
UI_DATA, 223	input, 388
argc, 224	invalid_input, 389
argv, 224	lark, 386

macaw, 386	Explicit_Eval, 233
magpie, 386	getActivity, 234
mola, 386	getActivityEnum, 235
monkfish, 386	getAdsorbIndex, 235
number_files, 388	getAdsorbentName, 235
path, 387	getAqueousIndex, 235
run_exec, 391	getAreaFactor, 234
run_executable, 391	getBulkDensity, 234
run_test, 391	getChargeDensity, 234
sandbox, 386	getIonicStrength, 235
scops_opt, 386	getMolarFactor, 234
scopsowl, 386	getNumberRxns, 235
shark, 386	getOldActivity, 234
skua, 386	getReaction, 233
skua_opt, 386	getSpecificArea, 234
TEST, 386	getSpecificMolality, 234
test, 387	getSurfaceCharge, 234
test loop, 390	getTotalMass, 234
trajectory, 386	getTotalVolume, 234
UI HPP , 385	getVolumeFactor, 234
valid_addon_options, 389	includeSurfaceCharge, 235
valid exec string, 388	Initialize_Object, 228
valid_input_execute, 390	isAreaBasis, 235
valid_input_main, 390	modifyDeltas, 228
valid_input_tests, 390	setActivityEnum, 229
valid_input_tests, 390 valid_options, 386	setActivityLitarii, 229
_ ·	
valid_test_string, 388	setAdsorbIndices, 228
version, 387	setAdsorbentName, 230
un	setAqueousIndex, 229
FINCH_DATA, 68	setAqueousIndexAuto, 229
unm1	setAreaBasisBool, 230
FINCH_DATA, 68	setAreaFactor, 229
unp1	setBasis, 230
FINCH_DATA, 68	setChargeDensity, 230
unregistered_name	setlonicStrength, 230
error.h, 268	setMolarFactor, 229
unstable_matrix	setSpecificArea, 229
error.h, 268	setSpecificMolality, 229
UnsteadyAdsList	setSurfaceCharge, 229
SHARK_DATA, 196	setSurfaceChargeBool, 230
UnsteadyAdsorption, 225	setTotalMass, 230
\sim UnsteadyAdsorption, 228	setTotalVolume, 230
activities_old, 235	setVolumeFactor, 229
ads_rxn, 235	UnsteadyAdsorption, 228
calculateActiveFraction, 231	updateActivities, 230
calculateAqueousChargeExchange, 231	UnsteadyList
calculateAreaFactors, 230	SHARK_DATA, 196
calculateEquilibria, 230	UnsteadyReaction, 236
calculateEquilibriumCorrection, 232	\sim UnsteadyReaction, 239
calculatePsi, 231	activation_energy, 246
calculateRates, 230	calculateEnergies, 242
calculateSurfaceChargeDensity, 231	calculateEquilibrium, 242
callSurfaceActivity, 231	calculateRate, 242
checkAqueousIndices, 228	checkSpeciesEnergies, 242
Display_Info, 228	Display_Info, 239
Eval_IC_Residual, 233	Eval_IC_Residual, 245
Eval_ReactionRate, 233	Eval ReactionRate, 244
Eval_Residual, 232	Eval_Residual, 244, 245
·· <u> </u>	, , ,

	Explicit_Eval, 245	lark.h, 288
	forward_rate, 245	updateActivities
	forward_ref_rate, 246	UnsteadyAdsorption, 230
	Get_ActivationEnergy, 244	upperHessenberg2Triangular
	Get_Affinity, 244	Matrix, 119
	Get_Energy, 243	upperHessenbergSolve
	Get_Enthalpy, 243	Matrix, 120
	Get_Entropy, 243	upperTriangularSolve
	Get_Equilibrium, 243	Matrix, 119
	Get_Forward, 243	us_ads_names
	Get_ForwardRef, 244	SHARK_DATA, 198
	Get_InitialValue, 243	user_data
	Get_MaximumValue, 243	DOGFISH_DATA, 56
	Get_Reverse, 244	MONKFISH_DATA, 136
	Get_ReverseRef, 244	SCOPSOWL_DATA, 184
	Get_Species_Index, 243	SKUA_DATA, 207
	Get_Stoichiometric, 243	user_input
	Get_TimeStep, 244	UI_DATA, 223
	haveEquilibrium, 242	uT
	HaveForRef, 246	FINCH_DATA, 63
	HaveForward, 246	uz_l_E
	haveForward, 243	FINCH_DATA, 67
	haveForwardRef, 242	uz_l_l
	haveRate, 242	FINCH_DATA, 67
	HaveRevRef, 246	uz_lm1_E
	HaveReverse, 246	FINCH_DATA, 68
	haveReverse, 243	uz_lm1_l
	haveReverseRef, 243	FINCH_DATA, 67
	initial_value, 245	uz_lp1_E
	Initialize_Object, 239	FINCH_DATA, 68
	max_value, 245	uz_lp1_l
	reverse_rate, 246	FINCH_DATA, 67
	reverse_ref_rate, 246	V
	Set_ActivationEnergy, 241	magpie.h, 306
	Set_Affinity, 241	V
	Set_Energy, 240	ARNOLDI_DATA, 23
	Set_Enthalpy, 240	BiCGSTAB_DATA, 34
	Set_EnthalpyANDEntropy, 240	CGS DATA, 37
	Set_Entropy, 240	GMRESRP_DATA, 83
	Set_Equilibrium, 240	mSPD_DATA, 139
	Set_Forward, 241	PJFNK DATA, 170
	Set_ForwardRef, 241	V0
	Set_InitialValue, 240	TRAJECTORY DATA, 221
	Set MaximumValue, 240	V RAD
	Set Reverse, 241	Trajectory.h, 382
	Set ReverseRef, 241	V THETA
	Set Species Index, 239	Trajectory.h, 382
	Set Stoichiometric, 240	V_separator
	Set_TimeStep, 242	TRAJECTORY_DATA, 221
	species index, 246	V_wire
	temperature_affinity, 246	TRAJECTORY_DATA, 221
	time_step, 246	vIC
	UnsteadyReaction, 239	FINCH DATA, 64
uo	,	VacuumPermittivity
	FINCH DATA, 64	shark.h, 349
Upd	_ · · · ·	valence_e
200	FINCH DATA, 65	Atom, 28
und	ate_arnoldi_solution	valid_act
apu	a.to_a.rioidi_ooidiloii	·

shark.h, 349	vnp1
valid_addon_options	FINCH_DATA, 68
ui.h, 389	VO
valid_exec_string	FINCH_DATA, 64
ui.h, 388	volume
valid_input_execute	MassBalance, 107
ui.h, 390	SHARK_DATA, 200
valid_input_main	volume_factors
ui.h, 390	AdsorptionReaction, 20
valid_input_tests	VolumeSTD
ui.h, 390	shark.h, 348
valid_mb	Vr
shark.h, 349	TRAJECTORY_DATA, 222
valid_options	Vt
ui.h, 386	TRAJECTORY_DATA, 222
valid_phase	
mola.h, 320	W
valid_surf_act	ARNOLDI_DATA, 23
shark.h, <u>350</u>	CGS_DATA, 37
valid test string	GMRESRP_DATA, 83
ui.h, 388	WaterRelPerm
Value_Type	shark.h, 349
ValueTypePair, 249	weightedAvg
value_type	gsta_opt.h, 280
UI_DATA, 223	
ValueTypePair, 247	Χ
~ValueTypePair, 248	TRAJECTORY_DATA, 222
assertType, 248	X
DisplayPair, 249	BiCGSTAB_DATA, 33
editPair, 248	CGS_DATA, 37
editValue, 248	GCR_DATA, 73
findType, 248	GMRESLP_DATA, 76
getBool, 249	GMRESRP_DATA, 82
getDouble, 249	GPAST_DATA, 84
getInt, 249	PCG_DATA, 162
getPair, 249	PJFNK_DATA, 170
getString, 249	QR_DATA, 174
getType, 249	x0
- · · ·	PICARD_DATA, 166
getValue, 249	X_new
operator=, 248	SHARK_DATA, 202
type, 249	X_old
Value_Type, 249	SHARK_DATA, 202
ValueTypePair, 248	xIC
Van_R	SCOPSOWL_PARAM_DATA, 190
Trajectory.h, 382	SKUA_PARAM, 212
vanAlbada_discretization	xk
finch.h, 275	BACKTRACK_DATA, 30
vector_out_of_bounds	xn
error.h, 268	SKUA_PARAM, 212
velocity	xnp1
MIXED_GAS, 123	SKUA_PARAM, 212
version	xsec_area
ui.h, 387	MassBalance, 107
Vk	SHARK_DATA, 200
ARNOLDI_DATA, 23	
GMRESRP_DATA, 82	Υ
vn	TRAJECTORY_DATA, 222
FINCH_DATA, 68	у

BiCGSTAB_DATA, 34		Doc_Map, 256
GMRESRP_DATA, 82		end, 254
GPAST_DATA, 84		getAnchoredDoc, 255
SCOPSOWL_DATA, 183		getDocFromHeadAlias, 255
SKUA_DATA, 206		getDocFromSubAlias, 255
y base		getDocMap, 254
SCOPSOWL OPT DATA, 188		getDocument, 254
SKUA_OPT_DATA, 211		operator(), 254
y_eff		operator=, 254
SKUA PARAM, 212		resetKeys, 255
Y initial		revalidateAllKeys, 255
_		size, 255
TRAJECTORY_DATA, 222		YamlWrapper, 254
YAML_CPP_TEST	νk	ramitvrapper, 254
yaml_wrapper.h, 396	yk	ADNOLDI DATA 22
YAML_WRAPPER_TESTS		ARNOLDI_DATA, 23
yaml_wrapper.h, 396	Z	
yaml_cpp_class, 250	_	
\sim yaml_cpp_class, 251		magpie.h, 305
cleanup, 251	Z	DIGGOTAD DATA OF
current_token, 252		BiCGSTAB_DATA, 34
DisplayContents, 251		CGS_DATA, 38
executeYamlRead, 251		PCG_DATA, 162
file_name, 251	Z_0	d
getYamlWrapper, 251		PCG_DATA, 162
input_file, 251	zero	_surf_diffusion
• —		scopsowl.h, 331
previous_token, 252	zero	_vector
readInputFile, 251		error.h, 268
setInputFile, 251	Zero	olnitialSolids
token_parser, 252		MassBalance, 108
yaml_cpp_class, 251		SHARK DATA, 201
yaml_wrapper, 251	zero	<u> </u>
yaml_object	2610	Matrix, 115
SHARK_DATA, 203	Zk	Matrix, 113
yaml_wrapper	۷ĸ	CMDECDD DATA 00
yaml_cpp_class, 251		GMRESRP_DATA, 82
yaml_wrapper.h, 391		
ALIAS, 396		
ANCHOR, 396		
allLower, 396		
BOOLEAN, 395		
DOUBLE, 395		
data_type, 395		
header_state, 395		
INT, 395		
isEven, 396		
NONE, 396		
•		
NONE, 396		
NONE, 396 STRING, 395		
NONE, 396 STRING, 395 UNKNOWN, 395		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396 YAML_WRAPPER_TESTS, 396		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396 YAML_WRAPPER_TESTS, 396 YamlWrapper, 252 ~YamlWrapper, 254		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396 YAML_WRAPPER_TESTS, 396 YamlWrapper, 252 ~YamlWrapper, 254 addDocKey, 255		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396 YAML_WRAPPER_TESTS, 396 YamlWrapper, 252 ~YamlWrapper, 254 addDocKey, 255 begin, 254, 255		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396 YAML_WRAPPER_TESTS, 396 YamlWrapper, 252 ~YamlWrapper, 254 addDocKey, 255 begin, 254, 255 changeKey, 255		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396 YAML_WRAPPER_TESTS, 396 YamlWrapper, 252 ~YamlWrapper, 254 addDocKey, 255 begin, 254, 255 changeKey, 255 clear, 255		
NONE, 396 STRING, 395 UNKNOWN, 395 YAML_CPP_TEST, 396 YAML_WRAPPER_TESTS, 396 YamlWrapper, 252 ~YamlWrapper, 254 addDocKey, 255 begin, 254, 255 changeKey, 255		