**GW** Analysis Tools

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# **Gravitational Waves Analysis Tools**

A suite of analysis tools useful for gravitational wave science. All code is written in C++, with some of the interface classes wrapped in Cython to allow for python-access.

### 1.1 Compatibility

Known to work with gcc/g++-7

Known to work with gcc/g++-9

Need nvcc - known to work with v9.1 of CUDA

### 1.2 Required Software

Required non-standard C libraries: FFTW3 ADOL-C GSL CUDA

Required non-standard Python packages: Cython

Required non-standard packages for documentation: Doxygen

### 1.3 Current Development

NOTE: currently using static parameters to share data between threads for mcmc\_gw.cpp. This could cause issues when running multiple samplers at the same time. Investigating further.

To do:

Change MCMC\_MH to use the more general threadPool class instead of a custom threadpool, incorporate job class and comparator

### 1.4 Installation

For proper compilation, update or create the enviornment variables CPATH, LIBRARY\_PATH, and LD\_LIBRARY — \_PATH, which should point to header files and lib files, respectively. Specifically, these variables should point to the above libraries.

Also, the PYTHONPATH environment variables must point to /gw\_analysis\_tools\_py/src because I can't figure how to get this shit to work.

In the root directory of the project, run 'make' to compile source files, create the library file and create the cython modules, and create the documentation.

To just create C++/C files, run 'make c'.

Run 'make test' to build a test program that will create an executable.

### 1.5 Supported Functionality

#### 1.5.1 Waveform Generation

IMRPhenomD, IMRPhenomPv2

### 1.5.2 Modified Gravity

ppE\_IMRPhenomD\_Inspiral ppE\_IMRPhenomDv2\_IMR ppE\_IMRPhenomPv2\_IMRPhenomPv2\_IMR

### 1.5.3 Fisher Analysis

utilizes the above waveform templates

### 1.5.4 MCMC Routines

Has a generic MCMC sampler, MCMC\_MH, that utilizes gaussian steps, differential evolution steps, and Fisher informed steps. Includes wrapping MCMC\_MH\_GW for GW specific sampling, currently only for one detector.

Includes log likelihood caclulation for implementation in other samplers.

### 1.6 Usage

### 1.6.1 Environment variables

The environment variable PYTHONPATH should include the directory \$(PROJECT\_DIR)

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#### 1.6.2 Include

To include header files, use -I\$(PROJECT\_DIRECTORY)/include

#### 1.6.3 Link

To link object files, use -L\$(PROJECT\_DIRECTORY)/lib -lgwat (the -L command is un-needed if you add /lib to the environment variable CPATH)

For dynamic linking, the following environment variables for Linux (MacOs) should be updated to include /lib - LD LIBRARY PATH (DYLD LIBRARY PATH)

For Cuda code: use -lcuda -lcudart

For Cuda, may need to link to /usr/local/cuda/lib64/ (or wherever this library is on your machine)

### 1.6.4 Python Importable Code

Two modules currently available:

1.6.4.1 gw\_analysis\_tools\_py.mcmc\_routines\_ext

mcmc\_routines\_ext.pyx wraps the log\_likelihood functions in mcmc\_routines.cpp

1.6.4.2 gw\_analysis\_tools\_py.waveform\_generator\_ext

waveform\_generator\_ext.pyx wraps the fourier\_waveform function in waveform\_generator.cpp

Also contains the SNR calculation function

### 1.6.4.3 Custom Waveforms

If adding waveforms and to have full accesibility:

Create class, using other waveforms as template – need interface to create full waveform (plus,cross polarization), and amplitude/phase

Add the option as a waveform to waveform\_generation.cpp, including the header file at the top of the waveform\_← generation.cpp file

For autodiff Fishers – write the class as a template with double and adouble types for all variables. Then write the necessary fisher subroutines (see fisher file to determine whats necessary)

For numerical Fishers - write finite difference method, following the template of the previous waveforms

For MCMC sampling – write mcmc\_fisher\_wrapper and mcmc\_likelihood\_wrapper options and write any necessary initialization in MCMC MH GW

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# gw\_analysis\_tools

A suite of tools useful for doing statistical studies on gravitational wave science, including routines useful in  $MC \leftarrow MC$  studies, wave template generation, Fisher analysis, etc. Written in C++ and wrapped in Cython for access in Python.

6 gw\_analysis\_tools

# Namespace Index

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3.1	Namespace	LIST
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Here is a list of all documer	nted namespaces v	with	brie	f descript	ions
-------------------------------	-------------------	------	------	------------	------

waveform_generator_ext	
Python wrapper for the waveform generation in waveform, generator cpp	15

8 Namespace Index

# **Hierarchical Index**

## 4.1 Class Hierarchy

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

$alpha\_coeffs < T > \dots \dots$
Comparator
comparator_ac_fft
comparator_ac_serial
Comparatorswap
default_comp< jobtype >
$epsilon\_coeffs < T > \hspace{0.5cm} \ldots \hspace{0.5cm} \ldots \hspace{0.5cm} 28$
fftw_outline
mcmc_routines_ext.fftw_outline_py
gen_params
waveform_generator_ext.gen_params_py
GPUplan
$IMRPhenomD < T > \dots \dots$
IMRPhenomPv2< T >
ppE_IMRPhenomPv2_Inspiral < T >
ppE_IMRPhenomPv2_IMR< T >
ppE IMRPhenomD Inspiral T >
dCS IMRPhenomD <t>19</t>
dCS_IMRPhenomD_log< T >
EdGB_IMRPhenomD <t>24</t>
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# **Class Index**

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# File Index

## 6.1 File List

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

gw_analysis_tools_py/src/mcmc_routines_ext.pyx
File that wraps the code in mcmc_gw.cpp, mcmc_sampler.cpp, mcmc_sampler_internals.cpp,
autocorrelation.cpp
gw_analysis_tools_py/src/waveform_generator_ext.pyx
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include/fisher.h
include/GWATConfig.h
include/IMRPhenomD.h
include/IMRPhenomP.h
include/mcmc_gw.h
include/mcmc_sampler.h
include/mcmc_sampler_internals.h
include/ppE_IMRPhenomD.h
include/ppE_IMRPhenomP.h
include/threadPool.h
include/util.h
include/waveform_generator.h
include/waveform_generator_C.h
include/waveform_util.h
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src/fisher.cpp
src/IMRPhenomD.cpp
src/IMRPhenomP.cpp
src/mcmc_gw.cpp
src/mcmc_sampler.cpp
src/mcmc_sampler_internals.cpp
src/ppE_IMRPhenomD.cpp
src/ppE_IMRPhenomP.cpp
src/util.cpp
src/waveform_generator.cpp
src/waveform_util.cpp

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# **Namespace Documentation**

### 7.1 waveform\_generator\_ext Namespace Reference

Python wrapper for the waveform generation in waveform generator.cpp.

#### Classes

· class gen\_params\_py

Python wrapper for the generation parameters structure, as defined in util.cpp.

#### **Functions**

- def **double** (:1] frequencies, string generation\_method, gen\_params\_py parameters):cdef double[::1] amplitude=np.ascontiguousarray(np.zeros((frequencies.size) double, dtype=np.float64, frequencies, frequencies, size, amplitude, generation\_method, parameters, params, :1] frequencies, string generation\_method, gen\_params\_py parameters):cdef double[::1] phase=np.ascontiguousarray(np.zeros((frequencies.size) double, dtype=np.float64, frequencies, frequencies, size, phase, generation\_method, parameters, params, :1] frequencies, string generation\_method, gen\_params\_py parameters):cdef double[::1] waveform\_plus\_\( \limes \) real=np.ascontiguousarray(np.zeros((frequencies.size) double, dtype=np.float64)

### **Variables**

- · complex128\_t
- ndim
- · waveform
- dtype
- **i** = i +1

### 7.1.1 Detailed Description

Python wrapper for the waveform generation in waveform\_generator.cpp.

# **Class Documentation**

## 8.1 alpha\_coeffs < T > Struct Template Reference

### **Public Attributes**

- T coeff1
- T coeff2
- T coeff3
- T coeff4
- T coeff5

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

• include/IMRPhenomP.h

## 8.2 Comparator Class Reference

Class to facilitate the comparing of chains for priority.

### **Public Member Functions**

• bool operator() (int i, int j)

### 8.2.1 Detailed Description

Class to facilitate the comparing of chains for priority.

3 levels of priority: 0 (high) 1 (default) 2 (low)

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

• src/mcmc\_sampler.cpp

18 Class Documentation

## 8.3 comparator\_ac\_fft Class Reference

comparator to sort ac-jobs

```
#include <autocorrelation.h>
```

### **Public Member Functions**

• bool operator() (threaded\_ac\_jobs\_fft t, threaded\_ac\_jobs\_fft k)

### 8.3.1 Detailed Description

comparator to sort ac-jobs

Starts with the longest jobs, then works down the list

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

• include/autocorrelation.h

## 8.4 comparator\_ac\_serial Class Reference

comparator to sort ac-jobs

```
#include <autocorrelation.h>
```

### **Public Member Functions**

• bool operator() (threaded\_ac\_jobs\_serial t, threaded\_ac\_jobs\_serial k)

### 8.4.1 Detailed Description

comparator to sort ac-jobs

Starts with the longest jobs, then works down the list

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

• include/autocorrelation.h

## 8.5 Comparatorswap Class Reference

**Public Member Functions** 

• bool **operator()** (int i, int j)

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

• src/mcmc\_sampler.cpp

## 8.6 dCS\_IMRPhenomD< T> Class Template Reference

Inheritance diagram for dCS\_IMRPhenomD< T >:



Collaboration diagram for dCS\_IMRPhenomD< T >:



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### **Public Member Functions**

virtual int construct\_waveform (T \*frequencies, int length, std::complex < T > \*waveform, source\_parameters < T > \*params)

Constructs the waveform as outlined by.

- virtual T dCS\_phase\_mod (source\_parameters < T > \*param)
- virtual T dCS\_phase\_factor (source\_parameters < T > \*param)
- virtual int construct\_amplitude (T \*frequencies, int length, T \*amplitude, source\_parameters< T > \*params)

  Constructs the Amplitude as outlined by IMRPhenomD.
- virtual int construct\_phase (T \*frequencies, int length, T \*phase, source\_parameters < T > \*params)
   Constructs the Phase as outlined by IMRPhenomD.

### 8.6.1 Member Function Documentation

### 8.6.1.1 construct\_amplitude()

Constructs the Amplitude as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output amplitude, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

#### 8.6.1.2 construct\_phase()

Constructs the Phase as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output phase, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

## 8.6.1.3 construct\_waveform()

Constructs the waveform as outlined by.

arguments: array of frequencies, length of that array, a complex array for the output waveform, and a source\_parameters structure

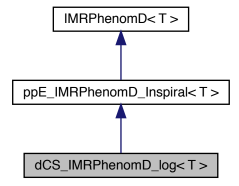
Reimplemented from IMRPhenomD< T >.

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

- include/ppE\_IMRPhenomD.h
- src/ppE\_IMRPhenomD.cpp

# 8.7 dCS\_IMRPhenomD\_log< T > Class Template Reference

Inheritance diagram for dCS\_IMRPhenomD\_log< T >:



Collaboration diagram for dCS\_IMRPhenomD\_log< T >:



#### **Public Member Functions**

virtual int construct\_waveform (T \*frequencies, int length, std::complex < T > \*waveform, source\_parameters < T > \*params)

Constructs the waveform as outlined by.

- virtual T dCS\_phase\_mod (source\_parameters< T > \*param)
- virtual T dCS phase factor (source parameters < T > \*param)
- virtual int construct\_amplitude (T \*frequencies, int length, T \*amplitude, source\_parameters< T > \*params)

  Constructs the Amplitude as outlined by IMRPhenomD.
- virtual int construct\_phase (T \*frequencies, int length, T \*phase, source\_parameters< T > \*params)

  Constructs the Phase as outlined by IMRPhenomD.

#### 8.7.1 Member Function Documentation

# 8.7.1.1 construct\_amplitude()

Constructs the Amplitude as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output amplitude, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

#### 8.7.1.2 construct\_phase()

Constructs the Phase as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output phase, and a source\_parameters structure

Reimplemented from IMRPhenomD < T >.

# 8.7.1.3 construct\_waveform()

Constructs the waveform as outlined by.

arguments: array of frequencies, length of that array, a complex array for the output waveform, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

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

- include/ppE IMRPhenomD.h
- src/ppE IMRPhenomD.cpp

# 8.8 default\_comp < jobtype > Class Template Reference

Default comparator for priority\_queue in threadPool – no comparison.

```
#include <threadPool.h>
```

#### **Public Member Functions**

• bool **operator()** (jobtype j, jobtype k)

# 8.8.1 Detailed Description

template < class jobtype > class default\_comp < jobtype >

Default comparator for priority\_queue in threadPool – no comparison.

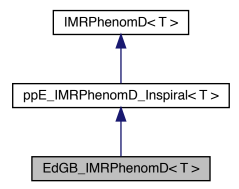
First in first out, not sorting

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

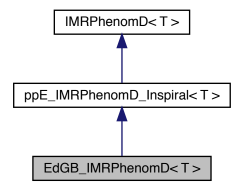
• include/threadPool.h

# 8.9 EdGB\_IMRPhenomD< T> Class Template Reference

Inheritance diagram for EdGB\_IMRPhenomD< T >:



Collaboration diagram for EdGB\_IMRPhenomD< T >:



## **Public Member Functions**

virtual int construct\_waveform (T \*frequencies, int length, std::complex < T > \*waveform, source\_parameters < T > \*params)

Constructs the waveform as outlined by.

- virtual T EdGB\_phase\_mod (source\_parameters< T > \*param)
- virtual T EdGB\_phase\_factor (source\_parameters< T > \*param)
- virtual int construct\_amplitude (T \*frequencies, int length, T \*amplitude, source\_parameters< T > \*params)

  Constructs the Amplitude as outlined by IMRPhenomD.
- virtual int construct\_phase (T \*frequencies, int length, T \*phase, source\_parameters< T > \*params)

  Constructs the Phase as outlined by IMRPhenomD.

## 8.9.1 Member Function Documentation

## 8.9.1.1 construct\_amplitude()

Constructs the Amplitude as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output amplitude, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

#### 8.9.1.2 construct\_phase()

Constructs the Phase as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output phase, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

## 8.9.1.3 construct\_waveform()

Constructs the waveform as outlined by.

arguments: array of frequencies, length of that array, a complex array for the output waveform, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

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

- include/ppE\_IMRPhenomD.h
- src/ppE\_IMRPhenomD.cpp

# 8.10 EdGB\_IMRPhenomD\_log< T > Class Template Reference

Inheritance diagram for EdGB\_IMRPhenomD\_log< T >:



Collaboration diagram for EdGB\_IMRPhenomD\_log< T >:



#### **Public Member Functions**

virtual int construct\_waveform (T \*frequencies, int length, std::complex < T > \*waveform, source\_parameters < T > \*params)

Constructs the waveform as outlined by.

- virtual T EdGB\_phase\_mod (source\_parameters < T > \*param)
- virtual T EdGB phase factor (source parameters< T > \*param)
- virtual int construct\_amplitude (T \*frequencies, int length, T \*amplitude, source\_parameters< T > \*params)

  Constructs the Amplitude as outlined by IMRPhenomD.
- virtual int construct\_phase (T \*frequencies, int length, T \*phase, source\_parameters < T > \*params)
   Constructs the Phase as outlined by IMRPhenomD.

#### 8.10.1 Member Function Documentation

# 8.10.1.1 construct\_amplitude()

Constructs the Amplitude as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output amplitude, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

## 8.10.1.2 construct\_phase()

Constructs the Phase as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output phase, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

## 8.10.1.3 construct\_waveform()

Constructs the waveform as outlined by.

arguments: array of frequencies, length of that array, a complex array for the output waveform, and a source\_parameters structure

Reimplemented from IMRPhenomD< T >.

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

- include/ppE\_IMRPhenomD.h
- src/ppE\_IMRPhenomD.cpp

# 8.11 epsilon\_coeffs < T > Struct Template Reference

**Public Attributes** 

- T coeff1
- T coeff2
- T coeff3
- T coeff4
- T coeff5

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

• include/IMRPhenomP.h

# 8.12 fftw\_outline Struct Reference

## **Public Attributes**

- fftw\_complex \* in
- fftw\_complex \* out
- fftw\_plan p

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

• include/util.h

# 8.13 mcmc\_routines\_ext.fftw\_outline\_py Class Reference

# **Public Member Functions**

- def \_\_init\_\_ (self, N)
- def \_\_reduce\_\_ (self)

# **Public Attributes**

• N

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

• gw\_analysis\_tools\_py/src/mcmc\_routines\_ext.pyx

# 8.14 gen\_params Struct Reference

# **Public Attributes**

- double mass1
- double mass2
- double Luminosity\_Distance
- double spin1 [3]
- double spin2 [3]
- double phic =0
- double tc =0
- int \* bppe
- double \* betappe
- int Nmod
- · double incl\_angle
- · double theta
- · double phi
- double RA
- double DEC
- · double gmst

- double psi =0
- bool NSflag
- double f\_ref =0
- double **phiRef** =0
- double thetaJN = -1
- double alpha0 = 0
- double zeta\_polariz = 0
- double **phi\_aligned** = 0
- double **chil** = 0
- double chip = 0
- bool sky\_average
- gsl\_spline \* **Z\_DL\_spline\_ptr** =NULL
- gsl\_interp\_accel \* **Z\_DL\_accel\_ptr** = NULL
- std::string cosmology ="PLANCK15"

# 8.14.1 Member Data Documentation

```
8.14.1.1 betappe
```

double\* gen\_params::betappe

ppE coefficient for the phase modification - vector for multiple modifications

# 8.14.1.2 bppe

int\* gen\_params::bppe

ppE b parameter (power of the frequency) - vector for multiple modifications

#### 8.14.1.3 f\_ref

double gen\_params:: $f_ref = 0$ 

Reference frequency for PhenomPv2

# 8.14.1.4 incl\_angle

double gen\_params::incl\_angle

\*angle between angular momentum and the total momentum

# 8.14.1.5 Luminosity\_Distance

double gen\_params::Luminosity\_Distance

Luminosity distance to the source

```
8.14.1.6 mass1
double gen_params::mass1
mass of the larger body in Solar Masses
8.14.1.7 mass2
double gen_params::mass2
mass of the smaller body in Solar Masses
8.14.1.8 Nmod
int gen_params::Nmod
Number of phase modificatinos
8.14.1.9 NSflag
bool gen_params::NSflag
BOOL flag for early termination of NS binaries
8.14.1.10 phic
double gen_params::phic =0
coalescence phase of the binary
8.14.1.11 RA
double gen_params::RA
Equatorial coordinates of source
8.14.1.12 spin1
double gen_params::spin1[3]
Spin vector of the larger mass [Sx,Sy,Sz]
8.14.1.13 spin2
double gen_params::spin2[3]
Spin vector of the smaller mass [Sx,Sy,Sz]
```

## 8.14.1.14 tc

```
double gen_params::tc =0
```

coalescence time of the binary

#### 8.14.1.15 theta

```
double gen_params::theta
```

spherical angles for the source location relative to the detector

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

• include/util.h

# 8.15 waveform\_generator\_ext.gen\_params\_py Class Reference

Python wrapper for the generation parameters structure, as defined in util.cpp.

# 8.15.1 Detailed Description

Python wrapper for the generation parameters structure, as defined in util.cpp.

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

• gw\_analysis\_tools\_py/src/waveform\_generator\_ext.pyx

# 8.16 GPUplan Struct Reference

# **Public Attributes**

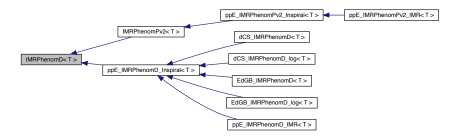
- · int device\_id
- double \* device data
- double \* host\_data
- int \* host\_lag
- int \* device\_lag
- int \* device lags
- int \* initial\_lag
- cudaStream\_t stream

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

• include/autocorrelation\_cuda.hu

# 8.17 IMRPhenomD < T > Class Template Reference

Inheritance diagram for IMRPhenomD< T >:



#### **Public Member Functions**

- virtual void **fisher\_calculation** (double \*frequency, int length, gen\_params \*parameters, double \*\*amplitude\_deriv, double \*\*phase\_deriv, double \*amplitude, int \*amp\_tapes, int \*phase\_tapes)
- virtual void change\_parameter\_basis (T \*old\_param, T \*new\_param, bool sky\_average)
   Convience method to change parameter basis between common Fisher parameters and the intrinsic parameters of
- IMRPhenomD.
   virtual void construct\_amplitude\_derivative (double \*frequencies, int length, int dimension, double \*\*amplitude derivative, source parameters< double > \*input params, int \*tapes=NULL)
  - Construct the derivative of the amplitude for a given source evaluated by the given frequency.
- virtual void construct\_phase\_derivative (double \*frequencies, int length, int dimension, double \*\*phase\_←
  derivative, source\_parameters< double > \*input\_params, int \*tapes=NULL)
  - Construct the derivative of the phase for a given source evaluated by the given frequency.
- virtual void amplitude tape (source parameters< double > \*input params, int \*tape)
  - Creates the tapes for derivatives of the amplitude.
- virtual void phase\_tape (source\_parameters < double > \*input\_params, int \*tape)
  - Creates the tapes for derivatives of phase.
- virtual int construct\_waveform (T \*frequencies, int length, std::complex < T > \*waveform, source\_parameters < T > \*params)
  - Constructs the waveform as outlined by.
- virtual std::complex < T > construct waveform (T frequency, source parameters < T > \*params)
  - overloaded method to evaluate the waveform for one frequency instead of an array
- virtual int construct\_amplitude (T \*frequencies, int length, T \*amplitude, source\_parameters < T > \*params)
   Constructs the Amplitude as outlined by IMRPhenomD.
- virtual int construct\_phase (T \*frequencies, int length, T \*phase, source\_parameters< T > \*params)

  Constructs the Phase as outlined by IMRPhenomD.
- virtual T build\_amp (T f, lambda\_parameters< T > \*lambda, source\_parameters< T > \*params, useful\_powers< T > \*pows, T \*amp\_coeff, T \*deltas)
  - constructs the IMRPhenomD amplitude for frequency f
- virtual T build\_phase (T f, lambda\_parameters< T > \*lambda, source\_parameters< T > \*params, useful\_powers< T > \*pows, T \*phase\_coeff)
  - constructs the IMRPhenomD phase for frequency f
- virtual T assign\_lambda\_param\_element (source\_parameters < T > \*source\_param, int i)
  - Calculate the lambda parameters from Khan et al for element i.
- virtual void assign\_lambda\_param (source\_parameters < T > \*source\_param, lambda\_parameters < T > \*lambda)

Wrapper for the Lambda parameter assignment that handles the looping.

virtual void precalc\_powers\_ins (T f, T M, useful\_powers< T > \*Mf\_pows)

Pre-calculate powers of Mf, to speed up calculations for the inspiral waveform (both amplitude and phase.

virtual void precalc\_powers\_PI (useful\_powers< T > \*PI\_pows)

Pre-calculate powers of pi, to speed up calculations for the inspiral phase.

virtual void precalc\_powers\_ins\_phase (T f, T M, useful\_powers< T > \*Mf\_pows)

Pre-calculate powers of Mf, to speed up calculations for the inspiral phase.

virtual void precalc\_powers\_ins\_amp (T f, T M, useful\_powers< T > \*Mf\_pows)

Pre-calculate powers of Mf, to speed up calculations for the inspiral amplitude.

 $\bullet \ \ virtual\ void\ assign\_pn\_amplitude\_coeff\ (source\_parameters < T > *source\_param,\ T\ *coeff)$ 

Calculates the static PN coeffecients for the amplitude.

virtual void assign\_static\_pn\_phase\_coeff (source\_parameters < T > \*source\_param, T \*coeff)

Calculates the static PN coeffecients for the phase - coeffecients 0,1,2,3,4,7.

- virtual void assign\_nonstatic\_pn\_phase\_coeff (source\_parameters< T > \*source\_param, T \*coeff, T f)

  Calculates the dynamic PN phase coefficients 5,6.
- virtual void assign\_nonstatic\_pn\_phase\_coeff\_deriv (source\_parameters < T > \*source\_param, T \*Dcoeff, T f)

Calculates the derivative of the dynamic PN phase coefficients 5,6.

virtual void post\_merger\_variables (source\_parameters < T > \*source\_param)

Calculates the post-merger ringdown frequency and dampening frequency.

• virtual T fpeak (source\_parameters < T > \*params, lambda\_parameters < T > \*lambda)

Solves for the peak frequency, where the waveform transitions from intermediate to merger-ringdown.

virtual T amp\_ins (T f, source\_parameters < T > \*param, T \*pn\_coeff, lambda\_parameters < T > \*lambda, useful powers < T > \*pow)

Calculates the scaled inspiral amplitude A/A0 for frequency f with precomputed powers of MF and PI.

- virtual T Damp\_ins (T f, source\_parameters < T > \*param, T \*pn\_coeff, lambda\_parameters < T > \*lambda)

  Calculates the derivative wrt frequency for the scaled inspiral amplitude A/A0 for frequency f.
- virtual T phase\_ins (T f, source\_parameters < T > \*param, T \*pn\_coeff, lambda\_parameters < T > \*lambda, useful\_powers < T > \*pow)

Calculates the inspiral phase for frequency f with precomputed powers of MF and PI for speed.

virtual T Dphase\_ins (T f, source\_parameters< T > \*param, T \*pn\_coeff, lambda\_parameters< T > \*lambda)

Calculates the derivative of the inspiral phase for frequency f.

virtual T amp\_mr (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)

Calculates the scaled merger-ringdown amplitude A/A0 for frequency f.

• virtual T phase\_mr (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)

Calculates the merger-ringdown phase for frequency f.

 $\bullet \ \ virtual \ T \ Damp\_mr \ (T \ f, source\_parameters < T > *param, lambda\_parameters < T > *lambda) \\$ 

Calculates the derivative wrt frequency for the scaled merger-ringdown amplitude A/A0 for frequency f.

virtual T Dphase\_mr (T f, source\_parameters< T > \*param, lambda\_parameters< T > \*lambda)

Calculates the derivative of the merger-ringdown phase for frequency f.

 $\bullet \ \ virtual \ T \ amp\_int \ (T \ f, source\_parameters < T > *param, lambda\_parameters < T > *lambda, T *deltas) \\$ 

Calculates the intermediate phase for frequency f.

- virtual T Dphase\_int (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)
   Calculates the derivative of the intermediate phase for frequency f.
- virtual void phase\_connection\_coefficients (source\_parameters< T > \*param, lambda\_parameters< T > \*lambda, T \*pn coeffs)

Calculates the phase connection coefficients alpha{0,1} and beta{0,1}.

- virtual T calculate\_beta1 (source\_parameters < T > \*param, lambda\_parameters < T > \*lambda, T \*pn ←
   \_coeffs)
- virtual T calculate\_beta0 (source\_parameters < T > \*param, lambda\_parameters < T > \*lambda, T \*pn ←
   \_coeffs)
- virtual T calculate\_alpha1 (source\_parameters< T > \*param, lambda\_parameters< T > \*lambda)
- virtual T calculate alpha0 (source parameters< T > \*param, lambda parameters< T > \*lambda)
- virtual void amp\_connection\_coeffs (source\_parameters < T > \*param, lambda\_parameters < T > \*lambda, T \*pn\_coeffs, T \*coeffs)

Solves for the connection coefficients to ensure the transition from inspiral to merger ringdown is continuous and smooth.

- virtual T calculate\_delta\_parameter\_0 (T f1, T f2, T f3, T v1, T v2, T v3, T dd1, T dd3, T M)
   Calculates the delta\_0 component.
- virtual T calculate\_delta\_parameter\_1 (T f1, T f2, T f3, T v1, T v2, T v3, T dd1, T dd3, T M)
   Calculates the delta\_1 component.
- virtual T calculate\_delta\_parameter\_2 (T f1, T f2, T f3, T v1, T v2, T v3, T dd1, T dd3, T M)
   Calculates the delta 2 component.
- virtual T calculate\_delta\_parameter\_3 (T f1, T f2, T f3, T v1, T v2, T v3, T dd1, T dd3, T M)
   Calculates the delta\_3 component.
- virtual T calculate\_delta\_parameter\_4 (T f1, T f2, T f3, T v1, T v2, T v3, T dd1, T dd3, T M)
   Calculates the delta 4 component.

#### 8.17.1 Member Function Documentation

## 8.17.1.1 amp\_ins()

Calculates the scaled inspiral amplitude A/A0 for frequency f with precomputed powers of MF and PI.

return a T

additional argument contains useful powers of MF and PI in structure userful\_powers

## 8.17.1.2 amp\_int()

Calculates the scaled intermediate range amplitude A/A0 for frequency f.

return a T

## 8.17.1.3 amp\_mr()

Calculates the scaled merger-ringdown amplitude A/A0 for frequency f.

return a T

## 8.17.1.4 amplitude\_tape()

Creates the tapes for derivatives of the amplitude.

For efficiency in long runs of large sets of fishers, the tapes can be precomputed and reused

## **Parameters**

input_params	source parameters structure of the desired source
tape	tape ids

 $\label{eq:local_$ 

# 8.17.1.5 assign\_nonstatic\_pn\_phase\_coeff()

Calculates the dynamic PN phase coefficients 5,6.

f is in Hz

# 8.17.1.6 assign\_nonstatic\_pn\_phase\_coeff\_deriv()

Calculates the derivative of the dynamic PN phase coefficients 5,6.

f is in Hz

## 8.17.1.7 build\_amp()

constructs the IMRPhenomD amplitude for frequency f

arguments: numerical parameters from Khan et al  $lambda_parameters$  structure, source\_parameters structure, useful\_powers<T> structure, PN parameters for the inspiral portions of the waveform, and the delta parameters for the intermediate region, numerically solved for using the amp\_connection\_coeffs function

## 8.17.1.8 build\_phase()

constructs the IMRPhenomD phase for frequency f

arguments: numerical parameters from Khan et al lambda\_parameters structure, source\_parameters structure, useful\_powers structure, PN parameters for the inspiral portions of the waveform

## 8.17.1.9 calculate\_delta\_parameter\_0()

Calculates the delta\_0 component.

Solved in Mathematica and imported to C

## 8.17.1.10 calculate\_delta\_parameter\_1()

Calculates the delta\_1 component.

Solved in Mathematica and imported to C

# 8.17.1.11 calculate\_delta\_parameter\_2()

Calculates the delta\_2 component.

Solved in Mathematica and imported to C

# 8.17.1.12 calculate\_delta\_parameter\_3()

Calculates the delta\_3 component.

Solved in Mathematica and imported to C

# 8.17.1.13 calculate\_delta\_parameter\_4()

Calculates the delta\_4 component.

Solved in Mathematica and imported to C

## 8.17.1.14 change\_parameter\_basis()

Convience method to change parameter basis between common Fisher parameters and the intrinsic parameters of IMRPhenomD.

Takes input array of old parameters and ouputs array of transformed parameters

# **Parameters**

old_param	array of old params, order {A0, tc, phic, chirpmass, eta, spin1, spin2}
new_param	output new array: order {m1,m2,DL, spin1,spin2,phic,tc}

# 8.17.1.15 construct\_amplitude()

Constructs the Amplitude as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output amplitude, and a source\_parameters structure

#### **Parameters**

frequencies	T array of frequencies the waveform is to be evaulated at
length	integer length of the input array of frequencies and the output array
amplitude	output T array for the amplitude
params	Structure of source parameters to be initilized before computation

Reimplemented in EdGB\_IMRPhenomD< T >, EdGB\_IMRPhenomD\_log< T >, dCS\_IMRPhenomD< T >, and dCS\_IMRPhenomD\_log< T >.

## 8.17.1.16 construct\_amplitude\_derivative()

Construct the derivative of the amplitude for a given source evaluated by the given frequency.

Order of output: dh/d \theta : \theta \el {A0,tc, phic, chirp mass, eta, symmetric spin, antisymmetric spin}

# Parameters

frequencies	input array of frequency
length	length of the frequency array
amplitude_derivative	< dimension of the fisher output array for all the derivatives double[dimension][length]
input_params	Source parameters structure for the source
tapes	int array of tape ids, if NULL, these will be calculated

 $Reimplemented \ in \ ppE\_IMRPhenomD\_IMR < T>, \ and \ ppE\_IMRPhenomD\_Inspiral < T>.$ 

# 8.17.1.17 construct\_phase()

Constructs the Phase as outlined by IMRPhenomD.

arguments: array of frequencies, length of that array, T array for the output phase, and a source\_parameters structure

#### **Parameters**

frequencies	T array of frequencies the waveform is to be evaluated at
length	integer length of the input and output arrays
phase	output T array for the phasee
params	structure of source parameters to be calculated before computation

Reimplemented in EdGB\_IMRPhenomD< T >, EdGB\_IMRPhenomD\_log< T >, dCS\_IMRPhenomD< T >, and dCS\_IMRPhenomD\_log< T >.

## 8.17.1.18 construct\_phase\_derivative()

Construct the derivative of the phase for a given source evaluated by the given frequency.

Order of output: dh/d \theta : \theta \el {A0,tc, phic, chirp mass, eta, symmetric spin, antisymmetric spin}

# **Parameters**

frequencies	input array of frequency
length	length of the frequency array
phase_derivative	< dimension of the fisher output array for all the derivatives double[dimension][length]
input_params	Source parameters structure for the source
tapes	int array of tape ids, if NULL, these will be calculated

Reimplemented in ppE\_IMRPhenomD\_IMR< T >, and ppE\_IMRPhenomD\_Inspiral< T >.

# **8.17.1.19** construct\_waveform() [1/2]

Constructs the waveform as outlined by.

arguments: array of frequencies, length of that array, a complex array for the output waveform, and a source\_parameters structure

#### **Parameters**

frequencies	T array of frequencies the waveform is to be evaluated at
length	integer length of the array of frequencies and the waveform
waveform	complex T array for the waveform to be output

Reimplemented in EdGB\_IMRPhenomD< T >, EdGB\_IMRPhenomD\_log< T >, dCS\_IMRPhenomD< T >, and dCS\_IMRPhenomD\_log< T >.

```
8.17.1.20 construct_waveform() [2/2]
```

overloaded method to evaluate the waveform for one frequency instead of an array

#### **Parameters**

frequency	T array of frequencies the waveform is to be evaluated at
-----------	---

# 8.17.1.21 Damp\_ins()

Calculates the derivative wrt frequency for the scaled inspiral amplitude A/A0 for frequency f.

This is an analytic derivative for the smoothness condition on the amplitude connection

return a T

## 8.17.1.22 Damp\_mr()

Calculates the derivative wrt frequency for the scaled merger-ringdown amplitude A/A0 for frequency f.

This is an analytic derivative for the smoothness condition on the amplitude connection

The analytic expression was obtained from Mathematica - See the mathematica folder for code

return a T

## 8.17.1.23 Dphase\_ins()

Calculates the derivative of the inspiral phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented in ppE\_IMRPhenomD\_Inspiral < T >, and ppE\_IMRPhenomPv2\_Inspiral < T >.

# 8.17.1.24 Dphase\_int()

Calculates the derivative of the intermediate phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented in ppE\_IMRPhenomD\_IMR< T >, and ppE\_IMRPhenomPv2\_IMR< T >.

## 8.17.1.25 Dphase\_mr()

Calculates the derivative of the merger-ringdown phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented in ppE\_IMRPhenomD\_IMR< T >, and ppE\_IMRPhenomPv2\_IMR< T >.

```
8.17.1.26 fpeak()
```

Solves for the peak frequency, where the waveform transitions from intermediate to merger-ringdown.

returns Hz

#### 8.17.1.27 phase\_connection\_coefficients()

Calculates the phase connection coefficients alpha{0,1} and beta{0,1}.

Note: these coefficients are stored in the lambda parameter structure, not a separate array

#### 8.17.1.28 phase\_ins()

Calculates the inspiral phase for frequency f with precomputed powers of MF and PI for speed.

return a T

extra argument of precomputed powers of MF and pi, contained in the structure useful\_powers<T>

Reimplemented in ppE\_IMRPhenomD\_Inspiral<T>, and ppE\_IMRPhenomPv2\_Inspiral<T>.

#### 8.17.1.29 phase\_int()

Calculates the intermediate phase for frequency f.

return a T

Reimplemented in ppE\_IMRPhenomD\_IMR< T >, and ppE\_IMRPhenomPv2\_IMR< T >.

# 8.17.1.30 phase\_mr()

Calculates the merger-ringdown phase for frequency f.

return a T

Reimplemented in ppE\_IMRPhenomD\_IMR< T >, and ppE\_IMRPhenomPv2\_IMR< T >.

#### 8.17.1.31 phase\_tape()

Creates the tapes for derivatives of phase.

For efficiency in long runs of large sets of fishers, the tapes can be precomputed and reused

# Parameters

input_params	source parameters structure of the desired source
tape	tape ids

Reimplemented in ppE\_IMRPhenomD\_IMR< T>, and ppE\_IMRPhenomD\_Inspiral< T>.

#### 8.17.1.32 post\_merger\_variables()

Calculates the post-merger ringdown frequency and dampening frequency.

Returns in Hz - assigns fRD to var[0] and fdamp to var[1]

# 8.17.1.33 precalc\_powers\_ins()

Pre-calculate powers of Mf, to speed up calculations for the inspiral waveform (both amplitude and phase.

It seems the pow() function is very slow, so to speed things up, powers of Mf will be precomputed and passed to the functions within the frequency loops

## 8.17.1.34 precalc\_powers\_ins\_amp()

Pre-calculate powers of Mf, to speed up calculations for the inspiral amplitude.

It seems the pow() function is very slow, so to speed things up, powers of Mf will be precomputed and passed to the functions within the frequency loops

# 8.17.1.35 precalc\_powers\_ins\_phase()

Pre-calculate powers of Mf, to speed up calculations for the inspiral phase.

It seems the pow() function is very slow, so to speed things up, powers of Mf will be precomputed and passed to the functions within the frequency loops

# 8.17.1.36 precalc\_powers\_PI()

Pre-calculate powers of pi, to speed up calculations for the inspiral phase.

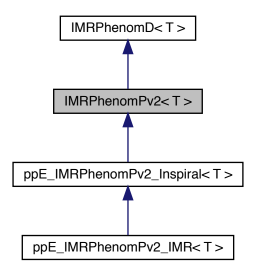
It seems the pow() function is very slow, so to speed things up, powers of PI will be precomputed and passed to the functions within the frequency loops

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

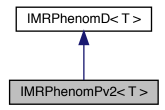
- include/IMRPhenomD.h
- src/IMRPhenomD.cpp

# 8.18 IMRPhenomPv2< T > Class Template Reference

Inheritance diagram for IMRPhenomPv2< T >:



Collaboration diagram for IMRPhenomPv2< T >:



# **Public Member Functions**

- virtual T alpha (T omega, T q, T chi2l, T chi2)
- virtual T epsilon (T omega, T q, T chi2l, T chi2)
- virtual void calculate\_euler\_coeffs (alpha\_coeffs< T > \*acoeffs, epsilon\_coeffs< T > \*ecoeffs, source\_parameters< T > \*params)

Pre calculate euler angle coefficients.

- virtual T d (int I, int mp, int m, T s)
- virtual int construct\_waveform (T \*frequencies, int length, std::complex< T > \*waveform\_plus, std
   ::complex< T > \*waveform\_cross, source\_parameters< T > \*params)

Constructs the waveform for IMRPhenomPv2 - uses IMRPhenomD, then twists up.

- virtual void **WignerD** (T d2[5], T dm2[5], useful\_powers< T > \*pows, source\_parameters< T > \*params)
- virtual void calculate\_twistup (T alpha, std::complex< T > \*hp\_factor, std::complex< T > \*hc\_factor, T d2[5], T dm2[5], sph\_harm< T > \*sph\_harm)
- virtual void calculate\_euler\_angles (T \*alpha, T \*epsilon, useful\_powers< T > \*pows, alpha\_coeffs< T > \*acoeffs, epsilon\_coeffs< T > \*ecoeffs)
- virtual void PhenomPv2\_Param\_Transform (source\_parameters < T > \*params)
- virtual void PhenomPv2\_Param\_Transform\_J (source\_parameters < T > \*params)
- virtual T L2PN (T eta, useful powers< T > \*pow)

#### 8.18.1 Member Function Documentation

#### 8.18.1.1 calculate\_euler\_coeffs()

Pre calculate euler angle coefficients.

Straight up stolen from LALsuite

# 8.18.1.2 construct\_waveform()

Constructs the waveform for IMRPhenomPv2 - uses IMRPhenomD, then twists up.

arguments: array of frequencies, length of that array, a complex array for the output waveform, and a source\_parameters structure

# **Parameters**

frequencies	T array of frequencies the waveform is to be evaluated at
length	integer length of the array of frequencies and the waveform
waveform_plus	complex T array for the plus polariaztion waveform to be output
waveform_cross	complex T array for the cross polarization waveform to be output

## 8.18.1.3 PhenomPv2\_Param\_Transform()

/Brief Parameter transformtion to precalculate needed parameters for PhenomP from source parameters

Pretty much stolen verbatim from lalsuite

Reimplemented in ppE\_IMRPhenomPv2\_IMR< T >, and ppE\_IMRPhenomPv2\_Inspiral< T >.

## 8.18.1.4 PhenomPv2\_Param\_Transform\_J()

/Brief Parameter transformtion to precalculate needed parameters for PhenomP from source parameters – assumed inclination of total angular momentum J is given, not orbital angular momentum (in source frame (Lhat == zhat)

Pretty much stolen verbatim from lalsuite

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

- include/IMRPhenomP.h
- src/IMRPhenomP.cpp

# 8.19 lambda parameters < T > Struct Template Reference

**Public Attributes** 

- T rho [4]
- T v2
- T gamma [4]
- T sigma [5]
- T beta [5]
- T alpha [7]

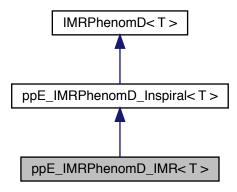
The documentation for this struct was generated from the following file:

• include/IMRPhenomD.h

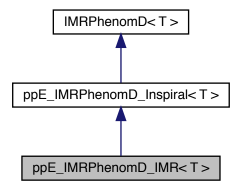
# 8.20 ppE\_IMRPhenomD\_IMR < T > Class Template Reference

#include <ppE\_IMRPhenomD.h>

Inheritance diagram for ppE\_IMRPhenomD\_IMR< T >:



Collaboration diagram for ppE\_IMRPhenomD\_IMR< T >:



# **Public Member Functions**

- virtual T Dphase\_mr (T f, source\_parameters< T > \*param, lambda\_parameters< T > \*lambda)
   Calculates the derivative of the merger-ringdown phase for frequency f.
- virtual T phase\_mr (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)
   Calculates the merger-ringdown phase for frequency f.
- virtual T phase\_int (T f, source\_parameters< T > \*param, lambda\_parameters< T > \*lambda)

Calculates the intermediate phase for frequency f.

virtual T Dphase\_int (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)

Calculates the derivative of the intermediate phase for frequency f.

- virtual void **fisher\_calculation** (double \*frequency, int length, gen\_params \*parameters, double \*\*amplitude\_deriv, double \*\*phase\_deriv, double \*amplitude, int \*amp\_tapes, int \*phase\_tapes)
- virtual void amplitude\_tape (source\_parameters< double > \*input\_params, int \*tape)

Creates the tapes for derivatives of the amplitude.

virtual void phase tape (source parameters< double > \*input params, int \*tape)

Creates the tapes for derivatives of phase.

 virtual void construct\_amplitude\_derivative (double \*frequencies, int length, int dimension, double \*\*amplitude\_derivative, source\_parameters< double > \*input\_params, int \*tapes=NULL)

Construct the derivative of the amplitude for a given source evaluated by the given frequency.

virtual void construct\_phase\_derivative (double \*frequencies, int length, int dimension, double \*\*phase\_←
derivative, source\_parameters< double > \*input\_params, int \*tapes=NULL)

Construct the derivative of the phase for a given source evaluated by the given frequency.

## 8.20.1 Detailed Description

```
\label{template} \begin{split} \text{template} &< \text{class T}> \\ \text{class ppE\_IMRPhenomD\_IMR} &< \text{T}> \end{split}
```

Class that extends the IMRPhenomD waveform to include non-GR terms in the full phase. This is an appropriate waveform choice for propagation effects

# 8.20.2 Member Function Documentation

# 8.20.2.1 amplitude\_tape()

Creates the tapes for derivatives of the amplitude.

For efficiency in long runs of large sets of fishers, the tapes can be precomputed and reused

# **Parameters**

input_params	source parameters structure of the desired source
tape	tape ids

Reimplemented from ppE\_IMRPhenomD\_Inspiral< T >.

## 8.20.2.2 construct\_amplitude\_derivative()

Construct the derivative of the amplitude for a given source evaluated by the given frequency.

Order of output: dh/d \theta : \theta \el {A0,tc, phic, chirp mass, eta, symmetric spin, antisymmetric spin}

#### **Parameters**

frequencies	input array of frequency
length	length of the frequency array
amplitude_derivative	< dimension of the fisher output array for all the derivatives double[dimension][length]
input_params	Source parameters structure for the source
tapes	int array of tape ids, if NULL, these will be calculated

Reimplemented from ppE\_IMRPhenomD\_Inspiral< T >.

## 8.20.2.3 construct\_phase\_derivative()

Construct the derivative of the phase for a given source evaluated by the given frequency.

Order of output: dh/d \theta: \theta \el {A0,tc, phic, chirp mass, eta, symmetric spin, antisymmetric spin}

## **Parameters**

frequencies	input array of frequency
length	length of the frequency array
phase_derivative	< dimension of the fisher output array for all the derivatives double[dimension][length]
input_params	Source parameters structure for the source
tapes	int array of tape ids, if NULL, these will be calculated

Reimplemented from ppE\_IMRPhenomD\_Inspiral< T >.

# 8.20.2.4 Dphase\_int()

Calculates the derivative of the intermediate phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented from IMRPhenomD< T >.

# 8.20.2.5 Dphase\_mr()

Calculates the derivative of the merger-ringdown phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented from IMRPhenomD< T >.

# 8.20.2.6 phase\_int()

Calculates the intermediate phase for frequency f.

return a T

Reimplemented from IMRPhenomD< T >.

# 8.20.2.7 phase\_mr()

Calculates the merger-ringdown phase for frequency f.

return a T

Reimplemented from IMRPhenomD< T >.

## 8.20.2.8 phase\_tape()

Creates the tapes for derivatives of phase.

For efficiency in long runs of large sets of fishers, the tapes can be precomputed and reused

# Parameters

input_params	source parameters structure of the desired source
tape	tape ids

Reimplemented from ppE\_IMRPhenomD\_Inspiral< T >.

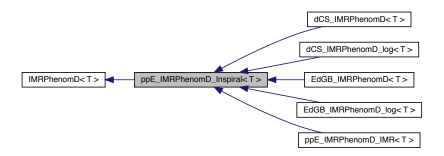
The documentation for this class was generated from the following files:

- include/ppE\_IMRPhenomD.h
- src/ppE\_IMRPhenomD.cpp

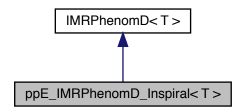
# 8.21 ppE\_IMRPhenomD\_Inspiral < T > Class Template Reference

```
#include <ppE_IMRPhenomD.h>
```

Inheritance diagram for ppE\_IMRPhenomD\_Inspiral< T >:



Collaboration diagram for ppE IMRPhenomD Inspiral < T >:



# **Public Member Functions**

virtual T phase\_ins (T f, source\_parameters < T > \*param, T \*pn\_coeff, lambda\_parameters < T > \*lambda, useful\_powers < T > \*pow)

Overloaded method for the inspiral portion of the phase.

virtual T Dphase\_ins (T f, source\_parameters< T > \*param, T \*pn\_coeff, lambda\_parameters< T > \*lambda)

Calculates the derivative of the inspiral phase for frequency f.

- virtual void **fisher\_calculation** (double \*frequency, int length, gen\_params \*parameters, double \*\*amplitude deriv, double \*\*phase deriv, double \*amplitude, int \*amp tapes, int \*phase tapes)
- virtual void amplitude\_tape (source\_parameters< double > \*input\_params, int \*tape)

Creates the tapes for derivatives of the amplitude.

• virtual void phase\_tape (source\_parameters< double > \*input\_params, int \*tape)

Creates the tapes for derivatives of phase.

 virtual void construct\_amplitude\_derivative (double \*frequencies, int length, int dimension, double \*\*amplitude\_derivative, source\_parameters< double > \*input\_params, int \*tapes=NULL)

Construct the derivative of the amplitude for a given source evaluated by the given frequency.

virtual void construct\_phase\_derivative (double \*frequencies, int length, int dimension, double \*\*phase\_←
derivative, source\_parameters< double > \*input\_params, int \*tapes=NULL)

Construct the derivative of the phase for a given source evaluated by the given frequency.

# 8.21.1 Detailed Description

```
\label{template} \begin{split} \text{template} \! < \! \text{class T} \! > \\ \text{class ppE\_IMRPhenomD\_Inspiral} \! < \! \text{T} \! > \end{split}
```

Class that extends the IMRPhenomD waveform to include non-GR terms in the inspiral portion of the phase. This is an appropriate waveform choice for generation effects, but not necessarily for propagation effects

## 8.21.2 Member Function Documentation

# 8.21.2.1 amplitude\_tape()

Creates the tapes for derivatives of the amplitude.

For efficiency in long runs of large sets of fishers, the tapes can be precomputed and reused

## **Parameters**

input_params	source parameters structure of the desired source
tape	tape ids

Reimplemented from IMRPhenomD< T >.

Reimplemented in ppE\_IMRPhenomD\_IMR< T >.

# 8.21.2.2 construct\_amplitude\_derivative()

Construct the derivative of the amplitude for a given source evaluated by the given frequency.

Order of output: dh/d \theta : \theta \el {A0,tc, phic, chirp mass, eta, symmetric spin, antisymmetric spin}

#### **Parameters**

frequencies	input array of frequency	
length length of the frequency array		
amplitude_derivative	tive < dimension of the fisher output array for all the derivatives double[dimension][length]	
input_params Source parameters structure for the source		
tapes int array of tape ids, if NULL, these will be calculated		

Reimplemented from IMRPhenomD < T >.

Reimplemented in ppE\_IMRPhenomD\_IMR< T >.

## 8.21.2.3 construct\_phase\_derivative()

Construct the derivative of the phase for a given source evaluated by the given frequency.

Order of output: dh/d \theta: \theta \el {A0,tc, phic, chirp mass, eta, symmetric spin, antisymmetric spin}

#### **Parameters**

frequencies	input array of frequency	
length	length of the frequency array	
phase_derivative	e < dimension of the fisher output array for all the derivatives double[dimension][length]	
input_params Source parameters structure for the source		
tapes int array of tape ids, if NULL, these will be calculated		

Reimplemented from IMRPhenomD< T >.

Reimplemented in ppE\_IMRPhenomD\_IMR< T >.

# 8.21.2.4 Dphase\_ins()

Calculates the derivative of the inspiral phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented from IMRPhenomD< T >.

# 8.21.2.5 phase\_tape()

Creates the tapes for derivatives of phase.

For efficiency in long runs of large sets of fishers, the tapes can be precomputed and reused

## **Parameters**

input_params	source parameters structure of the desired source
tape	tape ids

Reimplemented from IMRPhenomD< T >.

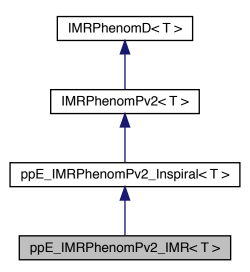
Reimplemented in ppE\_IMRPhenomD\_IMR< T >.

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

- include/ppE\_IMRPhenomD.h
- src/ppE\_IMRPhenomD.cpp

# 8.22 ppE\_IMRPhenomPv2\_IMR < T > Class Template Reference

Inheritance diagram for ppE\_IMRPhenomPv2\_IMR< T >:



Collaboration diagram for ppE\_IMRPhenomPv2\_IMR< T >:



# **Public Member Functions**

 $\bullet \ \ virtual \ T \ phase\_mr \ (T \ f, \ source\_parameters < T > *param, \ lambda\_parameters < T > *lambda) \\$ 

Calculates the merger-ringdown phase for frequency f.

- virtual T Dphase\_mr (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)
   Calculates the derivative of the merger-ringdown phase for frequency f.
- virtual T phase\_int (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)
   Calculates the intermediate phase for frequency f.
- virtual T Dphase\_int (T f, source\_parameters < T > \*param, lambda\_parameters < T > \*lambda)
   Calculates the derivative of the intermediate phase for frequency f.
- virtual void PhenomPv2 Param Transform (source parameters < T > \*params)

#### 8.22.1 Member Function Documentation

# 8.22.1.1 Dphase\_int()

Calculates the derivative of the intermediate phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented from IMRPhenomD< T >.

# 8.22.1.2 Dphase\_mr()

Calculates the derivative of the merger-ringdown phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented from IMRPhenomD< T >.

## 8.22.1.3 phase\_int()

Calculates the intermediate phase for frequency f.

return a T

Reimplemented from IMRPhenomD< T >.

#### 8.22.1.4 phase\_mr()

Calculates the merger-ringdown phase for frequency f.

return a T

Reimplemented from IMRPhenomD< T >.

## 8.22.1.5 PhenomPv2\_Param\_Transform()

```
\label{template} $$\operatorname{DPE_IMRPhenomPv2_IMR} < T >:: PhenomPv2_Param_Transform ($$\operatorname{source\_parameters} < T > * params ) [virtual]
```

/Brief Parameter transformtion to precalculate needed parameters for PhenomP from source parameters

Pretty much stolen verbatim from lalsuite

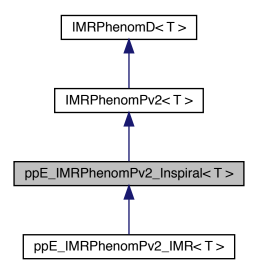
Reimplemented from ppE\_IMRPhenomPv2\_Inspiral < T >.

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

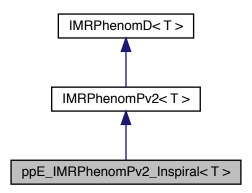
- include/ppE IMRPhenomP.h
- src/ppE\_IMRPhenomP.cpp

# 8.23 ppE\_IMRPhenomPv2\_Inspiral < T > Class Template Reference

Inheritance diagram for ppE\_IMRPhenomPv2\_Inspiral < T >:



Collaboration diagram for ppE\_IMRPhenomPv2\_Inspiral< T >:



# **Public Member Functions**

virtual T phase\_ins (T f, source\_parameters < T > \*param, T \*pn\_coeff, lambda\_parameters < T > \*lambda, useful\_powers < T > \*pow)

Calculates the inspiral phase for frequency f with precomputed powers of MF and PI for speed.

virtual T Dphase\_ins (T f, source\_parameters< T > \*param, T \*pn\_coeff, lambda\_parameters< T > \*lambda)

Calculates the derivative of the inspiral phase for frequency f.

virtual void PhenomPv2\_Param\_Transform (source\_parameters< T > \*params)

## 8.23.1 Member Function Documentation

#### 8.23.1.1 Dphase\_ins()

Calculates the derivative of the inspiral phase for frequency f.

For phase continuity and smoothness return a T

Reimplemented from IMRPhenomD< T >.

#### 8.23.1.2 phase\_ins()

Calculates the inspiral phase for frequency f with precomputed powers of MF and PI for speed.

return a T

extra argument of precomputed powers of MF and pi, contained in the structure useful\_powers<T>

Reimplemented from IMRPhenomD< T>.

## 8.23.1.3 PhenomPv2\_Param\_Transform()

/Brief Parameter transformtion to precalculate needed parameters for PhenomP from source parameters

Pretty much stolen verbatim from lalsuite

Reimplemented from IMRPhenomPv2< T >.

Reimplemented in ppE\_IMRPhenomPv2\_IMR< T >.

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

- include/ppE IMRPhenomP.h
- src/ppE\_IMRPhenomP.cpp

# 8.24 sampler Class Reference

```
#include <mcmc_sampler_internals.h>
```

# **Public Attributes**

- int types\_of\_steps = 5
- double \*\* step\_prob
- double \*\* prob boundaries
- double \* chain\_temps
- bool \* waiting
- int \* chain pos
- double swp\_freq
- int chain\_N
- int numThreads
- int N\_steps
- int dimension
- int min\_dim
- int max\_dim
- · bool fisher\_exist
- bool \* de\_primed
- int \* priority
- bool \* ref\_chain\_status
- bool prioritize\_cold\_chains = true
- double \*\*\* output
- · bool pool
- int progress =0
- bool show\_progress
- · int num threads
- int history\_length =500
- int history\_update =5
- int \* current\_hist\_pos

- double \*\*\* history
- int \*\*\* history\_status
- · double \* current\_likelihoods
- int \* check stepsize freq
- double \* max\_target\_accept\_ratio
- double \* min\_target\_accept\_ratio
- int \* gauss\_last\_accept\_ct
- int \* gauss\_last\_reject\_ct
- int \* de\_last\_accept\_ct
- int \* de\_last\_reject\_ct
- int \* fish\_last\_accept\_ct
- int \* fish\_last\_reject\_ct
- double \*\* randgauss width
- double \*\*\* fisher vecs
- double \*\* fisher\_vals
- int \* fisher\_update\_ct
- int fisher\_update\_number =200
- std::function< double(double \*, int \*, int, int)> lp
- std::function< double(double \*, int \*, int, int)> II
- std::function< void(double \*, int \*, int, double \*\*, int)> fish
- gsl\_rng \*\* rvec
- int \* nan counter
- int \* num\_gauss
- int \* num\_fish
- int \* num\_de
- int \* num\_mmala
- int \* num\_RJstep
- double time\_elapsed\_cpu
- double time\_elapsed\_wall
- double time\_elapsed\_cpu\_ac
- double time\_elapsed\_wall\_ac
- int \* fish\_accept\_ct
- int \* fish\_reject\_ct
- int \* de\_accept\_ct
- int \* de\_reject\_ct
- int \* gauss\_accept\_ct
- int \* gauss\_reject\_ct
- int \* mmala\_accept\_ct
- int \* mmala\_reject\_ct
- int \* RJstep\_accept\_ct
- int \* RJstep reject ct
- int \* swap\_accept\_ct
- int \* swap\_reject\_ct
- int \* step\_accept\_ct
- int \* step\_reject\_ctdouble \*\*\* II Ip output
- bool log\_II =false
- bool log\_lp =false
- int \* A
- bool PT\_alloc =false
- int \*\*\* param\_status
- bool RJMCMC =false
- std::function < void(double \*, double \*, int \*, int \*, int, int) > rj

# 8.24.1 Detailed Description

Class storing everything that defines an instance of the sampler

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

· include/mcmc sampler internals.h

# 8.25 source\_parameters < T > Struct Template Reference

# **Static Public Member Functions**

- static source\_parameters < T > populate\_source\_parameters (gen\_params \*param\_in)
   Builds the structure that shuttles source parameters between functions -updated version to incorporate structure argument.
- static source\_parameters < T > populate\_source\_parameters\_old (T mass1, T mass2, T Luminosity\_
   —
   Distance, T \*spin1, T \*spin2, T phi\_c, T t\_c, bool sky\_average)

Builds the structure that shuttles source parameters between functions- outdated in favor of structure argument.

## **Public Attributes**

- T mass1
- T mass2
- T M
- Tq
- T spin1z
- T spin2z
- T spin1x
- T spin2x
- T spin1y
- T spin2y
- T chirpmass
- T eta
- T chi s
- T chi\_a
- T chi eff
- T chi pn
- T DL
- T delta\_mass
- T fRD
- T fdamp
- T f1
- T f3
- T f1\_phase
- T f2\_phase
- T phic
- Ttc
- T A0
- Ts
- T chil
- T chip

- T f\_ref
- T phi\_aligned
- T incl angle
- T phiRef
- Talpha0
- T thetaJN
- T zeta\_polariz
- T \* betappe
- int \* bppe
- int Nmod
- T phi
- T theta
- T SP
- · TSL
- · bool sky average
- gsl\_spline \* **Z\_DL\_spline\_ptr** = NULL
- gsl\_interp\_accel \* **Z\_DL\_accel\_ptr** = NULL
- std::string cosmology

## 8.25.1 Member Function Documentation

# 8.25.1.1 populate\_source\_parameters()

Builds the structure that shuttles source parameters between functions -updated version to incorporate structure argument.

Populates the structure that is passed to all generation methods - contains all relavent source parameters

## 8.25.1.2 populate\_source\_parameters\_old()

Builds the structure that shuttles source parameters between functions- outdated in favor of structure argument.

Populates the structure that is passed to all generation methods - contains all relavent source parameters

## **Parameters**

mass1	mass of the larger body - in Solar Masses
mass2	mass of the smaller body - in Solar Masses
Luminosity_Distance	Luminosity Distance in Mpc
spin2	spin vector of the larger body {sx,sy,sz}
phi_c	spin vector of the smaller body {sx,sy,sz}
t_c	coalescence phase
sky_average	coalescence time

# 8.25.2 Member Data Documentation

```
template<class T>
T source_parameters< T >::chi_a
```

Antisymmetric spin combination

```
8.25.2.2 chi_eff
```

8.25.2.1 chi\_a

```
template<class T>
T source_parameters< T >::chi_eff
```

# Effective spin

# 8.25.2.3 chi\_pn

```
template<class T>
T source_parameters< T >::chi_pn
```

# PN spin

# 8.25.2.4 chi\_s

```
template<class T>
T source_parameters< T >::chi_s
```

# Symmetric spin combination

# 8.25.2.5 chirpmass

```
template<class T>
T source_parameters< T >::chirpmass
```

# Chirp mass of the binary

```
8.25.2.6 delta_mass
```

```
template<class T>
T source_parameters< T >::delta_mass
```

Delta mass comibination

## 8.25.2.7 DL

```
template<class T>
T source_parameters< T >::DL
```

Luminoisity Distance

## 8.25.2.8 eta

```
template<class T>
T source_parameters< T >::eta
```

Symmetric mass ratio

#### 8.25.2.9 f1

```
template<class T>
T source_parameters< T >::f1
```

Transition Frequency 1 for the amplitude

## 8.25.2.10 f1\_phase

```
template<class T>
T source_parameters< T >::fl_phase
```

Transition frequency 1 for the phase

# 8.25.2.11 f2\_phase

```
template<class T>
T source_parameters< T >::f2_phase
```

Transition frequency 2 for the phase

## 8.25.2.12 f3

```
template<class T>
T source_parameters< T >::f3
```

Transition Frequency 2 for the amplitude

```
8.25.2.13 fdamp
```

```
template<class T>
T source_parameters< T >::fdamp
```

Dampening frequency after merger

```
8.25.2.14 fRD
```

```
template<class T>
T source_parameters< T >::fRD
```

Ringdown frequency after merger

## 8.25.2.15 M

```
template<class T>
T source_parameters< T >::M
```

Total mass

#### 8.25.2.16 mass1

```
template<class T>
T source_parameters< T >::mass1
```

mass of the larger component

## 8.25.2.17 mass2

```
template<class T>
T source_parameters< T >::mass2
```

mass of the smaller component

# 8.25.2.18 Nmod

```
template<class T>
int source_parameters< T >::Nmod
```

Number of modifications to phase

# 8.25.2.19 phic

```
template<class T>
T source_parameters< T >::phic
```

# Coalescence phase

```
8.25.2.20 spin1x
template<class T>
T source_parameters< T >::spin1x
x-Spin component of the larger body
8.25.2.21 spin1y
template<class T>
T source_parameters< T >::spinly
y-Spin component of the larger body
8.25.2.22 spin1z
template<class T>
T source_parameters< T >::spin1z
z-Spin component of the larger body
8.25.2.23 spin2x
template<class T>
T source_parameters< T >::spin2x
x-Spin component of the smaller body
8.25.2.24 spin2y
{\tt template}{<}{\tt class} \ {\tt T}{>}
T source_parameters< T >::spin2y
y-Spin component of the smaller body
8.25.2.25 spin2z
template < class T >
T source_parameters< T >::spin2z
```

Generated by Doxygen

z-Spin component of the smaller body

# 8.25.2.26 tc

```
template<class T>
T source_parameters< T >::tc
```

## Coalescence time

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

- include/util.h
- src/util.cpp

# 8.26 sph\_harm< T > Struct Template Reference

# **Public Attributes**

- std::complex < T > Y22
- std::complex< T > Y21
- std::complex< T> Y20
- std::complex< T > Y2m1
- std::complex < T > Y2m2

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

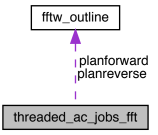
• include/util.h

# 8.27 threaded\_ac\_jobs\_fft Class Reference

Class to contain spectral method jobs.

```
#include <autocorrelation.h>
```

Collaboration diagram for threaded\_ac\_jobs\_fft:



# **Public Attributes**

- double \*\* data
- int \* length
- int \* start
- int \* end
- int dimension
- fftw\_outline \* planforward
- fftw\_outline \* planreverse
- int \* lag
- double \* target

# 8.27.1 Detailed Description

Class to contain spectral method jobs.

# 8.27.2 Member Data Documentation

# 8.27.2.1 dimension

int threaded\_ac\_jobs\_fft::dimension

Read only - end index

8.27.2.2 end

int\* threaded\_ac\_jobs\_fft::end

Read only - start index

8.27.2.3 lag

int\* threaded\_ac\_jobs\_fft::lag

fftw plan to use for spectral method

8.27.2.4 length

int\* threaded\_ac\_jobs\_fft::length

Read only - Data to use - full chain

## 8.27.2.5 planforward

```
fftw_outline* threaded_ac_jobs_fft::planforward
```

Read only - dimension being analyzed

# 8.27.2.6 planreverse

```
fftw_outline* threaded_ac_jobs_fft::planreverse
```

fftw plan to use for spectral method

# 8.27.2.7 start

```
int* threaded_ac_jobs_fft::start
```

Read only - length of total data

#### 8.27.2.8 target

```
double* threaded_ac_jobs_fft::target
```

# READ AND WRITE - final lag

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

• include/autocorrelation.h

# 8.28 threaded\_ac\_jobs\_serial Class Reference

Class to contain serial method jobs.

```
#include <autocorrelation.h>
```

# **Public Attributes**

- double \*\* data
- int \* length
- int \* start
- int \* end
- int dimension
- int \* lag
- double \* target

# 8.28.1 Detailed Description

Class to contain serial method jobs.

#### 8.28.2 Member Data Documentation

```
8.28.2.1 dimension
```

int threaded\_ac\_jobs\_serial::dimension

Read only – end index

8.28.2.2 end

int\* threaded\_ac\_jobs\_serial::end

Read only - start index

8.28.2.3 lag

int\* threaded\_ac\_jobs\_serial::lag

Read only - dimension being analyzed

8.28.2.4 length

int\* threaded\_ac\_jobs\_serial::length

Read only - Data to use - full chain

8.28.2.5 start

int\* threaded\_ac\_jobs\_serial::start

Read only - length of total data

8.28.2.6 target

double\* threaded\_ac\_jobs\_serial::target

READ AND WRITE - final lag

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

• include/autocorrelation.h

# 8.29 threadPool < jobtype, comparator > Class Template Reference

Class for creating a pool of threads to asynchronously distribute work.

```
#include <threadPool.h>
```

## **Public Member Functions**

threadPool (std::size\_t numThreads, std::function< void(int, jobtype)> work\_fn)

Constructor - starts thread pool running.

∼threadPool ()

Destructor - stops threads.

void enqueue (jobtype job\_id)

Places jobs in queue to wait for scheduling.

int get\_num\_threads ()

Get the number of threads being used by the thread pool.

• int get\_queue\_length ()

Get the current length of the job queue.

# 8.29.1 Detailed Description

```
{\tt template}{<} {\tt class\ jobtype=int,\ class\ comparator=default\_comp}{<} {\tt jobtype}{>}{>} {\tt class\ threadPool}{<} {\tt\ jobtype,\ comparator}{>}
```

Class for creating a pool of threads to asynchronously distribute work.

Template parameters:

jobtype defines a structure or class that represents a job or task

comparator defines how to compare jobs for sorting the list

Default options correspond to jobs being defined by an integer job\_id, and no sorting of the list (first in first out)

## 8.29.2 Member Function Documentation

## 8.29.2.1 enqueue()

Places jobs in queue to wait for scheduling.

job\_id is sorted if a comparator is provided

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

• include/threadPool.h

# 8.30 threadPool < jobtype, comparator > Class Template Reference

Class for creating a pool of threads to asynchronously distribute work.

```
#include <threadPool.h>
```

## **Public Member Functions**

threadPool (std::size\_t numThreads, std::function < void(int, jobtype) > work\_fn)

Constructor - starts thread pool running.

∼threadPool ()

Destructor - stops threads.

void enqueue (jobtype job\_id)

Places jobs in queue to wait for scheduling.

int get\_num\_threads ()

Get the number of threads being used by the thread pool.

• int get\_queue\_length ()

Get the current length of the job queue.

# 8.30.1 Detailed Description

```
{\tt template}{<} {\tt class\ jobtype=int,\ class\ comparator=default\_comp}{<} {\tt jobtype}{>}{>} {\tt class\ threadPool}{<} {\tt\ jobtype,\ comparator}{>}
```

Class for creating a pool of threads to asynchronously distribute work.

Template parameters:

jobtype defines a structure or class that represents a job or task

comparator defines how to compare jobs for sorting the list

Default options correspond to jobs being defined by an integer job\_id, and no sorting of the list (first in first out)

## 8.30.2 Member Function Documentation

## 8.30.2.1 enqueue()

Places jobs in queue to wait for scheduling.

job id is sorted if a comparator is provided

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

• include/threadPool.h

# 8.31 useful\_powers < T > Struct Template Reference

To speed up calculations within the for loops, we pre-calculate reoccuring powers of M\*F and Pi, since the pow() function is prohibatively slow.

```
#include <util.h>
```

# **Public Attributes**

- T MFthird
- T MFsixth
- T MF7sixth
- T MF2third
- T MF4third

- T MF7third
- T MFcube
- T MFminus 5third
- T MF3fourth
- double Plsquare
- · double Plcube
- · double Plthird
- double Pl2third
- double Pl4third
- double PI5third
- double PI7third
- double Plminus\_5third

# 8.31.1 Detailed Description

```
\label{eq:template} \begin{split} \text{template} &< \text{class T} > \\ \text{struct useful\_powers} &< \text{T} > \end{split}
```

To speed up calculations within the for loops, we pre-calculate reoccuring powers of M\*F and Pi, since the pow() function is prohibatively slow.

Powers of PI are initialized once, and powers of MF need to be calculated once per for loop (if in the inspiral portion).

use the functions precalc\_powers\_ins\_amp, precalc\_powers\_ins\_phase, precalc\_powers\_pi to initialize

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

• include/util.h

# **Chapter 9**

# **File Documentation**

# 9.1 gw\_analysis\_tools\_py/src/mcmc\_routines\_ext.pyx File Reference

File that wraps the code in mcmc\_gw.cpp, mcmc\_sampler.cpp, mcmc\_sampler\_internals.cpp, autocorrelation.cpp.

#### Classes

· class mcmc\_routines\_ext.fftw\_outline\_py

## **Functions**

- def mcmc\_routines\_ext.write\_auto\_corr\_file\_from\_data\_file\_py (string, autocorr\_filename, string, datafile, int, length, int, dimension, int, num\_segments, double, target\_corr, int, num\_threads)
- def mcmc\_routines\_ext.arange (string, autocorr\_filename, :1] data, int length, int dimension, int num\_
   segments, double target\_corr, int num\_threads):#Not ideal -- have to wrap the memview in a real c++array cdef double \*\*temparr=< double \*\* > malloc(sizeof(double \*double, length)
- def mcmc\_routines\_ext.allocate\_FFTW\_mem\_forward\_py (fftw\_outline\_py, plan, int, length)
- def mcmc\_routines\_ext.deallocate\_FFTW\_mem\_py (fftw\_outline\_py, plan)

# 9.1.1 Detailed Description

File that wraps the code in mcmc\_gw.cpp, mcmc\_sampler.cpp, mcmc\_sampler\_internals.cpp, autocorrelation.cpp.

# 9.2 gw\_analysis\_tools\_py/src/waveform\_generator\_ext.pyx File Reference

File that contains cython code to wrap the c++ library.

## Classes

class waveform\_generator\_ext.gen\_params\_py

Python wrapper for the generation parameters structure, as defined in util.cpp.

## **Namespaces**

· waveform generator ext

Python wrapper for the waveform generation in waveform\_generator.cpp.

## **Functions**

- def waveform generator ext.double (self, double, mass1, double, mass2, double, DL, spin1, spin2, double, phic, double, tc, :1] bppe, double[::1] betappe, int Nmod, double theta, double phi, double incl angle, double f\_ref, double phiRef, bool NSflag):self.params.mass1=mass1 self.params.mass2=mass2 self.← params.Luminosity Distance=DL self.params.spin1=spin1 self.params.spin2=spin2 self.params.phic=phic self.params.tc=tc self.params.bppe=&bppe[0] self.params.betappe=&betappe[0] self.params.Nmod=Nmod self.params.incl\_angle=incl\_angle self.params.theta=theta self.params.phi=phi self.params.f\_ref=f\_ref self.← params.phiRef=phiRef self.params.NSflag=NSflag ##Computes the waveform in Fourier space # @param frequencies The array of frequencies to use # @param generation\_method Method to use for the waveform generation # @param gen params py Parameters of the binary def fourier waveform py(double[::1] frequencies, string generation method, gen params py parameters):cdef double[::1] waveform real=np.← ascontiguousarray(np.zeros((frequencies.size) int, dtype=np.float64)
- def waveform generator ext.double (:1] frequencies, string generation method, gen params ← \_py parameters):cdef double[::1] amplitude=np.ascontiguousarray(np.zeros((frequencies.size) double, dtype=np.float64, frequencies, frequencies, size, amplitude, generation\_method, parameters, params, :1] frequencies, string generation\_method, gen\_params\_py parameters):cdef double[::1] phase=np.← ascontiguousarray(np.zeros((frequencies.size) double, dtype=np.float64, frequencies, frequencies, size, phase, generation\_method, parameters, params, :1] frequencies, string generation\_method, gen\_params\_py parameters):cdef double[::1] waveform plus real=np.ascontiguousarray(np.zeros((frequencies.size) double, dtvpe=np.float64)

## **Variables**

- · waveform generator ext.complex128 t
- · waveform generator ext.ndim
- · waveform generator ext.waveform
- waveform generator ext.dtvpe
- waveform\_generator\_ext.i = i +1

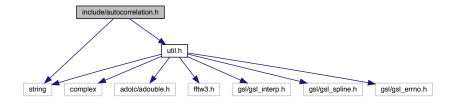
#### 9.2.1 **Detailed Description**

File that contains cython code to wrap the c++ library.

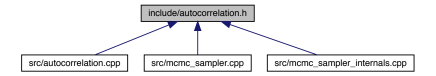
#### include/autocorrelation.h File Reference 9.3

```
#include <string>
#include "util.h"
```

Include dependency graph for autocorrelation.h:



This graph shows which files directly or indirectly include this file:



## Classes

· class threaded ac jobs fft

Class to contain spectral method jobs.

class threaded\_ac\_jobs\_serial

Class to contain serial method jobs.

class comparator\_ac\_fft

comparator to sort ac-jobs

class comparator\_ac\_serial

comparator to sort ac-jobs

## **Functions**

- void write\_auto\_corr\_file\_from\_data\_file (std::string autocorr\_filename, std::string datafile, int length, int dimension, int num\_segments, double target\_corr, int num\_threads)
- void write\_auto\_corr\_file\_from\_data (std::string autocorr\_filename, double \*\*data, int length, int dimension, int num\_segments, double target\_corr, int num\_threads)

Writes the autocorrelation file from a data array.

• void auto\_corr\_from\_data (double \*\*data, int length, int dimension, int \*\*output, int num\_segments, double target\_corr, int num\_threads)

Calculates the autocorrelation length for a set of data for a number of segments for each dimension – completely host code, utilitizes FFTW3 for longer chuncks of the chains.

void threaded \_ac\_spectral (int thread, threaded\_ac\_jobs\_fft job)

Internal routine to calculate an spectral autocorrelation job.

void threaded\_ac\_serial (int thread, threaded\_ac\_jobs\_serial job)

Internal routine to calculate an serial autocorrelation job.

• double auto\_correlation\_serial (double \*arr, int length, int start, double target)

Calculates the autocorrelation of a chain with the brute force method.

void auto\_correlation\_spectral (double \*chain, int length, double \*autocorr, fftw\_outline \*plan\_forw, fftw\_outline \*plan\_rev)

Wrapper function for convience – assumes the data array starts at 0.

• void auto\_correlation\_spectral (double \*chain, int length, int start, double \*autocorr, fftw\_outline \*plan\_forw, fftw\_outline \*plan\_rev)

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT – accepts FFTW plan as argument for plan reuse and multi-threaded applications.

void auto correlation spectral (double \*chain, int length, double \*autocorr)

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT.

• double auto correlation (double \*arr, int length, double tolerance)

OUTDATED - numerically finds autocorrelation length - not reliable.

double auto\_correlation\_serial\_old (double \*arr, int length)

OUTDATED Calculates the autocorrelation – less general version.

• double auto\_correlation\_grid\_search (double \*arr, int length, int box\_num=10, int final\_length=50, double target\_length=.01)

OUTDATED - Grid search method of computing the autocorrelation - unreliable.

double auto\_correlation\_internal (double \*arr, int length, int lag, double ave)

Internal function to compute the auto correlation for a given lag.

void auto\_corr\_intervals\_outdated (double \*data, int length, double \*output, int num\_segments, double accuracy)

Function that computes the autocorrelation length on an array of data at set intervals to help determine convergence.

void write\_auto\_corr\_file\_from\_data (std::string autocorr\_filename, double \*\*output, int intervals, int dimension, int N steps)

OUTDATED – writes autocorrelation lengths for a data array, but only with the serial method and only for a target correlation of .01.

void write\_auto\_corr\_file\_from\_data\_file (std::string autocorr\_filename, std::string output\_file, int intervals, int dimension, int N\_steps)

OUTDATED – writes autocorrelation lengths for a data file, but only with the serial method and only for a target correlation of .01.

# 9.3.1 Detailed Description

Autocorrelation header file

## 9.3.2 Function Documentation

#### 9.3.2.1 auto\_corr\_from\_data()

Calculates the autocorrelation length for a set of data for a number of segments for each dimension – completely host code, utilitizes FFTW3 for longer chuncks of the chains.

Takes in the data from a sampler, shape data[N\_steps][dimension]

Outputs lags that correspond to the target\_corr - shape output[dimension][num\_segments]

#### **Parameters**

	data	Input data
	length	length of input data
	dimension	dimension of data
out	output	array that stores the auto-corr lengths – array[num_segments]
	num_segments	number of segements to compute the auto-corr length
	target_corr	Autocorrelation for which the autocorrelation length is defined (lag of autocorrelation
		for which it equals the target_corr)
	num_threads	Total number of threads to use

## 9.3.2.2 auto\_corr\_intervals\_outdated()

Function that computes the autocorrelation length on an array of data at set intervals to help determine convergence.

outdated version - new version uses FFTs

## **Parameters**

	data	Input data
	length	length of input data
out	output	array that stores the auto-corr lengths – array[num_segments]
	num_segments	number of segements to compute the auto-corr length
	accuracy	longer chains are computed numerically, this specifies the tolerance

## 9.3.2.3 auto\_correlation\_grid\_search()

OUTDATED - Grid search method of computing the autocorrelation - unreliable.

Hopefully more reliable than the box-search method, which can sometimes get caught in a recursive loop when the stepsize isn't tuned, but also faster than the basic linear, serial search

## **Parameters**

arr	Input array to use for autocorrelation	
length	Length of input array	
box_num	number of boxes to use for each iteration, default is 10	
final_length number of elements per box at which the grid search ends and the serial calculation begin		
target_length	target correlation that corresponds to the returned lag	

## 9.3.2.4 auto\_correlation\_internal()

Internal function to compute the auto correlation for a given lag.

## 9.3.2.5 auto\_correlation\_serial()

Calculates the autocorrelation of a chain with the brute force method.

#### **Parameters**

arr	input array	
length	Length of input array	
start	starting index (probably 0)	
target	Target autocorrelation for which `'length'' is defined	

# 9.3.2.6 auto\_correlation\_spectral() [1/2]

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT – accepts FFTW plan as argument for plan reuse and multi-threaded applications.

Based on the Wiener-Khinchin Theorem.

Algorithm used from https://lingpipe-blog.com/2012/06/08/autocorrelation-fft-kiss-eigen/

NOTE the length used in initializing the fftw plans should be L = pow(2, std::ceil( std::log2(length) ) ) – the plans are padded so the total length is a power of two

Option to provide starting index for multi-dimension arrays in collapsed to one dimension

length is the length of the segment to be analyzed, not necessarily the dimension of the chain

## 9.3.2.7 auto\_correlation\_spectral() [2/2]

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT.

Based on the Wiener-Khinchin Theorem.

Algorithm used from https://lingpipe-blog.com/2012/06/08/autocorrelation-fft-kiss-eigen/

#### 9.3.2.8 threaded\_ac\_serial()

Internal routine to calculate an serial autocorrelation job.

Allows for a more efficient use of the threadPool class

## 9.3.2.9 threaded\_ac\_spectral()

Internal routine to calculate an spectral autocorrelation job.

Allows for a more efficient use of the threadPool class

# 9.3.2.10 write\_auto\_corr\_file\_from\_data()

```
void write_auto_corr_file_from_data (
    std::string autocorr_filename,
    double ** data,
    int length,
    int dimension,
    int num_segments,
    double target_corr,
    int num_threads )
```

Writes the autocorrelation file from a data array.

## **Parameters**

i didilictors		
autocorr_filename	Name of the file to write the autocorrelation to	
data	Input chains	
length	length of input data	
dimension	dimension of data	
Generaledsby Prygres	number of segements to compute the auto-corr length	
target_corr	Autocorrelation for which the autocorrelation length is defined (lag of autocorrelation for which it equals the target_corr)	
num_threads	Total number of threads to use	

## 9.3.2.11 write\_auto\_corr\_file\_from\_data\_file()

```
void write_auto_corr_file_from_data_file (
    std::string autocorr_filename,
    std::string datafile,
    int length,
    int dimension,
    int num_segments,
    double target_corr,
    int num_threads )
```

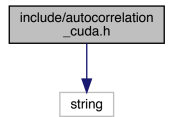
#### **Parameters**

length	length of input data	
dimension	dimension of data	
num_segments	number of segements to compute the auto-corr length	
target_corr	Autocorrelation for which the autocorrelation length is defined (lag of autocorrelation for which it equals the target_corr)	
num_threads	Total number of threads to use	

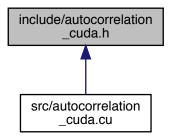
# 9.4 include/autocorrelation\_cuda.h File Reference

```
#include <string>
```

Include dependency graph for autocorrelation\_cuda.h:



This graph shows which files directly or indirectly include this file:



## **Macros**

• #define THREADS\_PER\_BLOCK 512

#### **Functions**

void write\_file\_auto\_corr\_from\_data\_file\_accel (std::string acfile, std::string chains\_file, int dimension, int N
 \_ steps, int num\_segments, double target\_corr)

Write data file for autocorrelation lengths of the data given a data file name, as written by the mcmc\_sampler.

• void write\_file\_auto\_corr\_from\_data\_accel (std::string acfile, double \*\*output, int dimension, int N\_steps, int num\_segments, double target\_corr)

Write data file given output chains, as formatted by the mcmc\_sampler.

• void <a href="mailto:auto\_corr\_from\_data\_accel">accel</a> (double \*\*output, int dimension, int N\_steps, int num\_segments, double target\_corr, double \*\*autocorr)

Find autocorrelation of data at different points in the chain length and output to autocorr.

void launch\_ac\_gpu (int device, int element, double \*\*data, int length, int dimension, double target\_corr, int num\_segments)

Launch the GPU kernel, formatted for the thread pool.

• void ac\_gpu\_wrapper (int thread, int job\_id)

Wrapper function for the thread pool.

· void auto\_correlation\_spectral\_accel (double \*chains, int length, double \*autocorr)

# 9.4.1 Detailed Description

Header file for CUDA accelerated algorithms

Currently, no algorithms are used in any other parts of the project, so if CUDA or CUDA-enabled devices are not available, this file can be skipped in compilation by commenting out the OBJECTSCUDA line in the makefile

## 9.4.2 Function Documentation

## 9.4.2.1 ac\_gpu\_wrapper()

Wrapper function for the thread pool.

# **Parameters**

thread	Host thread
job⊷	Job ID
_id	

# 9.4.2.2 auto\_corr\_from\_data\_accel()

Find autocorrelation of data at different points in the chain length and output to autocorr.

## **Parameters**

	output	Chain data input
	dimension	Dimension of the data
	N_steps	Number of steps in the data
	num_segments	number of segments to calculate the autocorrelation length
	target_corr	Target correlation ratio
out	autocorr	Autocorrelation lengths for the different segments

# 9.4.2.3 write\_file\_auto\_corr\_from\_data\_accel()

```
void write_file_auto_corr_from_data_accel (
    std::string acfile,
    double ** output,
    int dimension,
    int N_steps,
    int num_segments,
    double target_corr )
```

Write data file given output chains, as formatted by the mcmc\_sampler.

# **Parameters**

acfile	Output autocorrelation filename	
output	Chain data from MCMC_sampler	
dimension	Dimension of the data	
N_steps	Number of steps in the chain	
num_segments	Number of segments to check the autocorrelation length for each dimension	
target_corr	Target correlation ratio to use for the correlation length calculation	

# 9.4.2.4 write\_file\_auto\_corr\_from\_data\_file\_accel()

```
void write_file_auto_corr_from_data_file_accel (
    std::string acfile,
    std::string chains_file,
    int dimension,
    int N_steps,
    int num_segments,
    double target_corr )
```

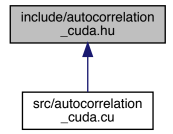
Write data file for autocorrelation lengths of the data given a data file name, as written by the mcmc\_sampler.

#### **Parameters**

acfile	Filename of the autocorrelation data	
chains_file	Filename of the data file for the chains	
dimension	Dimension of the data	
N_steps	Number of steps in the chain	
num_segments	Number of segments to check the autocorrelation length for each dimension	
target_corr	Target correlation ratio to use for the correlation length calculation	

# 9.5 include/autocorrelation\_cuda.hu File Reference

This graph shows which files directly or indirectly include this file:



## Classes

• struct GPUplan

#### **Functions**

• \_\_device\_ \_\_host\_\_ void auto\_corr\_internal (double \*arr, int length, int lag, double average, double \*corr, int start id)

Internal function to calculate the autocorrelation for a given lag Customized for the thread pool architecture, with extra arguments because of the way the memory is allocated.

• \_\_global\_\_ void auto\_corr\_internal\_kernal (double \*arr, int length, double average, int \*rho\_index, double target\_corr, double var, int start\_id)

Internal function to launch the CUDA kernel for a range of autocorrelations.

- void allocate\_gpu\_plan (GPUplan \*plan, int data\_length, int dimension, int num\_segments)
  - Allocates memory for autocorrelation-GPU structure.
- void deallocate\_gpu\_plan (GPUplan \*plan, int data\_length, int dimension, int num\_segments)

Deallocates memory for the autocorrelation-GPU structure.

Copy data to device before starting kernels.

## 9.5.1 Function Documentation

#### 9.5.1.1 allocate\_gpu\_plan()

Allocates memory for autocorrelation–GPU structure.

## **Parameters**

plan	Structure for GPU plan
data_length	Length of data
dimension	Dimension of the data
num_segments	Number of segments to calculate the autocorrelation length

# 9.5.1.2 auto\_corr\_internal()

```
int length,
int lag,
double average,
double * corr,
int start_id )
```

Internal function to calculate the autocorrelation for a given lag Customized for the thread pool architecture, with extra arguments because of the way the memory is allocated.

## **Parameters**

	arr	Input array of data
	length	Length of input array
	lag	Lag to be used to calculate the correlation
	average	Average of the array arr
out	corr	output correlation
	start_id	ID of location to start calculation – input arrary arr is assumed to be contiguous for multiple dimensions

## 9.5.1.3 auto\_corr\_internal\_kernal()

Internal function to launch the CUDA kernel for a range of autocorrelations.

# Correlation function used:

```
 rho(lag) = 1 \ / \ (length - lag) \ (arr[i+lag]-average) \ (arr[i]-average) \\ target\_corr = rho(rho\_index)/rho(0) = rho(rho\_index)/var
```

# **Parameters**

	arr	Input array of data
	length	Length of data array
	average	Average of input data
out	rho_index	Index of the lag that results ina correlation ratio target_corr
	target_corr	Target correlation ratio rho(lag)/rho(0) = target_corr
	var	Variance rho(0)
	start_id	Starting index to use for the data array arr

# 9.5.1.4 copy\_data\_to\_device()

Copy data to device before starting kernels.

## **Parameters**

plan	GPU plan
input_data	Input chain data
data_length	Length of data
dimension	Dimension of the data
num_segments	Number of segments to calculate the autocorrelation length

# 9.5.1.5 deallocate\_gpu\_plan()

Deallocates memory for the autocorrelation-GPU structure.

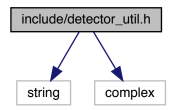
## **Parameters**

plan	Structure for the GPU plan
data_length	Length of data
dimension	Dimension of the data
num_segments	Number of segments to calculate the autocorrelation length

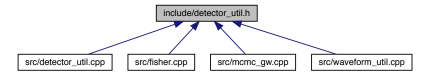
# 9.6 include/detector\_util.h File Reference

```
#include <string>
#include <complex>
```

Include dependency graph for detector\_util.h:



This graph shows which files directly or indirectly include this file:



## **Functions**

- void populate\_noise (double \*frequencies, std::string detector, double \*noise\_root, int length=0)
  - Function to populate the squareroot of the noise curve for various detectors.
- double aLIGO\_analytic (double f)
  - Analytic function approximating the PSD for aLIGO.
- $\operatorname{std}::\operatorname{complex} < \operatorname{double} > \operatorname{Q}$  (double theta, double phi, double iota)
  - Utility for the overall amplitude and phase shift for spin-aligned systems.
- double right\_interferometer\_cross (double theta, double phi)
  - Response function of a 90 deg interferometer for cross polarization.
- double right interferometer plus (double theta, double phi)
  - Response function of a 90 deg interferometer for plus polarization.
- double Hanford O1 fitted (double f)
  - Numerically fit PSD to the Hanford Detector's O1.
- void celestial\_horizon\_transform (double RA, double DEC, double gps\_time, std::string detector, double \*phi, double \*theta)
  - Transform from celestial coordinates to local horizontal coords.
- void derivative\_celestial\_horizon\_transform (double RA, double DEC, double gps\_time, std::string detector, double \*dphi\_dRA, double \*dtheta\_dRA, double \*dphi\_dDEC, double \*dtheta\_dDEC)
  - Numerical derivative of the transformation.
- double DTOA (double theta1, double theta2, std::string detector1, std::string detector2)
  - calculate difference in time of arrival (DTOA) for a given source location and 2 different detectors
- double radius\_at\_lat (double latitude, double elevation)

• void detector\_response\_functions\_equatorial (double D[3][3], double ra, double dec, double psi, double gmst, double \*Fplus, double \*Fcross)

Calculates the response coefficients for a detector with response tensor D for a source at RA, Dec, and psi.

• void detector\_response\_functions\_equatorial (std::string detector, double ra, double dec, double psi, double gmst, double \*Fplus, double \*Fcross)

Same as the other function, but for active detectors.

## **Variables**

- const double H LAT = 0.81079526383
- const double H LONG =-2.08405676917
- const double **H\_azimuth\_offset** = 2.199
- const double **H\_radius** = 6367299.93401105
- const double **H\_elevation** = 142.554
- const double L LAT = 0.53342313506
- const double L\_LONG =-1.58430937078
- const double L\_azimuth\_offset = 3.4557
- const double L\_radius = 6372795.50144497
- const double L\_elevation = -6.574
- const double V\_LAT = 0.76151183984
- const double V\_LONG =0.18333805213
- const double **V\_azimuth\_offset** = 1.239
- const double V\_radius = 6368051.92301
- const double V\_elevation = 51.884
- const double **RE\_polar** =6357e3
- const double **RE\_equatorial** = 6378e3
- const double Hanford\_D [3][3]
- const double Livingston\_D [3][3]
- const double Virgo\_D [3][3]

# 9.6.1 Detailed Description

Header file for all detector-specific utilities

### 9.6.2 Function Documentation

# 9.6.2.1 aLIGO\_analytic()

```
double aLIGO_analytic ( double f )
```

Analytic function approximating the PSD for aLIGO.

CITE (Will?)

## 9.6.2.2 celestial\_horizon\_transform()

Transform from celestial coordinates to local horizontal coords.

```
(RA,DEC) -> (altitude, azimuth)
```

Need gps\_time of transformation, as the horizontal coords change in time

detector is used to specify the lat and long of the local frame

## **Parameters**

RA	in RAD
DEC	in RAD
phi	in RAD
theta	in RAD

## 9.6.2.3 derivative\_celestial\_horizon\_transform()

Numerical derivative of the transformation.

Planned for use in Fisher calculations, but not currently implemented anywhere

#### **Parameters**

RA	in RAD
DEC	in RAD

# 9.6.2.4 detector\_response\_functions\_equatorial() [1/2]

```
void detector_response_functions_equatorial (
```

```
double D[3][3],
double ra,
double dec,
double psi,
double gmst,
double * Fplus,
double * Fcross )
```

Calculates the response coefficients for a detector with response tensor D for a source at RA, Dec, and psi.

## Taken from LALSuite

The response tensor for each of the operational detectors is precomputed in detector\_util.h, but to create a new tensor, follow the outline in Anderson et al 36 PRD 63 042003 (2001) Appendix B

## **Parameters**

	D	Detector Response tensor (3x3)
	ra	Right ascension in rad
	dec	Declination in rad
	psi	polarization angle in rad
	gmst	Greenwich mean sidereal time (rad)
out	Fplus	Fplus response coefficient
out	Fcross	Fcross response coefficient

# 9.6.2.5 detector\_response\_functions\_equatorial() [2/2]

```
void detector_response_functions_equatorial (
    std::string detector,
    double ra,
    double dec,
    double psi,
    double gmst,
    double * Fplus,
    double * Fcross )
```

Same as the other function, but for active detectors.

	detector	Detector
	ra	Right ascension in rad
	dec	Declination in rad
	psi	polarization angle in rad
	gmst	Greenwich mean sidereal time (rad)
out	Fplus	Fplus response coefficient
out	Fcross	Fcross response coefficient

## 9.6.2.6 DTOA()

calculate difference in time of arrival (DTOA) for a given source location and 2 different detectors

#### **Parameters**

theta1	spherical polar angle for detector 1 in RAD
theta2	spherical polar angle for detector 2 in RAD
detector1	name of detector one
detector2	name of detector two

## 9.6.2.7 Hanford\_O1\_fitted()

```
double Hanford_O1_fitted ( double f )
```

Numerically fit PSD to the Hanford Detector's O1.

CITE (Yunes?)

# 9.6.2.8 populate\_noise()

Function to populate the squareroot of the noise curve for various detectors.

If frequencies are left as NULL, standard frequency spacing is applied and the frequencies are returned, in which case the frequencies argument becomes an output array

Detector names must be spelled exactly

Detectors include: aLIGO\_analytic, Hanford\_O1\_fitted

frequencies	double array of frquencies (NULL)
detector	String to designate the detector noise curve to be used
noise_root	ouptput double array for the square root of the PSD of the noise of the specified detector
length	integer length of the output and input arrays

#### 9.6.2.9 Q()

Utility for the overall amplitude and phase shift for spin-aligned systems.

For spin aligned, all the extrinsic parameters have the effect of an overall amplitude modulation and phase shift

# 9.6.2.10 radius\_at\_lat()

/brief Analytic approximation of the radius from the center of earth to a given location

Just the raidus as a function of angles, modelling an oblate spheroid

#### **Parameters**

latitude	latitude in degrees
elevation	elevation in meters

# 9.6.2.11 right\_interferometer\_cross()

```
double right_interferometer_cross ( \label{eq:cross} \mbox{double } theta, \\ \mbox{double } phi \mbox{ )}
```

Response function of a 90 deg interferometer for cross polarization.

Theta and phi are local, horizontal coordinates relative to the detector

# 9.6.2.12 right\_interferometer\_plus()

Response function of a 90 deg interferometer for plus polarization.

Theta and phi are local, horizontal coordinates relative to the detector

# 9.6.3 Variable Documentation

#### 9.6.3.1 Hanford D

const double Hanford\_D[3][3]

## Initial value:

```
= {{-0.392632, -0.0776099, -0.247384}, {-0.0776099, 0.319499, 0.227988}, {-0.247384, 0.227988, 0.0730968}}
```

# 9.6.3.2 Livingston\_D

const double Livingston\_D[3][3]

## Initial value:

```
= {{0.411318, 0.14021, 0.247279}, {0.14021, -0.108998, -0.181597}, {0.247279, -0.181597, -0.302236}}
```

# 9.6.3.3 Virgo\_D

const double Virgo\_D[3][3]

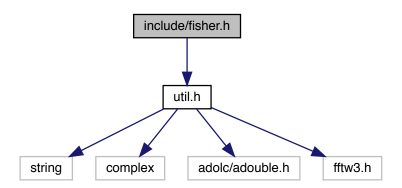
# Initial value:

```
= {{0.243903, -0.0990959, -0.232603}, {-0.0990959, -0.447841, 0.187841}, {-0.232603, 0.187841, 0.203979}}
```

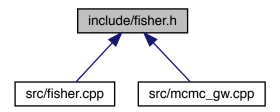
# 9.7 include/fisher.h File Reference

```
#include "util.h"
```

Include dependency graph for fisher.h:



This graph shows which files directly or indirectly include this file:



## **Functions**

• void fisher (double \*frequency, int length, string generation\_method, string detector, double \*\*output, int dimension, gen\_params \*parameters, int \*amp\_tapes=NULL, int \*phase\_tapes=NULL, double \*noise=N ∪ ULL)

Calculates the fisher matrix for the given arguments.

• void calculate\_derivatives (double \*\*amplitude\_deriv, double \*\*phase\_deriv, double \*amplitude, double \*frequencies, int length, string detector, string gen\_method, gen\_params \*parameters)

Abstraction layer for handling the case separation for the different waveforms.

 void fisher\_autodiff (double \*frequency, int length, string generation\_method, string detector, double \*\*output, int dimension, gen\_params \*parameters, int \*amp\_tapes=NULL, int \*phase\_tapes=NULL, double \*noise=NULL)

Calculates the fisher matrix for the given arguments to within numerical error using automatic differention - slower than the numerical version.

## 9.7.1 Function Documentation

# 9.7.1.1 calculate\_derivatives()

Abstraction layer for handling the case separation for the different waveforms.

## 9.7.1.2 fisher()

Calculates the fisher matrix for the given arguments.

## **Parameters**

length	if 0, standard frequency range for the detector is used
output	double [dimension][dimension]
amp_tapes	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method
phase_tapes	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method

# 9.7.1.3 fisher\_autodiff()

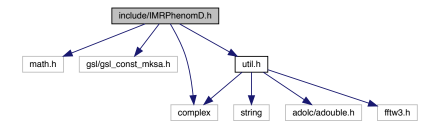
Calculates the fisher matrix for the given arguments to within numerical error using automatic differention - slower than the numerical version.

length	if 0, standard frequency range for the detector is used
output	double [dimension][dimension]
amp_tapes	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method
phase_tapes  Generated by Doxyge	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method

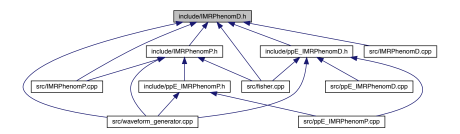
# 9.8 include/IMRPhenomD.h File Reference

```
#include <math.h>
#include <gsl/gsl_const_mksa.h>
#include <complex>
#include "util.h"
```

Include dependency graph for IMRPhenomD.h:



This graph shows which files directly or indirectly include this file:



# Classes

- struct lambda\_parameters < T >
- class IMRPhenomD< T >

# **Variables**

• const double lambda\_num\_params [19][11]

# 9.8.1 Detailed Description

Header file for utilities

# 9.8.2 Variable Documentation

## 9.8.2.1 lambda\_num\_params

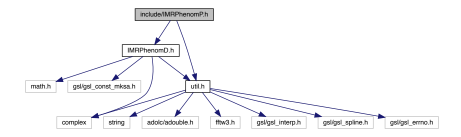
```
const double lambda_num_params[19][11]
```

Numerically calibrated parameters from arXiv:1508.07253 see the table in the data directory for labeled version

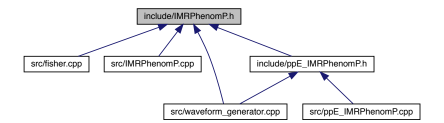
# 9.9 include/IMRPhenomP.h File Reference

```
#include "IMRPhenomD.h"
#include "util.h"
```

Include dependency graph for IMRPhenomP.h:



This graph shows which files directly or indirectly include this file:



# Classes

- struct alpha\_coeffs< T >
- struct epsilon coeffs< T >
- class IMRPhenomPv2< T >

# 9.9.1 Detailed Description

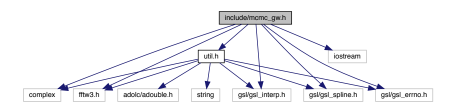
Header file for IMRPhenomP functions

Currently, only Pv2 is supported.

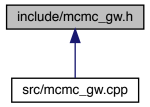
Wrapped around IMRPhenomD

# 9.10 include/mcmc\_gw.h File Reference

```
#include <complex>
#include <fftw3.h>
#include "util.h"
#include <iostream>
#include <gsl/gsl_interp.h>
#include <gsl/gsl_spline.h>
#include <gsl/gsl_errno.h>
Include dependency graph for mcmc gw.h:
```



This graph shows which files directly or indirectly include this file:



## **Functions**

• double maximized\_coal\_log\_likelihood\_IMRPhenomD (double \*frequencies, int length, std::complex< double > \*data, double \*noise, double SNR, double chirpmass, double symmetric\_mass\_ratio, double spin1, double spin2, bool NSflag, fftw\_outline \*plan)

Function to calculate the log Likelihood as defined by -1/2 (d-h|d-h) maximized over the extrinsic parameters phic and tc.

- double maximized\_coal\_log\_likelihood\_IMRPhenomD (double \*frequencies, size\_t length, double \*real\_
   data, double \*imag\_data, double \*noise, double SNR, double chirpmass, double symmetric\_mass\_ratio,
   double spin1, double spin2, bool NSflag)
- double maximized\_coal\_log\_likelihood\_IMRPhenomD (double \*frequencies, size\_t length, double \*real\_
   data, double \*imag\_data, double \*noise, double SNR, double chirpmass, double symmetric\_mass\_ratio,
   double spin1, double spin2, bool NSflag, fftw\_outline \*plan)
- double maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param (double \*frequencies, int length, std
   ::complex< double > \*data, double \*noise, double chirpmass, double symmetric\_mass\_ratio, double spin1,
   double spin2, double Luminosity\_Distance, double theta, double phi, double iota, bool NSflag, fftw\_outline
   \*plan)

- double maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param (double \*frequencies, size\_t length, double \*real\_data, double \*imag\_data, double \*noise, double chirpmass, double symmetric\_mass\_ratio, double spin1, double spin2, double Luminosity\_Distance, double theta, double phi, double iota, bool NSflag)
- double maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param (double \*frequencies, size\_t length, double \*real\_data, double \*imag\_data, double \*noise, double chirpmass, double symmetric\_mass\_ratio, double spin1, double spin2, double Luminosity\_Distance, double theta, double phi, double iota, bool NSflag, fftw outline \*plan)
- double maximized\_Log\_Likelihood\_aligned\_spin\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std::complex< double > \*detector\_response, size\_t length, fftw\_outline \*plan)

Maximized match over coalescence variables - returns log likelihood NOT NORMALIZED for aligned spins.

 double Log\_Likelihood (std::complex< double > \*data, double \*psd, double \*frequencies, size\_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw\_outline \*plan)

Unmarginalized log of the likelihood.

double maximized\_Log\_Likelihood\_unaligned\_spin\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std::complex< double > \*hplus, std::complex< double > \*hcross, size\_t length, fftw outline \*plan)

log likelihood function that maximizes over extrinsic parameters tc, phic, D, and phiRef, the reference frequency - for unaligned spins

double maximized\_Log\_Likelihood (std::complex < double > \*data, double \*psd, double \*frequencies, size ←
 \_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw\_outline \*plan)

routine to maximize over all extrinsic quantities and return the log likelihood

- double **maximized\_Log\_Likelihood** (double \*data\_real, double \*data\_imag, double \*psd, double \*frequencies, size\_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw outline \*plan)
- double maximized\_coal\_Log\_Likelihood (std::complex < double > \*data, double \*psd, double \*frequencies, size\_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw\_outline \*plan, double \*tc, double \*phic)

Function to maximize only over coalescence variables to and phic, returns the maximum values used.

- double maximized\_coal\_Log\_Likelihood\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std::complex< double > \*detector\_response, size\_t length, fftw\_outline \*plan, double \*tc, double \*phic)
- double Log\_Likelihood\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std
   ::complex< double > \*detector response, int length, fftw outline \*plan)

Internal function for the unmarginalized log of the likelihood.

void PTMCMC\_MH\_GW (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension, int chain\_id), int numThreads, bool pool, bool show\_prog, int num\_detectors, std::complex< double > \*\*data, double \*\*noise\_psd, double \*\*frequencies, int \*data\_length, double gps\_time, std::string \*detector, int Nmod, int \*bppe, std::string generation\_method, std::string statistics\_filename, std::string chain\_filename, std::string auto corr filename, std::string likelihood log filename, std::string checkpoint filename)

Wrapper for the MCMC\_MH function, specifically for GW analysis.

void PTMCMC\_MH\_dynamic\_PT\_alloc\_GW (double \*\*\*output, int dimension, int N\_steps, int chain\_N, int max\_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swpc\_freq, int t0, int nu, std::string chain\_distribution\_scheme, double(\*log\_prior)(double \*param, int dimension, int chain\_id), int numThreads, bool pool, bool show\_prog, int num\_detectors, std::complex< double > \*\*data, double \*\*noise\_psd, double \*\*frequencies, int \*data\_length, double gps\_time, std::string \*detectors, int Nmod, int \*bppe, std::string generation\_method, std::string statistics\_filename, std::string chain\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_filename)

Takes in an MCMC checkpoint file and continues the chain.

void continue\_PTMCMC\_MH\_GW (std::string start\_checkpoint\_file, double \*\*\*output, int dimension, int N←
 \_steps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension, int chain\_id), int numThreads, bool
 pool, bool show\_prog, int num\_detectors, std::complex< double > \*\*data, double \*\*noise\_psd, double
 \*\*frequencies, int \*data\_length, double gps\_time, std::string \*detector, int Nmod, int \*bppe, std::string
 generation\_method, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename,
 std::string likelihood\_log\_filename, std::string final\_checkpoint\_filename)

Takes in an MCMC checkpoint file and continues the chain.

void PTMCMC\_method\_specific\_prep (std::string generation\_method, int dimension, double \*seeding\_var, bool local\_seeding)

Unpacks MCMC parameters for method specific initiation.

- double MCMC\_likelihood\_extrinisic (bool save\_waveform, gen\_params \*parameters, std::string generation\_method, int \*data\_length, double \*\*frequencies, std::complex< double > \*\*data, double \*\*psd, std::string \*detectors, fftw\_outline \*fftw\_plans, int num\_detectors, double RA, double DEC, double gps time)
- $\bullet \ \ void \ \ \ \ MCMC\_fisher\_wrapper \ (double *param, int \ dimension, \ double **output, int \ chain\_id)\\$

Fisher function for MCMC for GW.

• double MCMC\_likelihood\_wrapper (double \*param, int dimension, int chain\_id)

log likelihood function for MCMC for GW

# 9.10.1 Detailed Description

Header file for the Graviational Wave specific MCMC routines

#### 9.10.2 Function Documentation

# 9.10.2.1 continue\_PTMCMC\_MH\_GW()

```
void continue_PTMCMC_MH_GW (
             std::string start_checkpoint_file,
             double *** output,
             int dimension.
             int N_steps,
             int swp_freq,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             int numThreads,
             bool pool,
             bool show_prog,
             int num detectors,
             std::complex< double > ** data,
             double ** noise_psd,
             double ** frequencies,
             int * data length,
             double gps_time,
             std::string * detectors,
             int Nmod.
             int * bppe,
             std::string generation_method,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string final_checkpoint_filename )
```

Takes in an MCMC checkpoint file and continues the chain.

Obviously, the user must be sure to correctly match the dimension, number of chains, the generation\_method, the prior function, the data, psds, freqs, and the detectors (number and name), and the gps\_time to the previous run, otherwise the behavior of the sampler is undefined.

numThreads and pool do not necessarily have to be the same

## 9.10.2.2 Log\_Likelihood()

Unmarginalized log of the likelihood.

## 9.10.2.3 Log\_Likelihood\_internal()

```
double Log_Likelihood_internal (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    std::complex< double > * detector_response,
    int length,
    fftw_outline * plan )
```

Internal function for the unmarginalized log of the likelihood.

```
.5 * ((h|h)-2(D|h))
```

## 9.10.2.4 maximized\_coal\_Log\_Likelihood()

```
double maximized_coal_Log_Likelihood (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    size_t length,
    gen_params * params,
    std::string detector,
    std::string generation_method,
    fftw_outline * plan,
    double * tc,
    double * phic )
```

Function to maximize only over coalescence variables to and phic, returns the maximum values used.

## 9.10.2.5 maximized\_coal\_log\_likelihood\_IMRPhenomD() [1/3]

Function to calculate the log Likelihood as defined by -1/2 (d-h|d-h) maximized over the extrinsic parameters phic and tc.

frequency array must be uniform spacing - this shouldn't be a problem when working with real data as DFT return uniform spacing

#### **Parameters**

chirpmass in solar masses

# $\textbf{9.10.2.6} \quad \textbf{maximized\_coal\_log\_likelihood\_IMRPhenomD()} \ \, \texttt{[2/3]}$

#### **Parameters**

chirpmass in solar masses

# 9.10.2.7 maximized\_coal\_log\_likelihood\_IMRPhenomD() [3/3]

```
size_t length,
double * real_data,
double * imag_data,
double * noise,
double SNR,
double chirpmass,
double symmetric_mass_ratio,
double spin1,
double spin2,
bool NSflag,
fftw_outline * plan )
```

#### **Parameters**

chirpmass	in solar masses
-----------	-----------------

## 9.10.2.8 maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param() [1/3]

# Parameters

```
chirpmass in solar masses
```

## 9.10.2.9 maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param() [2/3]

```
double spin2,
double Luminosity_Distance,
double theta,
double phi,
double iota,
bool NSflag )
```

#### **Parameters**

```
chirpmass in solar masses
```

## 9.10.2.10 maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param() [3/3]

```
double maximized_coal_log_likelihood_IMRPhenomD_Full_Param (
             double * frequencies,
             size_t length,
             double * real_data,
             double * imag_data,
             double * noise,
             double chirpmass,
             double symmetric_mass_ratio,
             double spin1,
             double spin2,
             double Luminosity_Distance,
             double theta,
             double phi,
             double iota,
             bool NSflag,
             fftw_outline * plan )
```

## **Parameters**

```
chirpmass in solar masses
```

# 9.10.2.11 maximized\_Log\_Likelihood()

```
double maximized_Log_Likelihood (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    size_t length,
    gen_params * params,
    std::string detector,
    std::string generation_method,
    fftw_outline * plan )
```

routine to maximize over all extrinsic quantities and return the log likelihood

 $\label{local-phic} \begin{tabular}{ll} $IMRPhenomD-maximizes over DL, phic, tc, \oldsymbol{local-phic}, \oldsymbol{local-phi$ 

#### 9.10.2.12 maximized\_Log\_Likelihood\_aligned\_spin\_internal()

```
double maximized_Log_Likelihood_aligned_spin_internal (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    std::complex< double > * detector_response,
    size_t length,
    fftw_outline * plan )
```

Maximized match over coalescence variables - returns log likelihood NOT NORMALIZED for aligned spins.

Note: this function is not properly normalized for an absolute comparison. This is made for MCMC sampling, so to minimize time, constant terms like (Data|Data), which would cancel in the Metropolis-Hasting ratio, are left out for efficiency

## 9.10.2.13 maximized\_Log\_Likelihood\_unaligned\_spin\_internal()

```
double maximized_Log_Likelihood_unaligned_spin_internal (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    std::complex< double > * hplus,
    std::complex< double > * hcross,
    size_t length,
    fftw_outline * plan )
```

log likelihood function that maximizes over extrinsic parameters tc, phic, D, and phiRef, the reference frequency - for unaligned spins

Ref: arXiv 1603.02444v2

#### 9.10.2.14 MCMC\_fisher\_wrapper()

Fisher function for MCMC for GW.

Wraps the fisher calculation in src/fisher.cpp and unpacks parameters correctly for common GW analysis

Supports all the method/parameter combinations found in MCMC MH GW

## 9.10.2.15 MCMC\_likelihood\_wrapper()

log likelihood function for MCMC for GW

Wraps the above likelihood functions and unpacks parameters correctly for common GW analysis

Supports all the method/parameter combinations found in MCMC\_MH\_GW

## 9.10.2.16 PTMCMC\_method\_specific\_prep()

```
void PTMCMC_method_specific_prep (
    std::string generation_method,
    int dimension,
    double * seeding_var,
    bool local_seeding )
```

Unpacks MCMC parameters for method specific initiation.

Populates seeding vector if non supplied, populates mcmc\_Nmod, populates mcmc\_log\_beta, populates mcmc\_← intrinsic

## 9.10.2.17 PTMCMC\_MH\_dynamic\_PT\_alloc\_GW()

```
void PTMCMC_MH_dynamic_PT_alloc_GW (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             int t0,
             int nu.
             std::string chain_distribution_scheme,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             int numThreads,
             bool pool,
             bool show_prog,
             int num_detectors,
             std::complex< double > ** data,
             double ** noise_psd,
             double ** frequencies,
             int * data_length,
             double gps_time,
             std::string * detectors,
             int Nmod,
             int * bppe,
             std::string generation_method,
             std::string statistics_filename,
             std::string chain_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_filename )
```

Takes in an MCMC checkpoint file and continues the chain.

Obviously, the user must be sure to correctly match the dimension, number of chains, the generation\_method, the prior function, the data, psds, freqs, and the detectors (number and name), and the gps\_time to the previous run, otherwise the behavior of the sampler is undefined.

numThreads and pool do not necessarily have to be the same

## 9.10.2.18 PTMCMC\_MH\_GW()

```
void PTMCMC_MH_GW (
             double *** output.
             int dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             int numThreads,
             bool pool,
             bool show_prog,
             int num_detectors,
             std::complex< double > ** data,
             double ** noise_psd,
             double ** frequencies,
             int * data_length,
             double gps_time,
             std::string * detectors,
             int Nmod,
             int * bppe,
             std::string generation_method,
             std::string statistics_filename,
             std::string chain filename.
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

Wrapper for the MCMC MH function, specifically for GW analysis.

Handles the details of setting up the MCMC sampler and wraps the fisher and log likelihood to conform to the format of the sampler

*NOTE* – This sampler is NOT thread safe. There is global memory declared for each call to MCMC\_MH\_GW, so separate samplers should not be run in the same process space

Supported parameter combinations:

```
IMRPhenomD - 4 dimensions - In chirpmass, eta, chi1, chi2
```

IMRPhenomD - 7 dimensions - In D\_L, tc, phic, In chirpmass, eta, chi1, chi2

IMRPhenomD - 8 dimensions - cos inclination, RA, DEC, In D L, In chirpmass, eta, chi1, chi2

 $\frac{dCS\_IMRPhenomD\_log}{dCS\_IMRPhenomD\_log} - 8 \ dimensions - cos \ inclination, \ RA, \ DEC, \ In \ D\_L, \ In \ chirpmass, \ eta, \ chi1, \ chi2, \ In \ \ lapha^2 \ (the \ coupling \ parameter)$ 

dCS\_IMRPhenomD- 8 dimensions – cos inclination, RA, DEC, In D\_L, In chirpmass, eta, chi1, chi2,  $\alpha^2$  (the coupling parameter)

dCS\_IMRPhenomD\_root\_alpha- 8 dimensions - cos inclination, RA, DEC, In D\_L, In chirpmass, eta, chi1, chi2, \sqrt \alpha (in km) (the coupling parameter)

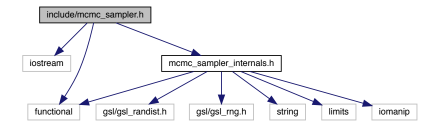
IMRPhenomPv2 - 9 dimensions - cos J\_N, In chirpmass, eta, |chi1|, |chi1|, theta\_1, theta\_2, phi\_1, phi\_2

## **Parameters**

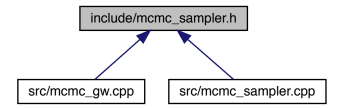
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.11 include/mcmc\_sampler.h File Reference

```
#include <iostream>
#include <functional>
#include "mcmc_sampler_internals.h"
Include dependency graph for mcmc_sampler.h:
```



This graph shows which files directly or indirectly include this file:



# **Functions**

- void mcmc\_step\_threaded (int j)
- void mcmc\_swap\_threaded (int i, int j)

void RJPTMCMC\_MH\_internal (double \*\*\*output, int \*\*\*parameter\_status, int max\_dimension, int min← \_dimension, int N\_steps, int chain\_N, double \*initial\_pos, int \*initial\_status, double \*seeding\_var, double \*chain\_temps, int swp\_freq, std::function< double(double \*, int \*, int, int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double \*, int \*, int, double \*\*, int)>fisher, std::function< void(double \*, double \*, int \*, int \*, int, int)> RJ\_proposal, int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)

Generic reversable jump sampler, where the likelihood, prior, and reversable jump proposal are parameters supplied by the user.

- void RJPTMCMC\_MH (double \*\*\*output, int \*\*\*parameter\_status, int max\_dimension, int min\_dimension, int N\_steps, int chain\_N, double \*initial\_pos, int \*initial\_status, double \*seeding\_var, double \*chain\_temps, int swp\_freq, double(\*log\_prior)(double \*param, int \*status, int max\_dimension, int chain\_id), double(\*log \_\_\_\_\_likelihood)(double \*param, int \*status, int max\_dimension, int chain\_id), void(\*fisher)(double \*param, int \*status, int max\_dimension, double \*param, int \*status, int max\_dimension, double \*status, int chain\_id), void(\*RJ\_proposal)(double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed\_status, int max\_dimension, int chain\_id), int num \_\_\_\_\_ Threads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)
- void PTMCMC\_MH\_dynamic\_PT\_alloc\_internal (double \*\*\*output, int dimension, int N\_steps, int chain 
   \_N, int max\_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, int t0, int nu, std::string chain\_distribution\_scheme, std::function< double(double \*, int \*, int, int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double \*, int \*, int, double \*\*, int)>fisher, int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)

Dyanmically tunes an MCMC for optimal spacing. step width, and chain number.

- void PTMCMC\_MH\_dynamic\_PT\_alloc (double \*\*\*output, int dimension, int N\_steps, int chain\_N, int max
   \_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp
   \_freq, int t0, int nu, std::string chain\_distribution\_scheme, double(\*log\_prior)(double \*param, int dimension, int chain\_id), double(\*log\_likelihood)(double \*param, int dimension, int chain\_id), void(\*fisher)(double \*param, int dimension, double \*\*fisher, int chain\_id), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)
- void PTMCMC\_MH\_dynamic\_PT\_alloc (double \*\*\*output, int dimension, int N\_steps, int chain\_N, int max
   \_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_
   freq, int t0, int nu, std::string chain\_distribution\_scheme, double(\*log\_prior)(double \*param, int dimension), double(\*log\_likelihood)(double \*param, int dimension), void(\*fisher)(double \*param, int dimension, double \*stisher), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_
   filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)
- void continue\_PTMCMC\_MH (std::string start\_checkpoint\_file, double \*\*\*output, int N\_steps, int swp\_← freq, double(\*log\_prior)(double \*param, int dimension, int chain\_id), double(\*log\_likelihood)(double \*param, int dimension, int chain\_id), void(\*fisher)(double \*param, int dimension, double \*\*fisher, int chain\_id), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string end\_checkpoint\_file)
- void continue\_PTMCMC\_MH (std::string start\_checkpoint\_file, double \*\*\*output, int N\_steps, int swp\_
   freq, double(\*log\_prior)(double \*param, int dimension), double(\*log\_likelihood)(double \*param, int dimension), void(\*fisher)(double \*param, int dimension, double \*\*fisher), int numThreads, bool pool, bool show
   \_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string end\_checkpoint\_file)
- void PTMCMC\_MH\_loop (sampler \*sampler)

Internal function that runs the actual loop for the sampler.

- void PTMCMC\_MH\_step\_incremental (sampler \*sampler, int increment)
  - Internal function that runs the actual loop for the sampler increment version.
- void PTMCMC\_MH (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension), double(\*log\_likelihood)(double \*param, int dimension), void(\*fisher)(double \*param, int dimension, double \*\*fisher), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_
  filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_filename)
- void PTMCMC\_MH (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension, int

chain\_id), double(\*log\_likelihood)(double \*param, int dimension, int chain\_id), void(\*fisher)(double \*param, int dimension, double \*\*fisher, int chain\_id), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_
filename, std::string checkpoint\_filename)

void PTMCMC\_MH\_internal (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_←
pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, std::function< double(double \*, int \*, int,
int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double
\*, int \*, int, double \*\*, int)>fisher, int numThreads, bool pool, bool show\_prog, std::string statistics\_←
filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std
::string checkpoint\_filename)

Generic sampler, where the likelihood, prior are parameters supplied by the user.

• void continue\_PTMCMC\_MH\_internal (std::string start\_checkpoint\_file, double \*\*\*output, int N\_steps, int swp\_freq, std::function< double(double \*, int \*, int, int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double \*, int \*, int, double \*\*, int)>fisher, int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_cilename, std::string likelihood\_log\_filename, std::string end\_checkpoint\_file)

Routine to take a checkpoint file and begin a new chain at said checkpoint.

# 9.11.1 Detailed Description

Header file for mcmc sampler

#### 9.11.2 Function Documentation

```
9.11.2.1 continue_PTMCMC_MH() [1/2]
```

```
void continue PTMCMC MH (
             std::string start_checkpoint_file,
             double *** output,
             int N_steps,
             int swp freq,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             double(*)(double *param, int dimension, int chain_id) log_likelihood,
             void(*)(double *param, int dimension, double **fisher, int chain_id) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string end_checkpoint_file )
```

	start_checkpoint_file	File for starting checkpoint
out	output	output array, dimensions: output[chain_N][N_steps][dimension]
	N_steps	Number of new steps to take
	swp_freq	frequency of swap attempts between temperatures
	log_prior	Funcion pointer for the log_prior

## **Parameters**

log_likelihood	Function pointer for the log_likelihood
fisher	Function pointer for the fisher - if NULL, fisher steps are not used
numThreads	Number of threads to use
pool	Boolean for whether to use deterministic'' vsstochastic" sampling
show_prog	Boolean for whether to show progress or not (turn off for cluster runs
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
end_checkpoint_file	Filename to output data for checkpoint at the end of the continued run, if empty string, not saved

# **9.11.2.2** continue\_PTMCMC\_MH() [2/2]

```
void continue_PTMCMC_MH (
           std::string start_checkpoint_file,
            double *** output,
            int N_steps,
             int swp_freq,
             double(*)(double *param, int dimension) log_prior,
             double(*)(double *param, int dimension) log_likelihood,
             void(*)(double *param, int dimension, double **fisher) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string end_checkpoint_file )
```

	start_checkpoint_file	File for starting checkpoint
out	output	output array, dimensions: output[chain_N][N_steps][dimension]
	N_steps	Number of new steps to take
	swp_freq	frequency of swap attempts between temperatures
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use
	pool	Boolean for whether to use deterministic'' vsstochastic" sampling
	show_prog	Boolean for whether to show progress or not (turn off for cluster runs
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output

## **Parameters**

	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	end_checkpoint_file	Filename to output data for checkpoint at the end of the continued run, if empty string, not saved

# 9.11.2.3 continue\_PTMCMC\_MH\_internal()

```
void continue_PTMCMC_MH_internal (
            std::string start_checkpoint_file,
             double *** output,
             int N_steps,
             int swp_freq,
             std::function< double(double *, int *, int, int) > log_prior,
             std::function< double(double *, int *, int, int)> log_likelihood,
             std::function< void(double *, int *, int, double **, int)> fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string end_checkpoint_file )
```

Routine to take a checkpoint file and begin a new chain at said checkpoint.

See MCMC\_MH\_internal for more details of parameters (pretty much all the same)

	start_checkpoint_file	File for starting checkpoint
out	output	output array, dimensions: output[chain_N][N_steps][dimension]
	N_steps	Number of new steps to take
	swp_freq	frequency of swap attempts between temperatures
	log_prior	std::function for the log_prior function – takes double *position, int dimension, int chain_id
	log_likelihood	std::function for the log_likelihood function – takes double *position, int dimension, int chain_id
	fisher	std::function for the fisher function – takes double *position, int dimension, double **output_fisher, int chain_id
	numThreads	Number of threads to use
	pool	Boolean for whether to use deterministic'' vsstochastic sampling
	show_prog	Boolean for whether to show progress or not (turn off for cluster runs
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty
		string to skip  Generated by Doxyge
	end_checkpoint_file	Filename to output data for checkpoint at the end of the continued run, if empty string, not saved

## **9.11.2.4 PTMCMC\_MH()** [1/2]

```
void PTMCMC_MH (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             double(*)(double *param, int dimension) log_prior,
             double(*)(double *param, int dimension) log_likelihood,
             void(*)(double *param, int dimension, double **fisher) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_filename )
```

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have >2 threads)
	show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	checkpoint_filename	Filename to output data for checkpoint, if empty string, not saved

# **9.11.2.5 PTMCMC\_MH()** [2/2]

```
void PTMCMC_MH (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             double(*)(double *param, int dimension, int chain_id) log_likelihood,
             void(*)(double *param, int dimension, double **fisher, int chain_id) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_filename )
```

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have >2 threads)
	show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	checkpoint_filename	Filename to output data for checkpoint, if empty string, not saved

## 9.11.2.6 PTMCMC\_MH\_dynamic\_PT\_alloc() [1/2]

```
void PTMCMC_MH_dynamic_PT_alloc (
            double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             int t0,
             int nu,
             std::string chain_distribution_scheme,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             double(*)(double *param, int dimension, int chain_id) log_likelihood,
             void(*)(double *param, int dimension, double **fisher, int chain_id) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

out	output	Output chains, shape is double[max chain N,
		N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain AFTER chain allocation
	chain_N	Maximum number of chains to use
	max_chain_N_thermo_ensemble	Maximum number of chains to use in the thermodynamic ensemble (may use less)
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Final chain temperatures used – should be shape double[chain_N]
	swp_freq	the frequency with which chains are swapped
	t0	Time constant of the decay of the chain dynamics ( $\sim$ 1000)
	nu	Initial amplitude of the dynamics (~100)
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have $>$ 2 threads)
	show_prog	boolean whether to print out progress (for example, should be se to `'false" if submitting to a cluster)
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output

## **Parameters**

likelihood_log_file	Ame Filename to write the log_likelihood and log_prior at each step –
	use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

## 9.11.2.7 PTMCMC\_MH\_dynamic\_PT\_alloc() [2/2]

```
void PTMCMC_MH_dynamic_PT_alloc (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             int t0,
             int nu,
             std::string chain_distribution_scheme,
             double(*)(double *param, int dimension) log_prior,
             double(*)(double *param, int dimension) log_likelihood,
             void(*)(double *param, int dimension, double **fisher) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

out	output	Output chains, shape is double[max_chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain AFTER chain allocation
	chain_N	Maximum number of chains to use
	max_chain_N_thermo_ensemble	Maximum number of chains to use in the thermodynamic ensemble (may use less)
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Final chain temperatures used – should be shape double[chain_N]
	swp_freq	the frequency with which chains are swapped
	tO	Time constant of the decay of the chain dynamics (∼1000)
	nu	Initial amplitude of the dynamics (~100)
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood

#### **Parameters**

fisher	Function pointer for the fisher - if NULL, fisher steps are not used
numThreads	Number of threads to use (=1 is single threaded)
pool	boolean to use stochastic chain swapping (MUST have $>$ 2 threads)
show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.11.2.8 PTMCMC\_MH\_dynamic\_PT\_alloc\_internal()

```
void PTMCMC_MH_dynamic_PT_alloc_internal (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             int t0,
             int nu,
             std::string chain_distribution_scheme,
             std::function< double(double *, int *, int, int) > log_prior,
             std::function< double(double *, int *, int, int) > log_likelihood,
             std::function< void(double *, int *, int, double **, int)> fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

Dyanmically tunes an MCMC for optimal spacing. step width, and chain number.

NOTE: nu, and t0 parameters determine the dynamics, so these are important quantities. nu is related to how many swap attempts it takes to substantially change the temperature ladder, why t0 determines the length of the total dyanimcally period. Moderate initial choices would be 10 and 1000, respectively.

Based on arXiv:1501.05823v3

Currently, Chain number is fixed

max\_chain\_N\_thermo\_ensemble sets the maximium number of chains to use to in successively hotter chains to cover the likelihood surface while targeting an optimal swap acceptance target\_swp\_acc.

max\_chain\_N determines the total number of chains to run once thermodynamic equilibrium has been reached. This results in chains being added after the initial PT dynamics have finished according to chain\_distribution\_\circ} scheme.

If no preference, set  $max\_chain\_N\_thermo\_ensemble = max\_chain\_N = numThreads = (number of cores (number of threads if hyperthreaded))— this will most likely be the most optimal configuration. If the number of cores on the system is low, you may want to use <math>n*numThreads$  for some integer n instead, depending on the system.

chain\_distribution\_scheme:

"cold": All chains are added at T=1 (untempered)

"refine": Chains are added between the optimal temps geometrically – this may be a good option as it will be a good approximation of the ideal distribution of chains, while keeping the initial dynamical time low

"double": Chains are added in order of rising temperature that mimic the distribution achieved by the earier PT dynamics

"half\_ensemble": For every cold chain added, half of the ensemble is added again. Effectively, two cold chains for every ensemble

out	output	Output chains, shape is double[max_chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain AFTER chain allocation
	chain_N	Maximum number of chains to use
	max_chain_N_thermo_ensemble	Maximum number of chains to use in the thermodynamic ensemble (may use less)
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
out	chain_temps	Final chain temperatures used – should be shape double[chain_N]
	swp_freq	the frequency with which chains are swapped
	tO	Time constant of the decay of the chain dynamics (~1000)
	nu	Initial amplitude of the dynamics (~100)
	log_prior	std::function for the log_prior function – takes double *position, int dimension, int chain_id
	log_likelihood	std::function for the log_likelihood function – takes double *position, int dimension, int chain_id
	fisher	std::function for the fisher function – takes double *position, int dimension, double **output_fisher, int chain_id
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have $>$ 2 threads)
	show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

#### 9.11.2.9 PTMCMC\_MH\_internal()

```
void PTMCMC_MH_internal (
             double *** output,
             int dimension.
             int N_steps,
             int chain N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             std::function< double(double *, int *, int, int) > log_prior,
             std::function< double(double *, int *, int, int) > log_likelihood,
             std::function< void(double *, int *, int, double **, int) > fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto corr filename.
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

Generic sampler, where the likelihood, prior are parameters supplied by the user.

Base of the sampler, generic, with user supplied quantities for most of the samplers properties

Uses the Metropolis-Hastings method, with the option for Fisher/MALA steps if the Fisher routine is supplied.

3 modes to use -

single threaded (numThreads = 1) runs single threaded

multi-threaded `'deterministic'' (numThreads>1; pool = false) progresses each chain in parallel for swp\_freq steps, then waits for all threads to complete before swapping temperatures in sequenctial order (j, j+1) then (j+1, j+2) etc (sequenctially)

multi-threaded `'stochastic'' (numThreads>2; pool = true) progresses each chain in parallel by queueing each temperature and evaluating them in the order they were submitted. Once finished, the threads are queued to swap, where they swapped in the order they are submitted. This means the chains are swapped randomly, and the chains do NOT finish at the same time. The sampler runs until the the 0th chain reaches the step number

Note on limits: In the prior function, if a set of parameters should be disallowed, return -std::numeric\_ $\leftarrow$  limits<double>::infinity() - (this is in the limits> file in std)

Format for the auto\_corr file (compatable with csv, dat, txt extensions): each row is a dimension of the cold chain, with the first row being the lengths used for the auto-corr calculation:

```
lengths: length1, length2...
```

dim1: length1, length2...

Format for the chain file (compatable with csv, dat, txt extensions): each row is a step, each column a dimension:

Step1: dim1, dim2, ...

Step2: dim1 , dim2 , ...

Statistics\_filename : should be txt extension

checkpoint\_file: This file saves the final position of all the chains, as well as other metadata, and can be loaded by the function <FUNCTION> to continue the chain from the point it left off. Not meant to be read by humans, the data order is custom to this software library. An empty string ("") means no checkpoint will be saved. For developers, the contents are:

dimension, # of chains

temps of chains

Stepping widths of all chains

Final position of all chains

## **Parameters**

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	std::function for the log_prior function – takes double *position, int dimension, int chain_id
	log_likelihood	std::function for the log_likelihood function – takes double *position, int dimension, int chain_id
	fisher	std::function for the fisher function – takes double *position, int dimension, double **output_fisher, int chain_id
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have >2 threads)
	show_prog	boolean whether to print out progress (for example, should be set to `false' if submitting to a cluster)
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.11.2.10 PTMCMC\_MH\_loop()

Internal function that runs the actual loop for the sampler.

#### 9.11.2.11 PTMCMC\_MH\_step\_incremental()

Internal function that runs the actual loop for the sampler – increment version.

The regular loop function runs for the entire range, this increment version will only step `increment' steps – asynchronous: steps are measured by the cold chains NEEDS TO CHANGE

## 9.11.2.12 RJPTMCMC\_MH()

```
void RJPTMCMC_MH (
            double *** output,
             int *** parameter_status,
             int max dimension,
             int min_dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             int * initial_status,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             double(*)(double *param, int *status, int max_dimension, int chain_id) log_prior,
             double(*)(double *param, int *status, int max_dimension, int chain_id) <math>log\_{\leftarrow}
likelihood,
             void(*)(double *param, int *status, int max_dimension, double **fisher, int chain↔
_id) fisher,
             void(*)(double *current_param, double *proposed_param, int *current_status, int
*proposed_status, int max_dimension, int chain_id) RJ_proposal,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
out	parameter_status	Parameter status for each step corresponding to the output chains, shape is double[chain_N, N_steps,dimension]
	max_dimension	maximum dimension of the parameter space being explored – only consideration is memory, as memory scales with dimension. Keep this reasonable, unless memory is REALLY not an issue
	min_dimension	minimum dimension of the parameter space being explored >=1
	N_steps	Number of total steps to be taken, per chain

#### **Parameters**

chain_N	Number of chains
initial_pos	Initial position in parameter space - shape double[dimension]
initial_status	Initial status of the parameters in the initial position in parameter space - shape int[max_dim]
seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[max_dimension] – initial seeding of zero corresponds to the dimension turned off initially
chain_temps	Double array of temperatures for the chains
swp_freq	the frequency with which chains are swapped
RJ_proposal	std::function for the log_likelihood function – takes double *position, int *param_status,int dimension, int chain_id
numThreads	Number of threads to use (=1 is single threaded)
pool	boolean to use stochastic chain swapping (MUST have >2 threads)
show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.11.2.13 RJPTMCMC\_MH\_internal()

```
void RJPTMCMC_MH_internal (
            double *** output,
             int *** parameter_status,
             int max_dimension,
             int min_dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             int * initial_status,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             std::function< double(double *, int *, int, int) > log_prior,
             std::function< double(double *, int *, int, int)> log_likelihood,
             std::function< void(double *, int *, int, double **, int)> fisher,
             std::function< void(double *, double *, int *, int *, int, int) > RJ_proposal,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

Generic reversable jump sampler, where the likelihood, prior, and reversable jump proposal are parameters supplied by the user.

Note: Using a min\_dimension tells the sampler that there is a base model'', and that the dimensions from min\_dim to max\_dim are small" corrections to that model. This helps inform some of the proposal algorithms and speeds up computation. If using discrete models with no overlap of variables (ie model A or model B), set min\_dim to 0. Even if reusing certain parameters, if the extra dimensions don't describe 'small' deviations, it's probably best to set min dim to 0.

Currently, no dynamic PT option, as it would be too many free parameters for the sampler to converge to a reasonable temperature distribution in a reasonable amount of time. Best use case, use the PTMCMC\_MH\_dyanmic\_PT for the `'base'' dimension space, and use that temperature ladder.

Base of the sampler, generic, with user supplied quantities for most of the samplers properties

Uses the Metropolis-Hastings method, with the option for Fisher/MALA steps if the Fisher routine is supplied.

3 modes to use -

single threaded (numThreads = 1) runs single threaded

multi-threaded `'deterministic'' (numThreads>1; pool = false) progresses each chain in parallel for swp\_freq steps, then waits for all threads to complete before swapping temperatures in sequenctial order (j, j+1) then (j+1, j+2) etc (sequenctially)

multi-threaded `'stochastic' (numThreads>2; pool = true) progresses each chain in parallel by queueing each temperature and evaluating them in the order they were submitted. Once finished, the threads are queued to swap, where they swapped in the order they are submitted. This means the chains are swapped randomly, and the chains do NOT finish at the same time. The sampler runs until the the 0th chain reaches the step number

Note on limits: In the prior function, if a set of parameters should be disallowed, return -std::numeric\_\(-\circ\) limits<double>::infinity() - (this is in the limits> file in std)

The parameter array uses the dimensions [0,min\_dim] always, and [min\_dim, max\_dim] in RJPTMCMC fashion

Format for the auto\_corr file (compatable with csv, dat, txt extensions): each row is a dimension of the cold chain, with the first row being the lengths used for the auto-corr calculation:

lengths: length1, length2 ...

dim1: length1, length2...

Format for the chain file (compatable with csv, dat, txt extensions): each row is a step, each column a dimension:

Step1: dim1, dim2, ..., max\_dim, param\_status1, param\_status2, ...

Step2: dim1 , dim2 , ..., max\_dim, param\_status1, param\_status2, ...

Statistics filename: should be txt extension

checkpoint\_file: This file saves the final position of all the chains, as well as other metadata, and can be loaded by the function <FUNCTION> to continue the chain from the point it left off. Not meant to be read by humans, the data order is custom to this software library. An empty string ("") means no checkpoint will be saved. For developers, the contents are:

dimension, # of chains

temps of chains

Stepping widths of all chains

Final position of all chains

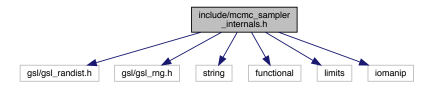
### **Parameters**

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
out	parameter_status	Parameter status for each step corresponding to the output chains, shape is double[chain_N, N_steps,dimension]
	max_dimension	maximum dimension of the parameter space being explored – only consideration is memory, as memory scales with dimension. Keep this reasonable, unless memory is REALLY not an issue
	min_dimension	minimum dimension of the parameter space being explored >=1
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	initial_status	Initial status of the parameters in the initial position in parameter space - shape int[max_dim]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[max_dimension] – initial seeding of zero corresponds to the dimension turned off initially
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	std::function for the log_prior function – takes double *position, int *param_status, int dimension, int chain_id
	log_likelihood	std::function for the log_likelihood function – takes double *position, int *param_status,int dimension, int chain_id
	fisher	std::function for the fisher function – takes double *position, int *param_status,int dimension, double **output_fisher, int chain_id
	RJ_proposal	std::function for the log_likelihood function – takes double *position, int *param_status,int dimension, int chain_id
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have >2 threads)
	show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

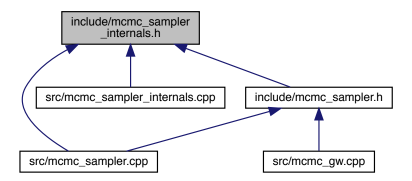
# 9.12 include/mcmc\_sampler\_internals.h File Reference

```
#include <gsl/gsl_randist.h>
#include <gsl/gsl_rng.h>
#include <string>
#include <functional>
#include <limits>
#include <iomanip>
```

Include dependency graph for mcmc\_sampler\_internals.h:



This graph shows which files directly or indirectly include this file:



## Classes

· class sampler

### **Functions**

• int mcmc\_step (sampler \*sampler, double \*current\_param, double \*next\_param, int \*current\_status, int \*next\_status, int chain\_number)

interface function between the sampler and the internal step functions

void gaussian\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_

 status, int \*proposed\_status, int chain\_id)

Straight gaussian step.

• void fisher\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed\_status, int chain\_index)

Fisher informed gaussian step.

- void update\_fisher (sampler \*sampler, double \*current\_param, int \*param\_status, int chain\_index)
- void mmala\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed\_status)

MMALA informed step - Currently not supported.

void diff\_ev\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed\_status, int chain\_id)

differential evolution informed step

- void RJ\_smooth\_history (sampler \*sampler, int \*current\_param\_status, int base\_history\_id, double \*eff\_←
  history\_coord, int \*eff\_history\_status, int chain\_id)
- void RJ\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed status, int chain number)

RJ-proposal step for trans-dimensional MCMCs.

void chain\_swap (sampler \*sampler, double \*\*\*output, int \*\*\*param\_status, int step\_num, int \*swp\_

 accepted, int \*swp rejected)

subroutine to perform chain comparison for parallel tempering

int single\_chain\_swap (sampler \*sampler, double \*chain1, double \*chain2, int \*chain1\_status, int \*chain1\_status, int \*chain1\_status, int T1 index, int T2 index)

subroutine to actually swap two chains

void assign probabilities (sampler \*sampler, int chain index)

update and initiate probabilities for each variety of step

Copies contents of one chain to another.

• bool check sampler status (sampler \*samplerptr)

Checks the status of a sampler for the stochastic sampling.

void update\_step\_widths (sampler \*samplerptr, int chain\_id)

Updates the step widths, shooting for 20% acceptance ratios for each type of step.

- void allocate sampler mem (sampler \*sampler)
- void deallocate\_sampler\_mem (sampler \*sampler)
- void update history (sampler \*sampler, double \*new params, int \*new param status, int chain index)
- void write\_stat\_file (sampler \*sampler, std::string filename)
- void write\_checkpoint\_file (sampler \*sampler, std::string filename)

Routine that writes metadata and final positions of a sampler to a checkpoint file.

void load\_checkpoint\_file (std::string check\_file, sampler \*sampler)

load checkpoint file into sampler struct

• void load temps checkpoint file (std::string check file, double \*temps, int chain N)

load temperatures from checkpoint file

- void assign\_ct\_p (sampler \*sampler, int step, int chain\_index)
- void assign\_ct\_m (sampler \*sampler, int step, int chain\_index)
- void assign\_initial\_pos (sampler \*samplerptr, double \*initial\_pos, int \*initial\_status, double \*seeding\_var)
- double PT dynamical timescale (int t0, int nu, int t)

Timescale of the PT dynamics.

• void update\_temperatures (sampler \*samplerptr, int t0, int nu, int t)

updates the temperatures for a sampler such that all acceptance rates are equal

• void initiate\_full\_sampler (sampler \*sampler\_new, sampler \*sampler\_old, int chain\_N\_thermo\_ensemble, int chain\_N, std::string chain\_allocation\_scheme)

For the dynamic PT sampler, this is the function that starts the full sampler with the max number of chains.

### **Variables**

• const double **limit** inf = -std::numeric limits<double>::infinity()

### 9.12.1 Detailed Description

Internal functions of the generic MCMC sampler (nothing specific to GW)

### 9.12.2 Function Documentation

### 9.12.2.1 assign\_probabilities()

update and initiate probabilities for each variety of step

Type 0: Gaussian step

Type 1: Differential Evolution step

Type 2: MMALA step (currently not supported)

Type 3: Fisher step

### 9.12.2.2 chain\_swap()

subroutine to perform chain comparison for parallel tempering

The total output file is passed, and the chains are swapped sequentially

This is the routine for `'Deterministic" sampling (parallel or sequential, but not pooled)

### **Parameters**

sampler	sampler struct
output	output vector containing chains
param_status	Parameter status
step_num	current step number

### 9.12.2.3 check\_sampler\_status()

```
bool check_sampler_status (
          sampler * samplerptr )
```

Checks the status of a sampler for the stochastic sampling.

Just loops through the ref\_chain\_status variables

### 9.12.2.4 diff\_ev\_step()

differential evolution informed step

Differential evolution uses the past history of the chain to inform the proposed step:

Take the difference of two random, accepted previous steps and step along that with some step size, determined by a gaussian

### **Parameters**

	sampler	Sampler struct
	current_param	current position in parameter space
out	proposed_param	Proposed position in parameter space

### 9.12.2.5 fisher\_step()

Fisher informed gaussian step.

#### **Parameters**

	sampler	Sampler struct
	current_param	current position in parameter space
out	proposed_param	Proposed position in parameter space

### 9.12.2.6 gaussian\_step()

```
int * current_status,
int * proposed_status,
int chain_id )
```

Straight gaussian step.

### **Parameters**

	sampler	Sampler struct
	current_param	current position in parameter space
out	proposed_param	Proposed position in parameter space

### 9.12.2.7 initiate\_full\_sampler()

```
void initiate_full_sampler (
          sampler * sampler_new,
          sampler * sampler_old,
          int chain_N_thermo_ensemble,
          int chain_N,
          std::string chain_allocation_scheme )
```

For the dynamic PT sampler, this is the function that starts the full sampler with the max number of chains.

The output file will be reused, but the positions are set back to zero (copying the current position to position zero)

Assumes the output, chain\_temps have been allocated in memory for the final number of chains chain\_N and steps N steps

Allocates memory for the new sampler sampler\_new -> it's the user's responsibility to deallocate with deallocate \( \cdot \) \_sampler\_mem

#### **Parameters**

sampler_old	Dynamic sampler
chain_N_thermo_ensemble	Number of chains used in the thermodynamic ensemble
chain_N	Number of chains to use in the static sampler
chain_allocation_scheme	Scheme to use to allocate any remaining chains

### 9.12.2.8 load\_checkpoint\_file()

load checkpoint file into sampler struct

NOTE - allocate\_sampler called in function - MUST deallocate manually

NOTE - sampler->chain\_temps allocated internally - MUST free manually

### 9.12.2.9 load\_temps\_checkpoint\_file()

```
void load_temps_checkpoint_file (
    std::string check_file,
    double * temps,
    int chain_N )
```

load temperatures from checkpoint file

Assumed the temps array is already allocated in memory for the correct number of chains

Just a utility routine to read temperatures from checkpoint file

It would be easy to read in the chain number and allocate memory in the function, but I prefer to leave allocation/deallocation up to the client

### 9.12.2.10 mmala\_step()

MMALA informed step - Currently not supported.

### **Parameters**

		sampler	Sampler struct
		current_param	current position in parameter space
out <i>proposed_param</i>		proposed_param	Proposed position in parameter space

### 9.12.2.11 PT\_dynamical\_timescale()

Timescale of the PT dynamics.

kappa in the the language of arXiv:1501.05823v3

### **Parameters**

tO	Timescale of the dyanmics	
nu	Initial amplitude (number of steps to base dynamics on)	
t	current time	

### 9.12.2.12 RJ\_smooth\_history()

### **Parameters**

	sampler	Current sampler
	current_param_status	Current parameters to match
	base_history_id	Original history element
out	eff_history_coord	Modified history coord
out	eff_history_status	Modified History status
	chain_id	Chain ID of the current chain

### 9.12.2.13 RJ\_step()

```
void RJ_step (
          sampler * sampler,
          double * current_param,
          double * proposed_param,
          int * current_status,
          int * proposed_status,
          int chain_number )
```

RJ-proposal step for trans-dimensional MCMCs.

This extra step may seem unnecessary, I'm just adding it in in case the extra flexibility is useful in the future for preprocessing of the chain before sending it to the user's RJ\_proposal

### **Parameters**

	sampler	sampler
	current_param	current coordinates in parameter space
out	proposed_param	Proposed coordinates in parameter space
	current_status	Current status of parameters
out	proposed_status	Proposed status of parameters
	chain_number	chain mumber

### 9.12.2.14 single\_chain\_swap()

subroutine to actually swap two chains

This is the more general subroutine, which just swaps the two chains passed to the function

#### **Parameters**

sampler	sampler structure	
chain1	parameter position of chain that could be changed	
chain2	chain that is not swapped, but provides parameters to be swapped by the other chain	
chain1_status	Parameter status array for chain1	
chain2_status Parameter status array for chain2		
T1_index number of chain swappe in chain_temps		
T2_index	number of chain swapper in chain_temps	

### 9.12.2.15 transfer\_chain()

Copies contents of one chain to another.

Transfers id\_source in samplerptr\_source to id\_dest samplerptr\_dest

NOTE: This copies the VALUE, not the reference. This could be expensive, so use with caution

id\_dest is ERASED

samplerptr\_dest and samplerptr\_source MUST have the same dimension, the same sampling details (like having or not having a fisher) etc

samplerptr\_dest must be previously allocated properly

As output is the largest transfer by far, the transfer\_output flag can be used to allow the user to handle that manually.

### 9.12.2.16 update\_temperatures()

updates the temperatures for a sampler such that all acceptance rates are equal

Follows the algorithm outlined in arXiv:1501.05823v3

Fixed temperatures for the first and last chain

used in MCMC\_MH\_dynamic\_PT\_alloc\_internal

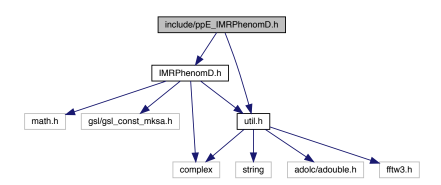
For defined results, this should be used while the sampler is using non-pooling methods

### 9.12.2.17 write\_checkpoint\_file()

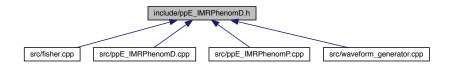
Routine that writes metadata and final positions of a sampler to a checkpoint file.

## 9.13 include/ppE\_IMRPhenomD.h File Reference

```
#include "IMRPhenomD.h"
#include "util.h"
Include dependency graph for ppE_IMRPhenomD.h:
```



This graph shows which files directly or indirectly include this file:

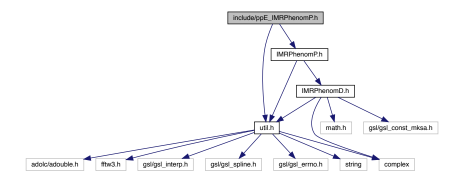


### **Classes**

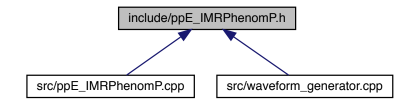
- class ppE\_IMRPhenomD\_Inspiral< T >
- class ppE\_IMRPhenomD\_IMR< T >
- class dCS\_IMRPhenomD\_log< T >
- class dCS\_IMRPhenomD< T >
- class EdGB IMRPhenomD log< T >
- class EdGB\_IMRPhenomD< T >

## 9.14 include/ppE\_IMRPhenomP.h File Reference

```
#include "util.h"
#include "IMRPhenomP.h"
Include dependency graph for ppE_IMRPhenomP.h:
```



This graph shows which files directly or indirectly include this file:

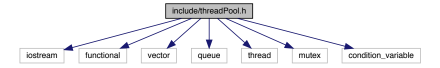


## Classes

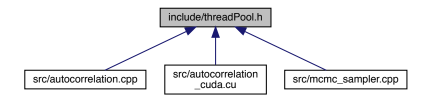
- class ppE\_IMRPhenomPv2\_Inspiral< T >
- class ppE\_IMRPhenomPv2\_IMR< T >

### 9.15 include/threadPool.h File Reference

```
#include <iostream>
#include <functional>
#include <vector>
#include <queue>
#include <thread>
#include <mutex>
#include <condition_variable>
Include dependency graph for threadPool.h:
```



This graph shows which files directly or indirectly include this file:



## **Classes**

- class default\_comp< jobtype >
  - Default comparator for priority\_queue in threadPool no comparison.
- class threadPool< jobtype, comparator >

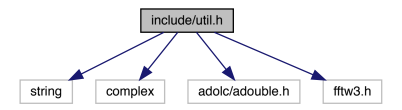
Class for creating a pool of threads to asynchronously distribute work.

### 9.15.1 Detailed Description

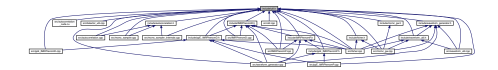
Header file (declarations and definitions because of template functions) for the implementation of a generic thread pool

### 9.16 include/util.h File Reference

```
#include <string>
#include <complex>
#include "adolc/adouble.h"
#include <fftw3.h>
#include <gsl/gsl_interp.h>
#include <gsl/gsl_spline.h>
#include <gsl/gsl_errno.h>
Include dependency graph for util.h:
```



This graph shows which files directly or indirectly include this file:



### Classes

- · struct fftw outline
- struct  $sph_harm < T >$
- struct gen\_params
- struct useful powers

To speed up calculations within the for loops, we pre-calculate reoccuring powers of M\*F and Pi, since the pow() function is prohibatively slow.

• struct source\_parameters< T >

## **Macros**

- #define PBSTR "|||||||||||||
- #define PBWIDTH 60

#### **Functions**

void initiate\_LumD\_Z\_interp (gsl\_interp\_accel \*\*Z\_DL\_accel\_ptr, gsl\_spline \*\*Z\_DL\_spline\_ptr)

Function that uses the GSL libraries to interpolate pre-calculated Z-D L data.

void free\_LumD\_Z\_interp (gsl\_interp\_accel \*\*Z\_DL\_accel\_ptr, gsl\_spline \*\*Z\_DL\_spline\_ptr)

Frees the allocated interpolation function.

- adouble Z\_from\_DL\_interp (adouble DL, gsl\_interp\_accel \*Z\_DL\_accel\_ptr, gsl\_spline \*Z\_DL\_spline\_ptr)
- double Z from\_DL\_interp (double DL, gsl\_interp\_accel \*Z\_DL\_accel\_ptr, gsl\_spline \*Z\_DL\_spline\_ptr)
- double Z from DL (double DL, std::string cosmology)

Calculates the redshift given the luminosity distance.

double DL\_from\_Z (double Z, std::string cosmology)

Calculates the luminosity distance given the redshift.

• double cosmology\_interpolation\_function (double x, double \*coeffs, int interp\_degree)

Custom interpolation function used in the cosmology calculations.

double cosmology\_lookup (std::string cosmology)

Helper function for mapping cosmology name to an internal index.

adouble Z from DL (adouble DL, std::string cosmology)

Calculates the redshift given the luminosity distance adouble version for ADOL-C implementation.

adouble DL\_from\_Z (adouble Z, std::string cosmology)

Calculates the luminosity distance given the redshift adouble version for ADOL-C implementation.

adouble cosmology interpolation function (adouble x, double \*coeffs, int interp degree)

Custom interpolation function used in the cosmology calculations adouble version for ADOL-C.

void printProgress (double percentage)

routine to print the progress of a process to the terminal as a progress bar

void allocate FFTW mem forward (fftw outline \*plan, int length)

Allocate memory for FFTW3 methods used in a lot of inner products input is a locally defined structure that houses all the pertinent data.

• void allocate\_FFTW\_mem\_reverse (fftw\_outline \*plan, int length)

Allocate memory for FFTW3 methods used in a lot of inner products –INVERSE input is a locally defined structure that houses all the pertinent data.

void deallocate FFTW mem (fftw outline \*plan)

deallocates the memory used for FFTW routines

double \*\* allocate\_2D\_array (int dim1, int dim2)

Utility to malloc 2D array.

- int \*\* allocate\_2D\_array\_int (int dim1, int dim2)
- void deallocate\_2D\_array (double \*\*array, int dim1, int dim2)

Utility to free malloc'd 2D array.

- void deallocate\_2D\_array (int \*\*array, int dim1, int dim2)
- double \*\*\* allocate 3D array (int dim1, int dim2, int dim3)

Utility to malloc 3D array.

- int \*\*\* allocate\_3D\_array\_int (int dim1, int dim2, int dim3)
- void deallocate\_3D\_array (double \*\*\*array, int dim1, int dim2, int dim3)

Utility to free malloc'd 2D array.

• void deallocate\_3D\_array (int \*\*\*array, int dim1, int dim2, int dim3)

Utility to free malloc'd 2D array.

void read file (std::string filename, double \*\*output, int rows, int cols)

Utility to read in data.

void read\_file (std::string filename, double \*output)

Utility to read in data (single dimension vector)

• void read\_LOSC\_data\_file (std::string filename, double \*output, double \*data\_start\_time, double \*duration, double \*fs)

Read data file from LIGO Open Science Center.

• void read\_LOSC\_PSD\_file (std::string filename, double \*\*output, int rows, int cols)

Read PSD file from LIGO Open Science Center.

• void allocate\_LOSC\_data (std::string \*data\_files, std::string psd\_file, int num\_detectors, int psd\_length, int data\_file\_length, double trigger\_time, std::complex< double > \*\*data, double \*\*psds, double \*\*freqs)

Prepare data for MCMC directly from LIGO Open Science Center.

- void free\_LOSC\_data (std::complex < double > \*\*data, double \*\*psds, double \*\*freqs, int num\_detectors, int length)
- void tukey\_window (double \*window, int length, double alpha)

Tukey window function for FFTs.

void write file (std::string filename, double \*\*input, int rows, int cols)

Utility to write 2D array to file.

void write file (std::string filename, double \*input, int length)

Utility to write 1D array to file.

double calculate\_eta (double mass1, double mass2)

Calculates the symmetric mass ration from the two component masses.

- adouble calculate eta (adouble mass1, adouble mass2)
- double calculate chirpmass (double mass1, double mass2)

Calculates the chirp mass from the two component masses.

- adouble calculate\_chirpmass (adouble mass1, adouble mass2)
- double calculate mass1 (double chirpmass, double eta)

Calculates the larger mass given a chirp mass and symmetric mass ratio.

- adouble calculate\_mass1 (adouble chirpmass, adouble eta)
- double calculate\_mass2 (double chirpmass, double eta)

Calculates the smaller mass given a chirp mass and symmetric mass ratio.

- adouble calculate\_mass2 (adouble chirpmass, adouble eta)
- void celestial\_horizon\_transform (double RA, double DEC, double gps\_time, double LONG, double LAT, double \*phi, double \*theta)

Utility to transform from celestial coord RA and DEC to local horizon coord for detector response functions.

double gps\_to\_GMST (double gps\_time)

 $\textit{Utility to transform from gps time to GMST} \quad \texttt{https://aa.usno.navy.mil/faq/docs/GAST.php.} \\$ 

• double gps\_to\_JD (double gps\_time)

Utility to transform from gps to JD.

void transform\_cart\_sph (double \*cartvec, double \*sphvec)

utility to transform a vector from cartesian to spherical (radian)

void transform\_sph\_cart (double \*sphvec, double \*cartvec)

utility to transform a vector from spherical (radian) to cartesian

• template<class T >

T trapezoidal sum uniform (double delta x, int length, T \*integrand)

Trapezoidal sum rule to approximate discrete integral - Uniform spacing.

• template<class T >

T trapezoidal sum (double \*delta x, int length, T \*integrand)

Trapezoidal sum rule to approximate discrete integral - Non-Uniform spacing.

• template<class T >

T simpsons\_sum (double delta\_x, int length, T \*integrand)

Simpsons sum rule to approximate discrete integral - Uniform spacing.

long factorial (long num)

Local function to calculate a factorial.

double pow\_int (double base, int power)

Local power function, specifically for integer powers.

- adouble pow\_int (adouble base, int power)
- template<class T >

std::complex< T > cpolar (T mag, T phase)

```
    template < class T >
        std::complex < T > XLALSpinWeightedSphericalHarmonic (T theta, T phi, int s, int I, int m)
```

• double cbrt\_internal (double base)

Fucntion that just returns the cuberoot.

• adouble cbrt\_internal (adouble base)

Fucntion that just returns the cuberoot ADOL-C doesn't have the cbrt function (which is faster), so have to use the power function.

### **Variables**

- const double gamma\_E = 0.5772156649015328606065120900824024310421
- const double c = 299792458.
- const double G = 6.674e-11\*(1.98855e30)
- const double MSOL\_SEC =4.925491025543575903411922162094833998e-6
- const double MPC\_SEC = 3.085677581491367278913937957796471611e22/c

### 9.16.1 Detailed Description

General utilities (functions and structures) independent of modelling method

### 9.16.2 Function Documentation

#### 9.16.2.1 allocate 2D array()

Utility to malloc 2D array.

## 9.16.2.2 allocate\_3D\_array()

Utility to malloc 3D array.

### 9.16.2.3 allocate\_LOSC\_data()

```
void allocate_LOSC_data (
    std::string * data_files,
    std::string psd_file,
    int num_detectors,
    int psd_length,
    int data_file_length,
    double trigger_time,
    std::complex< double > ** data,
    double ** psds,
    double ** freqs )
```

Prepare data for MCMC directly from LIGO Open Science Center.

Trims data for Tobs (determined by PSD file) 3/4\*Tobs in front of trigger, and 1/4\*Tobs behind

Currently, default to sampling frequency and observation time set by PSD - cannot be customized

Output is in order of PSD columns - string vector of detectos MUST match order of PSD cols

Output shapes—psds = [num\_detectors][psd\_length] data = [num\_detectors][psd\_length]

freqs = [num\_detectors][psd\_length]

Total observation time = 1/( freq[i] - freq[i-1]) (from PSD file)

Sampling frequency fs = max frequency from PSD file

ALLOCATES MEMORY - must be freed to prevent memory leak

### Parameters

	data_files	Vector of strings for each detector file from LOSC
psd_file		String of psd file from LOSC
	num_detectors	Number of detectors to use
	psd_length	Length of the PSD file (number of rows of DATA)
	data_file_length	Length of the data file (number of rows of DATA)
	trigger_time	Time for the signal trigger (GPS)
out	data	Output array of data for each detector
out	psds	Output array of psds for each detector
out	freqs	Output array of freqs for each detector

### 9.16.2.4 calculate\_chirpmass()

Calculates the chirp mass from the two component masses.

The output units are whatever units the input masses are

### 9.16.2.5 calculate\_mass1()

Calculates the larger mass given a chirp mass and symmetric mass ratio.

Units of the output match the units of the input chirp mass

### 9.16.2.6 calculate\_mass2()

Calculates the smaller mass given a chirp mass and symmetric mass ratio.

Units of the output match the units of the input chirp mass

### 9.16.2.7 celestial\_horizon\_transform()

Utility to transform from celestial coord RA and DEC to local horizon coord for detector response functions.

Outputs are the spherical polar angles defined by North as 0 degrees azimuth and the normal to the earth as 0 degree polar

#### **Parameters**

	RA	Right acsension (rad)
	DEC	Declination (rad)
	gps_time	GPS time
	LONG	Longitude (rad)
	LAT	Latitude (rad)
out	phi	horizon azimuthal angle (rad)
out	theta	horizon polar angle (rad)

### 9.16.2.8 cosmology\_interpolation\_function()

```
{\tt double\ cosmology\_interpolation\_function\ (}
```

```
double x,
double * coeffs,
int interp_degree )
```

Custom interpolation function used in the cosmology calculations.

Power series in half power increments of x, up to 11/2. powers of x

### 9.16.2.9 deallocate\_2D\_array()

Utility to free malloc'd 2D array.

#### 9.16.2.10 deallocate\_3D\_array() [1/2]

Utility to free malloc'd 2D array.

#### 9.16.2.11 deallocate\_3D\_array() [2/2]

```
void deallocate_3D_array (
          int *** array,
          int dim1,
          int dim2,
          int dim3 )
```

Utility to free malloc'd 2D array.

### 9.16.2.12 DL\_from\_Z()

Calculates the luminosity distance given the redshift.

Based on Astropy.cosmology calculations – see python script in the ./data folder of the project – numerically calculated given astropy.cosmology's definitions ( http://docs.astropy.org/en/stable/cosmology/) and used scipy.optimize to fit to a power series, stepping in half powers of Z. These coefficients are then output to a header file (D\_Z\_config.h) which are used here to calculate distance. Custom cosmologies etc can easily be acheived by editing the python script D\_Z\_config.py, the c++ functions do not need modification. They use whatever data is available in the header file. If the functional form of the fitting function changes, these functions DO need to change.

5 cosmological models are available (this argument must be spelled exactly):

PLANCK15, PLANCK13, WMAP9, WMAP7, WMAP5

### 9.16.2.13 free\_LOSC\_data()

/brief Free data allocated by prep\_LOSC\_data function

### 9.16.2.14 initiate\_LumD\_Z\_interp()

Function that uses the GSL libraries to interpolate pre-calculated Z-D\_L data.

Initiates the requried functions – GSL interpolation requires allocating memory before hand

## 9.16.2.15 pow\_int()

Local power function, specifically for integer powers.

Much faster than the std version, because this is only for integer powers

### 9.16.2.16 printProgress()

**9.16.2.17** read\_file() [1/2]

routine to print the progress of a process to the terminal as a progress bar

Call everytime you want the progress printed

int rows,

int cols )

Takes filename, and assigns to output[rows][cols]

double \*\* output,

File must be comma separated doubles

Utility to read in data.

### **Parameters**

	filename	input filename, relative to execution directory
out	output	array to store output, dimensions rowsXcols
	rows	first dimension
	cols	second dimension

Utility to read in data (single dimension vector)

Takes filename, and assigns to output[i\*rows + cols]

Output vector must be long enough, no check is done for the length

File must be comma separated doubles

### **Parameters**

	filename	input filename, relative to execution directory
out	output	output array, assumed to have the proper length of total items

```
9.16.2.19 read_LOSC_data_file()
```

Read data file from LIGO Open Science Center.

Convenience function for cutting off the first few lines of text

### **Parameters**

	filename	input filename
out	output	Output data
out	data_start_time	GPS start time of the data in file
out	duration	Duration of the signal
out	fs	Sampling frequency of the data

```
9.16.2.20 read_LOSC_PSD_file()
```

Read PSD file from LIGO Open Science Center.

Convenience function for cutting off the first few lines of text

### 9.16.2.21 simpsons\_sum()

Simpsons sum rule to approximate discrete integral - Uniform spacing.

More accurate than the trapezoidal rule, but must be uniform

### 9.16.2.22 transform\_cart\_sph()

utility to transform a vector from cartesian to spherical (radian)

order:

cart: x, y, z

spherical: r, polar, azimuthal

#### 9.16.2.23 transform\_sph\_cart()

utility to transform a vector from spherical (radian) to cartesian

order:

cart: x, y, z

spherical: r, polar, azimuthal

### 9.16.2.24 trapezoidal\_sum()

Trapezoidal sum rule to approximate discrete integral - Non-Uniform spacing.

This version is slower than the uniform version, but will handle non-uniform spacing

### 9.16.2.25 trapezoidal\_sum\_uniform()

Trapezoidal sum rule to approximate discrete integral - Uniform spacing.

This version is faster than the general version, as it has half the function calls

Something may be wrong with this function - had an overall offset for real data that was fixed by using the simpsons rule - not sure if this was because of a boost in accuracy or because something is off with the trapezoidal sum

### 9.16.2.26 tukey\_window()

Tukey window function for FFTs.

As defined by https://en.wikipedia.org/wiki/Window\_function

Utility to write 2D array to file.

Grid of data, comma separated

Grid has rows rows and cols columns

### **Parameters**

filename	Filename of output file, relative to execution directory	
input	Input 2D array pointer array[rows][cols]	
rows	First dimension of array	
cols	second dimension of array	

Utility to write 1D array to file.

Single column of data

### **Parameters**

filename	Filename of output file, relative to execution directory	
input	input 1D array pointer array[length]	
length length of array		

### 9.16.2.29 XLALSpinWeightedSphericalHarmonic()

## Shamelessly stolen from LALsuite

## Parameters

theta	polar angle (rad)	
phi azimuthal angle (rac		
s	spin weight	
1	mode number l	
m	mode number m	

### 9.16.2.30 Z\_from\_DL()

Calculates the redshift given the luminosity distance.

Based on Astropy.cosmology calculations – see python script in the ./data folder of the project – numerically calculated given astropy.cosmology's definitions ( <a href="http://docs.astropy.org/en/stable/cosmology/">http://docs.astropy.org/en/stable/cosmology/</a>) and used scipy.optimize to fit to a power series, stepping in half powers of DL. These coefficients are then output to a header file (D\_Z\_config.h) which are used here to calculate redshift. Custom cosmologies etc can easily be acheived by editing the python script D\_Z\_config.py, the c++ functions do not need modification. They use whatever data is available in the header file.

5 cosmological models are available (this argument must be spelled exactly, although case insensitive):

PLANCK15, PLANCK13, WMAP9, WMAP7, WMAP5

Function that returns Z from a given luminosity Distance - only Planck15

adouble version for ADOL-C calculations

Function that returns Z from a given luminosity Distance – only Planck15

### 9.16.3 Variable Documentation

```
9.16.3.1 c

const double c = 299792458.
```

Speed of light m/s

### 9.16.3.2 G

```
const double G = 6.674e - 11*(1.98855e30)
```

Gravitational constant in m\*\*3/(s\*\*2 SolMass)

### 9.16.3.3 gamma\_E

```
const double gamma_E = 0.5772156649015328606065120900824024310421
```

Euler number

### 9.16.3.4 MPC\_SEC

```
const double MPC_SEC = 3.085677581491367278913937957796471611e22/c
```

consts.kpc.to('m')\*1000/c Mpc in sec

### 9.16.3.5 MSOL\_SEC

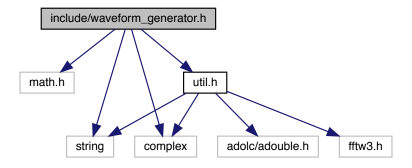
```
const double MSOL_SEC =4.925491025543575903411922162094833998e-6
```

G/c\*\*3 seconds per solar mass

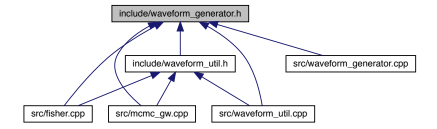
## 9.17 include/waveform\_generator.h File Reference

```
#include <math.h>
#include "util.h"
#include <complex>
#include <string>
```

Include dependency graph for waveform\_generator.h:



This graph shows which files directly or indirectly include this file:



#### **Functions**

- int fourier\_waveform (double \*frequencies, int length, std::complex< double > \*waveform\_plus, std
   ::complex< double > \*waveform\_cross, std::string generation\_method, gen\_params \*parameters)
- int fourier\_waveform (double \*frequencies, int length, double \*waveform\_plus\_real, double \*waveform
   \_plus\_imag, double \*waveform\_cross\_real, double \*waveform\_cross\_imag, std::string generation\_method,
   gen\_params \*parameters)
- int **fourier\_waveform** (double \*frequencies, int length, std::complex< double > \*waveform, std::string generation\_method, gen\_params \*parameters)
- int **fourier\_waveform** (double \*frequencies, int length, double \*waveform\_real, double \*waveform\_imag, std::string generation\_method, gen\_params \*parameters)
- int **fourier\_amplitude** (double \*frequencies, int length, double \*amplitude, std::string generation\_method, gen\_params \*parameters)
- int **fourier\_phase** (double \*frequencies, int length, double \*phase, std::string generation\_method, gen\_params \*parameters)

## 9.18 include/waveform generator C.h File Reference

### **Functions**

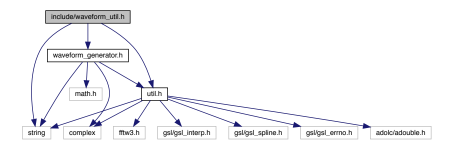
- int fourier\_waveformC (double \*frequencies, int length, double \*waveform\_plus\_real, double \*waveform\_cross\_real, double \*waveform\_cross\_imag, char \*generation\_method, double mass1, double mass2, double DL, double spin1x, double spin1y, double spin1z, double spin2x, double spin2x, double spin2z, double spin2z, double tc, double f\_ref, double phiRef, double \*ppE\_beta, int \*ppE\_b, int Nmod, double incl\_angle, double theta, double phi)
- int **fourier\_amplitudeC** (double \*frequencies, int length, double \*amplitude, char \*generation\_method, double mass1, double mass2, double DL, double spin1x, double spin1y, double spin1z, double spin2x, double spin2x, double incl\_angle, double theta, double phi)
- int fourier\_phaseC (double \*frequencies, int length, double \*phase, char \*generation\_method, double mass1, double mass2, double DL, double spin1x, double spin1y, double spin1z, double spin2x, double spin2x, double spin2x, double spin2z, double phic, double tc, double f\_ref, double phiRef, double \*ppE\_beta, int \*ppE\_b, int Nmod, double incl\_angle, double theta, double phi)
- void initiate\_LumD\_Z\_interp\_C ()
- void free\_LumD\_Z\_interp\_C ()

### 9.18.1 Detailed Description

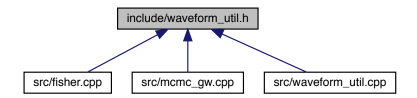
Header file for the C wrapping of the waveform\_generation.cpp

## 9.19 include/waveform\_util.h File Reference

```
#include "waveform_generator.h"
#include "util.h"
#include <string>
Include dependency graph for waveform util.h:
```



This graph shows which files directly or indirectly include this file:



### **Functions**

- double data\_snr\_maximized\_extrinsic (double \*frequencies, int length, std::complex< double > \*data, double \*psd, std::string detector, std::string generation\_method, gen\_params \*param)
  - Utility to calculate the snr of a fourier transformed data stream while maximizing over the coalescence parameters phic and tc.
- double data\_snr\_maximized\_extrinsic (double \*frequencies, int length, double \*data\_real, double \*data\_
  imag, double \*psd, std::string detector, std::string generation\_method, gen\_params \*param)
  - Light wrapper for the data\_snr\_maximized\_extrinsic method.
- double calculate\_snr (std::string detector, std::complex< double > \*waveform, double \*frequencies, int length)
  - Caclulates the snr given a detector and waveform (complex) and frequencies.
- int fourier\_detector\_response (double \*frequencies, int length, std::complex< double > \*hplus, std
   ::complex< double > \*hcross, std::complex< double > \*detector\_response, double theta, double phi, std
   ::string detector)
- int fourier\_detector\_response (double \*frequencies, int length, std::complex< double > \*hplus, std
   ::complex< double > \*hcross, std::complex< double > \*detector\_response, double theta, double phi, double psi, std::string detector)

• int fourier\_detector\_response\_equatorial (double \*frequencies, int length, std::complex< double > \*hplus, std::complex< double > \*hcross, std::complex< double > \*detector\_response, double ra, double dec, double psi, double gmst, std::string detector)

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary.

• int fourier\_detector\_response\_equatorial (double \*frequencies, int length, std::complex< double > \*response, std::string detector, std::string generation\_method, gen\_params \*parameters)

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary for equatorial coordinates.

• int fourier\_detector\_amplitude\_phase (double \*frequencies, int length, double \*amplitude, double \*phase, std::string detector, std::string generation\_method, gen\_params \*parameters)

Calculates the amplitude (magnitude) and phase (argument) of the response of a given detector.

### 9.19.1 Detailed Description

Header file for waveform specific utilites

#### 9.19.2 Function Documentation

### 9.19.2.1 calculate\_snr()

Caclulates the snr given a detector and waveform (complex) and frequencies.

This function computes the un-normalized snr: \sqrt( ( H | H ) )

#### **Parameters**

detector	detector name - must match the string of populate_noise precisely	
waveform	complex waveform	
frequencies	double array of frequencies that the waveform is evaluated at	
length	ength length of the above two arrays	

```
9.19.2.2 data_snr_maximized_extrinsic() [1/2]
```

```
int length,
std::complex< double > * data,
double * psd,
std::string detector,
std::string generation_method,
gen_params * param )
```

Utility to calculate the snr of a fourier transformed data stream while maximizing over the coalescence parameters phic and tc.

The gen\_params structure holds the parameters for the template to be used (the maximimum likelihood parameters)

#### **Parameters**

frequencies	Frequencies used by data	
length	length of the data	
data	input data in the fourier domain	
psd	PSD for the detector that created the data	
detector	Name of the detector –See noise_util for options	
generation_method	Generation method for the template – See waveform_generation.cpp for options	
param	gen_params structure for the template	

### 9.19.2.3 data\_snr\_maximized\_extrinsic() [2/2]

Light wrapper for the data\_snr\_maximized\_extrinsic method.

Splits the data into real and imaginary, so all the arguments are C-safe

### **Parameters**

- aramotoro		
frequencies	Frequencies used by data	
length	length of the data	
data_real	input data in the fourier domain – real part	
data_imag	input data in the fourier domain – imaginary part	
psd	PSD for the detector that created the data	
detector	Name of the detector –See noise_util for options	
generation_method	Generation method for the template – See waveform_generation.cpp for options	
param	gen_params structure for the template	

### 9.19.2.4 fourier\_detector\_amplitude\_phase()

Calculates the amplitude (magnitude) and phase (argument) of the response of a given detector.

This is for general waveforms, and will work for precessing waveforms

Not as fast as non-precessing, but that can't be helped. MUST include plus/cross polarizations

### 9.19.2.5 fourier\_detector\_response() [1/3]

#### **Parameters**

	frequencies	array of frequencies corresponding to waveform
	length	length of frequency/waveform arrays
	hcross	precomputed cross polarization of the waveform
out	detector_response	detector response
	theta	polar angle (rad) theta in detector frame
	phi	azimuthal angle (rad) phi in detector frame
	detector	detector - list of supported detectors in noise_util

### 9.19.2.6 fourier\_detector\_response() [2/3]

#### **Parameters**

	frequencies	array of frequencies corresponding to waveform
	length	length of frequency/waveform arrays
	hcross	precomputed cross polarization of the waveform
out	detector_response	detector response
	theta	polar angle (rad) theta in detector frame
	phi	azimuthal angle (rad) phi in detector frame
	psi	polarization angle (rad) phi in detector frame
	detector	detector - list of supported detectors in noise_util

### 9.19.2.7 fourier\_detector\_response() [3/3]

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

This puts the responsibility on the user to pass the necessary parameters

Detector options include classic interferometers like LIGO/VIRGO (coming soon: ET and LISA)

This is a wrapper that combines generation with response functions: if producing mulitple responses for one waveform (ie stacking Hanford, Livingston, and VIRGO), it will be considerably more efficient to calculate the waveform once, then combine each response manually

#### **Parameters**

	frequencies	double array of frequencies for the waveform to be evaluated at
	length	integer length of all the arrays
out	response	complex array for the output plus polarization waveform
	generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY
	parameters	structure containing all the source parameters

### 9.19.2.8 fourier\_detector\_response\_equatorial() [1/2]

```
int length,
std::complex< double > * hplus,
std::complex< double > * hcross,
std::complex< double > * detector_response,
double ra,
double dec,
double psi,
double gmst,
std::string detector )
```

#### **Parameters**

	frequencies	array of frequencies corresponding to waveform
	length	length of frequency/waveform arrays
	hcross	precomputed cross polarization of the waveform
out	detector_response	detector response
	ra	Right Ascension in rad
	dec	Declination in rad
	psi	polarization angle (rad)
	gmst	greenwich mean sidereal time
	detector	detector - list of supported detectors in noise_util

### 9.19.2.9 fourier\_detector\_response\_equatorial() [2/2]

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary for equatorial coordinates.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

This puts the responsibility on the user to pass the necessary parameters

Detector options include classic interferometers like LIGO/VIRGO (coming soon: ET and LISA)

This is a wrapper that combines generation with response functions: if producing mulitple responses for one waveform (ie stacking Hanford, Livingston, and VIRGO), it will be considerably more efficient to calculate the waveform once, then combine each response manually

### **Parameters**

	frequencies	double array of frequencies for the waveform to be evaluated at
	length	integer length of all the arrays
out	response	complex array for the output plus polarization waveform
	generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY
	parameters	structure containing all the source parameters

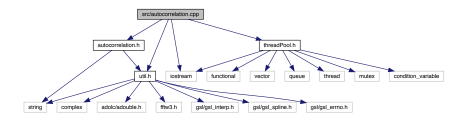
Generated by Doxygen

### 9.20 README.dox File Reference

## 9.21 src/autocorrelation.cpp File Reference

```
#include "autocorrelation.h"
#include "util.h"
#include "threadPool.h"
#include <iostream>
```

Include dependency graph for autocorrelation.cpp:



### **Macros**

• #define MAX SERIAL 200000

### **Functions**

- void write\_auto\_corr\_file\_from\_data\_file (std::string autocorr\_filename, std::string datafile, int length, int dimension, int num\_segments, double target\_corr, int num\_threads)
- void write\_auto\_corr\_file\_from\_data (std::string autocorr\_filename, double \*\*data, int length, int dimension, int num\_segments, double target\_corr, int num\_threads)

Writes the autocorrelation file from a data array.

• void auto\_corr\_from\_data (double \*\*data, int length, int dimension, int \*\*output, int num\_segments, double target\_corr, int num\_threads)

Calculates the autocorrelation length for a set of data for a number of segments for each dimension – completely host code, utilitizes FFTW3 for longer chuncks of the chains.

void threaded\_ac\_spectral (int thread, threaded\_ac\_jobs\_fft job)

Internal routine to calculate an spectral autocorrelation job.

void threaded\_ac\_serial (int thread, threaded\_ac\_jobs\_serial job)

Internal routine to calculate an serial autocorrelation job.

double auto\_correlation\_serial (double \*arr, int length, int start, double target)

Calculates the autocorrelation of a chain with the brute force method.

void auto\_correlation\_spectral (double \*chain, int length, double \*autocorr, fftw\_outline \*plan\_forw, fftw outline \*plan rev)

Wrapper function for convience - assumes the data array starts at 0.

 void auto\_correlation\_spectral (double \*chain, int length, int start, double \*autocorr, fftw\_outline \*plan\_forw, fftw\_outline \*plan\_rev)

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT – accepts FFTW plan as argument for plan reuse and multi-threaded applications.

void auto\_correlation\_spectral (double \*chain, int length, double \*autocorr)

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT.

• double auto\_correlation (double \*arr, int length, double tolerance)

OUTDATED - numerically finds autocorrelation length - not reliable.

• double auto\_correlation\_serial\_old (double \*arr, int length)

OUTDATED Calculates the autocorrelation – less general version.

double auto\_correlation\_grid\_search (double \*arr, int length, int box\_num, int final\_length, double target\_
 length)

OUTDATED - Grid search method of computing the autocorrelation - unreliable.

double auto\_correlation\_internal (double \*arr, int length, int lag, double ave)

Internal function to compute the auto correlation for a given lag.

void auto\_corr\_intervals\_outdated (double \*data, int length, double \*output, int num\_segments, double accuracy)

Function that computes the autocorrelation length on an array of data at set intervals to help determine convergence.

• void write\_auto\_corr\_file\_from\_data (std::string auto\_corr\_filename, double \*\*output, int intervals, int dimension, int N\_steps)

OUTDATED – writes autocorrelation lengths for a data array, but only with the serial method and only for a target correlation of .01.

void write\_auto\_corr\_file\_from\_data\_file (std::string auto\_corr\_filename, std::string output\_file, int intervals, int dimension, int N steps)

OUTDATED – writes autocorrelation lengths for a data file, but only with the serial method and only for a target correlation of .01.

### 9.21.1 Detailed Description

Turns out calculating the autocorrelation is more complicated if you want to do it fast, so it gets its own file now

### 9.21.2 Macro Definition Documentation

```
9.21.2.1 MAX SERIAL
```

```
#define MAX_SERIAL 200000
```

Max length of array to use serial calculation

#### 9.21.3 Function Documentation

#### 9.21.3.1 auto\_corr\_from\_data()

Calculates the autocorrelation length for a set of data for a number of segments for each dimension – completely host code, utilitizes FFTW3 for longer chuncks of the chains.

Takes in the data from a sampler, shape data[N\_steps][dimension]

Outputs lags that correspond to the target\_corr – shape output[dimension][num\_segments]

#### **Parameters**

	data	Input data	
	length	length of input data	
	dimension	dimension of data	
out	output	array that stores the auto-corr lengths – array[num_segments]	
	num_segments	number of segements to compute the auto-corr length	
	target_corr	Autocorrelation for which the autocorrelation length is defined (lag of autocorrelation for which it equals the target_corr)	
	num_threads	Total number of threads to use	

# 9.21.3.2 auto\_corr\_intervals\_outdated()

Function that computes the autocorrelation length on an array of data at set intervals to help determine convergence.

outdated version - new version uses FFTs

#### **Parameters**

	data	Input data
	length	length of input data
out	output array that stores the auto-corr lengths – array[num_segments]	
	num_segments   number of segements to compute the auto-corr length	
	accuracy	longer chains are computed numerically, this specifies the tolerance

## 9.21.3.3 auto\_correlation\_grid\_search()

OUTDATED – Grid search method of computing the autocorrelation – unreliable.

Hopefully more reliable than the box-search method, which can sometimes get caught in a recursive loop when the stepsize isn't tuned, but also faster than the basic linear, serial search

#### **Parameters**

arr	Input array to use for autocorrelation
length	Length of input array
box_num	number of boxes to use for each iteration, default is 10
final_length	number of elements per box at which the grid search ends and the serial calculation begins
target_length	target correlation that corresponds to the returned lag

## 9.21.3.4 auto\_correlation\_internal()

Internal function to compute the auto correlation for a given lag.

# 9.21.3.5 auto\_correlation\_serial()

Calculates the autocorrelation of a chain with the brute force method.

## **Parameters**

arr	input array
length	Length of input array
start	starting index (probably 0)
target	Target autocorrelation for which `'length" is defined

## 9.21.3.6 auto\_correlation\_spectral() [1/2]

```
fftw_outline * plan_forw,
fftw_outline * plan_rev )
```

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT – accepts FFTW plan as argument for plan reuse and multi-threaded applications.

Based on the Wiener-Khinchin Theorem.

```
Algorithm used from https://lingpipe-blog.com/2012/06/08/autocorrelation-fft-kiss-eigen/
```

NOTE the length used in initializing the fftw plans should be L = pow(2, std::ceil( std::log2(length) ) ) – the plans are padded so the total length is a power of two

Option to provide starting index for multi-dimension arrays in collapsed to one dimension

length is the length of the segment to be analyzed, not necessarily the dimension of the chain

```
9.21.3.7 auto_correlation_spectral() [2/2]
```

Faster approximation of the autocorrelation of a chain. Implements FFT/IFFT.

Based on the Wiener-Khinchin Theorem.

Algorithm used from https://lingpipe-blog.com/2012/06/08/autocorrelation-fft-kiss-eigen/

#### 9.21.3.8 threaded\_ac\_serial()

Internal routine to calculate an serial autocorrelation job.

Allows for a more efficient use of the threadPool class

# 9.21.3.9 threaded\_ac\_spectral()

Internal routine to calculate an spectral autocorrelation job.

Allows for a more efficient use of the threadPool class

# 9.21.3.10 write\_auto\_corr\_file\_from\_data()

```
void write_auto_corr_file_from_data (
    std::string autocorr_filename,
    double ** data,
    int length,
    int dimension,
    int num_segments,
    double target_corr,
    int num_threads )
```

Writes the autocorrelation file from a data array.

#### **Parameters**

autocorr_filename	Name of the file to write the autocorrelation to
data	Input chains
length	length of input data
dimension	dimension of data
num_segments	number of segements to compute the auto-corr length
target_corr	Autocorrelation for which the autocorrelation length is defined (lag of autocorrelation for which it equals the target_corr)
num_threads	Total number of threads to use

#### 9.21.3.11 write\_auto\_corr\_file\_from\_data\_file()

```
void write_auto_corr_file_from_data_file (
    std::string autocorr_filename,
    std::string datafile,
    int length,
    int dimension,
    int num_segments,
    double target_corr,
    int num_threads )
```

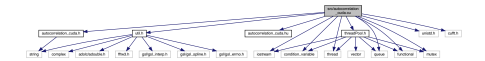
#### **Parameters**

length	length of input data	
dimension	dimension of data	
num_segments	number of segements to compute the auto-corr length	
target_corr	Autocorrelation for which the autocorrelation length is defined (lag of autocorrelation for which it equals the target_corr)	
num_threads	Total number of threads to use	

# 9.22 src/autocorrelation\_cuda.cu File Reference

```
#include "autocorrelation_cuda.h"
#include "autocorrelation_cuda.hu"
#include "util.h"
#include <iostream>
#include <condition_variable>
#include <thread>
#include <queue>
#include <functional>
#include <mutex>
#include <unistd.h>
#include <threadPool.h>
#include <cufft.h>
```

Include dependency graph for autocorrelation\_cuda.cu:



#### **Functions**

\_\_device\_\_ \_\_host\_\_ void auto\_corr\_internal (double \*arr, int length, int lag, double average, double \*corr, int start\_id)

Internal function to calculate the autocorrelation for a given lag Customized for the thread pool architecture, with extra arguments because of the way the memory is allocated.

• \_\_global\_\_ void auto\_corr\_internal\_kernal (double \*arr, int length, double average, int \*rho\_index, double target\_corr, double var, int start\_id)

Internal function to launch the CUDA kernel for a range of autocorrelations.

void write\_file\_auto\_corr\_from\_data\_file\_accel (std::string acfile, std::string chains\_file, int dimension, int N
 \_ steps, int num\_segments, double target\_corr)

Write data file for autocorrelation lengths of the data given a data file name, as written by the mcmc\_sampler.

• void write\_file\_auto\_corr\_from\_data\_accel (std::string acfile, double \*\*chains, int dimension, int N\_steps, int num segments, double target corr)

Write data file given output chains, as formatted by the mcmc\_sampler.

 void auto\_corr\_from\_data\_accel (double \*\*output, int dimension, int N\_steps, int num\_segments, double target\_corr, double \*\*autocorr)

Find autocorrelation of data at different points in the chain length and output to autocorr.

void ac\_gpu\_wrapper (int thread, int job\_id)

Wrapper function for the thread pool.

• void <a href="mailto:launch\_ac\_gpu">launch\_ac\_gpu</a> (int device, int element, double \*\*data, int length, int dimension, double target\_corr, int num segments)

Launch the GPU kernel, formatted for the thread pool.

• void allocate gpu plan (GPUplan \*plan, int data length, int dimension, int num segments)

Allocates memory for autocorrelation-GPU structure.

• void deallocate\_gpu\_plan (GPUplan \*plan, int data\_length, int dimension, int num\_segments)

Deallocates memory for the autocorrelation-GPU structure.

Copy data to device before starting kernels.

## **Variables**

GPUplan \* plans\_global

#### 9.22.1 Function Documentation

# 9.22.1.1 ac\_gpu\_wrapper()

```
void ac_gpu_wrapper (
          int thread,
          int job_id )
```

Wrapper function for the thread pool.

## **Parameters**

thread	Host thread
job⊷	Job ID
_id	

# 9.22.1.2 allocate\_gpu\_plan()

Allocates memory for autocorrelation-GPU structure.

#### **Parameters**

plan	Structure for GPU plan
data_length	Length of data
dimension	Dimension of the data
num_segments	Number of segments to calculate the autocorrelation length

## 9.22.1.3 auto\_corr\_from\_data\_accel()

Find autocorrelation of data at different points in the chain length and output to autocorr.

	output	Chain data input
	dimension	Dimension of the data
	N_steps	Number of steps in the data
	num_segments	number of segments to calculate the autocorrelation length
	target_corr	Target correlation ratio
out	autocorr	Autocorrelation lengths for the different segments

#### 9.22.1.4 auto\_corr\_internal()

Internal function to calculate the autocorrelation for a given lag Customized for the thread pool architecture, with extra arguments because of the way the memory is allocated.

#### **Parameters**

	arr	Input array of data
	length	Length of input array
	lag	Lag to be used to calculate the correlation
	average	Average of the array arr
out	corr	output correlation
	start_id	ID of location to start calculation – input arrary arr is assumed to be contiguous for multiple
		dimensions

## 9.22.1.5 auto\_corr\_internal\_kernal()

Internal function to launch the CUDA kernel for a range of autocorrelations.

Correlation function used:

```
\label{eq:rho(lag) = 1 / (length - lag) \sum (arr[i+lag]-average) (arr[i]-average)} \\
```

```
target\_corr = rho(rho\_index)/rho(0) = rho(rho\_index)/var
```

	arr	Input array of data
	length	Length of data array
	average	Average of input data
out	rho_index Index of the lag that results ina correlation ratio target_cor	
	target_corr	
	var	Variance rho(0)
	start_id	Starting index to use for the data array arr

# 9.22.1.6 copy\_data\_to\_device()

Copy data to device before starting kernels.

## **Parameters**

plan	GPU plan
input_data	Input chain data
data_length	Length of data
dimension	Dimension of the data
num_segments	Number of segments to calculate the autocorrelation length

# 9.22.1.7 deallocate\_gpu\_plan()

Deallocates memory for the autocorrelation-GPU structure.

## **Parameters**

plan	Structure for the GPU plan	
data_length	Length of data	
dimension Dimension of the data		
num_segments	Number of segments to calculate the autocorrelation length	

# 9.22.1.8 write\_file\_auto\_corr\_from\_data\_accel()

```
int num_segments,
double target_corr )
```

Write data file given output chains, as formatted by the mcmc\_sampler.

#### **Parameters**

acfile	Output autocorrelation filename	
chains	Chain data from MCMC_sampler	
dimension	Dimension of the data	
N_steps	Number of steps in the chain	
num_segments	Number of segments to check the autocorrelation length for each dimension	
target_corr	Target correlation ratio to use for the correlation length calculation	

## 9.22.1.9 write\_file\_auto\_corr\_from\_data\_file\_accel()

```
void write_file_auto_corr_from_data_file_accel (
    std::string acfile,
    std::string chains_file,
    int dimension,
    int N_steps,
    int num_segments,
    double target_corr )
```

Write data file for autocorrelation lengths of the data given a data file name, as written by the mcmc\_sampler.

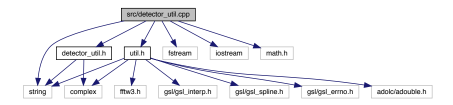
## **Parameters**

acfile	Filename of the autocorrelation data	
chains_file	Filename of the data file for the chains	
dimension	Dimension of the data	
N_steps	Number of steps in the chain	
num_segments	Number of segments to check the autocorrelation length for each dimension	
target_corr	Target correlation ratio to use for the correlation length calculation	

# 9.23 src/detector\_util.cpp File Reference

```
#include "detector_util.h"
#include "util.h"
#include <fstream>
#include <iostream>
#include <string>
#include <math.h>
```

Include dependency graph for detector\_util.cpp:



#### **Functions**

- void populate\_noise (double \*frequencies, std::string detector, double \*noise\_root, int length)
- double aLIGO analytic (double f)

Analytic function approximating the PSD for aLIGO.

double Hanford O1 fitted (double f)

Numerically fit PSD to the Hanford Detector's O1.

• std::complex< double > Q (double theta, double phi, double iota)

Utility for the overall amplitude and phase shift for spin-aligned systems.

Function to populate the squareroot of the noise curve for various detectors.

• double right interferometer plus (double theta, double phi)

Response function of a 90 deg interferometer for plus polarization.

double right\_interferometer\_cross (double theta, double phi)

Response function of a 90 deg interferometer for cross polarization.

• void celestial\_horizon\_transform (double RA, double DEC, double gps\_time, std::string detector, double \*phi, double \*theta)

Transform from celestial coordinates to local horizontal coords.

• void derivative\_celestial\_horizon\_transform (double RA, double DEC, double gps\_time, std::string detector, double \*dphi\_dRA, double \*dtheta\_dRA, double \*dphi\_dDEC, double \*dtheta\_dDEC)

Numerical derivative of the transformation.

- double DTOA (double theta1, double theta2, std::string detector1, std::string detector2)
  - calculate difference in time of arrival (DTOA) for a given source location and 2 different detectors
- double radius\_at\_lat (double latitude, double elevation)
- void detector\_response\_functions\_equatorial (double D[3][3], double ra, double dec, double psi, double gmst, double \*Fplus, double \*Fcross)

Calculates the response coefficients for a detector with response tensor D for a source at RA, Dec, and psi.

• void detector\_response\_functions\_equatorial (std::string detector, double ra, double dec, double psi, double gmst, double \*Fplus, double \*Fcross)

Same as the other function, but for active detectors.

## 9.23.1 Detailed Description

Routines to construct noise curves for various detectors and for detector specific utilities for response functions and coordinate transformations

### 9.23.2 Function Documentation

### 9.23.2.1 aLIGO\_analytic()

```
double aLIGO_analytic ( \label{eq:double_f} \mbox{double } f \mbox{ )}
```

Analytic function approximating the PSD for aLIGO.

CITE (Will?)

## 9.23.2.2 celestial\_horizon\_transform()

Transform from celestial coordinates to local horizontal coords.

```
(RA,DEC) -> (altitude, azimuth)
```

Need gps\_time of transformation, as the horizontal coords change in time

detector is used to specify the lat and long of the local frame

# Parameters

RA	in RAD
DEC	in RAD
phi	in RAD
theta	in RAD

## 9.23.2.3 derivative\_celestial\_horizon\_transform()

Numerical derivative of the transformation.

Planned for use in Fisher calculations, but not currently implemented anywhere

#### **Parameters**

RA	in RAD
DEC	in RAD

## 9.23.2.4 detector\_response\_functions\_equatorial() [1/2]

Calculates the response coefficients for a detector with response tensor D for a source at RA, Dec, and psi.

## Taken from LALSuite

The response tensor for each of the operational detectors is precomputed in detector\_util.h, but to create a new tensor, follow the outline in Anderson et al 36 PRD 63 042003 (2001) Appendix B

#### **Parameters**

	D	Detector Response tensor (3x3)
ra Right ascension in rad		Right ascension in rad
	dec	Declination in rad
	psi	polarization angle in rad
	gmst	Greenwich mean sidereal time (rad)
out	Fplus	Fplus response coefficient
out	Fcross	Fcross response coefficient

### 9.23.2.5 detector\_response\_functions\_equatorial() [2/2]

```
void detector_response_functions_equatorial (
    std::string detector,
    double ra,
    double dec,
    double psi,
    double gmst,
    double * Fplus,
    double * Fcross )
```

Same as the other function, but for active detectors.

#### **Parameters**

	detector	Detector
	ra	Right ascension in rad
	dec	Declination in rad
	psi	polarization angle in rad
	gmst	Greenwich mean sidereal time (rad)
out	Fplus	Fplus response coefficient
out	Fcross	Fcross response coefficient

## 9.23.2.6 DTOA()

calculate difference in time of arrival (DTOA) for a given source location and 2 different detectors

#### **Parameters**

theta1	spherical polar angle for detector 1 in RAD
theta2	spherical polar angle for detector 2 in RAD
detector1	name of detector one
detector2	name of detector two

# 9.23.2.7 Hanford\_O1\_fitted()

```
double Hanford_01_fitted ( double f )
```

Numerically fit PSD to the Hanford Detector's O1.

CITE (Yunes?)

## 9.23.2.8 populate\_noise()

Function to populate the squareroot of the noise curve for various detectors.

If frequencies are left as NULL, standard frequency spacing is applied and the frequencies are returned, in which case the frequencies argument becomes an output array

Detector names must be spelled exactly

Detectors include: aLIGO\_analytic, Hanford\_O1\_fitted

#### **Parameters**

frequencies	double array of frquencies (NULL)
detector	String to designate the detector noise curve to be used
noise_root	ouptput double array for the square root of the PSD of the noise of the specified detector
length	integer length of the output and input arrays

#### 9.23.2.9 Q()

Utility for the overall amplitude and phase shift for spin-aligned systems.

For spin aligned, all the extrinsic parameters have the effect of an overall amplitude modulation and phase shift

# 9.23.2.10 radius\_at\_lat()

/brief Analytic approximation of the radius from the center of earth to a given location

Just the raidus as a function of angles, modelling an oblate spheroid

### **Parameters**

latitude	latitude in degrees
elevation	elevation in meters

## 9.23.2.11 right\_interferometer\_cross()

Response function of a 90 deg interferometer for cross polarization.

Theta and phi are local, horizontal coordinates relative to the detector

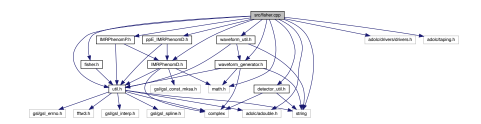
### 9.23.2.12 right\_interferometer\_plus()

Response function of a 90 deg interferometer for plus polarization.

Theta and phi are local, horizontal coordinates relative to the detector

# 9.24 src/fisher.cpp File Reference

```
#include <fisher.h>
#include <adolc/adouble.h>
#include <adolc/drivers/drivers.h>
#include <adolc/taping.h>
#include <math.h>
#include <string>
#include "util.h"
#include "detector_util.h"
#include "IMRPhenomD.h"
#include "IMRPhenomP.h"
#include "yppE_IMRPhenomD.h"
#include "waveform_generator.h"
#include "waveform_util.h"
Include dependency graph for fisher.cpp:
```



## **Functions**

• void fisher (double \*frequency, int length, string generation\_method, string detector, double \*\*output, int dimension, gen\_params \*parameters, int \*amp\_tapes, int \*phase\_tapes, double \*noise)

Calculates the fisher matrix for the given arguments.

• void calculate\_derivatives (double \*\*amplitude\_deriv, double \*\*phase\_deriv, double \*amplitude, double \*frequencies, int length, string detector, string gen\_method, gen\_params \*parameters)

Abstraction layer for handling the case separation for the different waveforms.

• void fisher\_autodiff (double \*frequency, int length, string generation\_method, string detector, double \*\*output, int dimension, gen\_params \*parameters, int \*amp\_tapes, int \*phase\_tapes, double \*noise)

Calculates the fisher matrix for the given arguments to within numerical error using automatic differention - slower than the numerical version.

# 9.24.1 Detailed Description

All subroutines associated with waveform differentiation and Fisher analysis

# 9.24.2 Function Documentation

## 9.24.2.1 calculate\_derivatives()

Abstraction layer for handling the case separation for the different waveforms.

#### 9.24.2.2 fisher()

Calculates the fisher matrix for the given arguments.

length	if 0, standard frequency range for the detector is used
output	double [dimension][dimension]
amp_tapes	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method
phase_tapes	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method

#### 9.24.2.3 fisher\_autodiff()

Calculates the fisher matrix for the given arguments to within numerical error using automatic differention - slower than the numerical version.

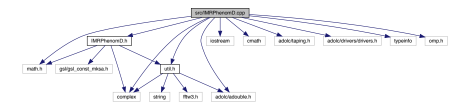
#### **Parameters**

length	if 0, standard frequency range for the detector is used
output	double [dimension][dimension]
amp_tapes	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method
phase_tapes	if speed is required, precomputed tapes can be used - assumed the user knows what they're doing, no checks done here to make sure that the number of tapes matches the requirement by the generation_method

# 9.25 src/IMRPhenomD.cpp File Reference

```
#include "IMRPhenomD.h"
#include "util.h"
#include <math.h>
#include <iostream>
#include <complex>
#include <cmath>
#include <adolc/adouble.h>
#include <adolc/taping.h>
#include <adolc/drivers/drivers.h>
#include <typeinfo>
#include <omp.h>
```

Include dependency graph for IMRPhenomD.cpp:



#### **Macros**

• #define omp ignore

## **Variables**

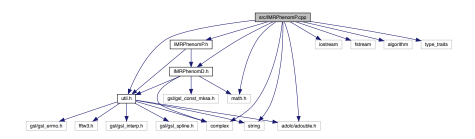
• double log\_64 = 4.15888308336

# 9.25.1 Detailed Description

File that includes all the low level functions that go into constructing the waveform

# 9.26 src/IMRPhenomP.cpp File Reference

```
#include "IMRPhenomP.h"
#include <iostream>
#include <fstream>
#include <string>
#include <complex>
#include "IMRPhenomD.h"
#include "util.h"
#include <adolc/adouble.h>
#include <math.h>
#include <algorithm>
#include <type_traits>
Include dependency graph for IMRPhenomP.cpp:
```



## **Macros**

- #define ROTATEZ(angle, vx, vy, vz)
- #define **ROTATEY**(angle, vx, vy, vz)

#### **Variables**

• const double **sqrt\_6** = 2.44948974278317788

# 9.26.1 Detailed Description

Source code for IMRPhenomP

#### 9.26.2 Macro Definition Documentation

#### 9.26.2.1 ROTATEY

#### Value:

```
tmp1 = vx*cos(angle) + vz*sin(angle);\
tmp2 = - vx*sin(angle) + vz*cos(angle);\
vx = tmp1;\
vz = tmp2
```

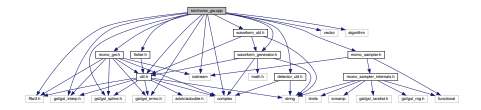
# 9.26.2.2 ROTATEZ

## Value:

```
tmp1 = vx*cos(angle) - vy*sin(angle);\
tmp2 = vx*sin(angle) + vy*cos(angle);\
vx = tmp1;\
vy = tmp2
```

# 9.27 src/mcmc\_gw.cpp File Reference

```
#include "mcmc_gw.h"
#include "waveform_generator.h"
#include "util.h"
#include "detector_util.h"
#include "waveform_util.h"
#include "fisher.h"
#include "mcmc_sampler.h"
#include <iostream>
#include <vector>
#include <complex>
#include <fftw3.h>
#include <algorithm>
#include <gsl/gsl_interp.h>
#include <gsl/gsl_spline.h>
#include <gsl/gsl_errno.h>
Include dependency graph for mcmc_gw.cpp:
```



#### **Functions**

double maximized\_coal\_log\_likelihood\_IMRPhenomD (double \*frequencies, int length, std::complex< double > \*data, double \*noise, double SNR, double chirpmass, double symmetric\_mass\_ratio, double spin1, double spin2, bool NSflag, fftw\_outline \*plan)

Function to calculate the log Likelihood as defined by -1/2 (d-h|d-h) maximized over the extrinsic parameters phic and tc.

- double maximized\_coal\_log\_likelihood\_IMRPhenomD (double \*frequencies, size\_t length, double \*real\_
   data, double \*imag\_data, double \*noise, double SNR, double chirpmass, double symmetric\_mass\_ratio, double spin1, double spin2, bool NSflag)
- double maximized\_coal\_log\_likelihood\_IMRPhenomD (double \*frequencies, size\_t length, double \*real\_
  data, double \*imag\_data, double \*noise, double SNR, double chirpmass, double symmetric\_mass\_ratio,
  double spin1, double spin2, bool NSflag, fftw outline \*plan)
- double maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param (double \*frequencies, int length, std
   ::complex < double > \*data, double \*noise, double chirpmass, double symmetric\_mass\_ratio, double spin1,
   double spin2, double Luminosity\_Distance, double theta, double phi, double iota, bool NSflag, fftw\_outline
   \*plan)
- double maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param (double \*frequencies, size\_t length, double \*real\_data, double \*imag\_data, double \*noise, double chirpmass, double symmetric\_mass\_ratio, double spin1, double spin2, double Luminosity\_Distance, double theta, double phi, double iota, bool NSflag)
- double maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param (double \*frequencies, size\_t length, double \*real\_data, double \*imag\_data, double \*noise, double chirpmass, double symmetric\_mass\_ratio, double spin1, double spin2, double Luminosity\_Distance, double theta, double phi, double iota, bool NSflag, fftw\_outline \*plan)
- double maximized\_Log\_Likelihood (std::complex < double > \*data, double \*psd, double \*frequencies, size ←
   \_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw\_outline \*plan)

routine to maximize over all extrinsic quantities and return the log likelihood

- double maximized\_Log\_Likelihood (double \*data\_real, double \*data\_imag, double \*psd, double \*frequencies, size\_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw\_outline \*plan)
- double maximized\_coal\_Log\_Likelihood (std::complex < double > \*data, double \*psd, double \*frequencies, size\_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw\_outline \*plan, double \*tc, double \*phic)

Function to maximize only over coalescence variables to and phic, returns the maximum values used.

- double maximized\_coal\_Log\_Likelihood\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std::complex< double > \*detector\_response, size\_t length, fftw\_outline \*plan, double \*tc, double \*phic)
- double Log\_Likelihood (std::complex< double > \*data, double \*psd, double \*frequencies, size\_t length, gen\_params \*params, std::string detector, std::string generation\_method, fftw\_outline \*plan)

Unmarginalized log of the likelihood.

• double maximized\_Log\_Likelihood\_aligned\_spin\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std::complex< double > \*detector\_response, size\_t length, fftw\_outline \*plan)

Maximized match over coalescence variables - returns log likelihood NOT NORMALIZED for aligned spins.

double maximized\_Log\_Likelihood\_unaligned\_spin\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std::complex< double > \*hplus, std::complex< double > \*hcross, size\_t length, fftw\_outline \*plan)

log likelihood function that maximizes over extrinsic parameters tc, phic, D, and phiRef, the reference frequency - for unaligned spins

double Log\_Likelihood\_internal (std::complex< double > \*data, double \*psd, double \*frequencies, std
 ::complex< double > \*detector\_response, int length, fftw\_outline \*plan)

Internal function for the unmarginalized log of the likelihood.

void PTMCMC\_MH\_GW (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension, int chain\_id), int numThreads, bool pool, bool show\_prog, int num\_detectors, std::complex< double > \*\*data, double \*\*noise\_psd, double \*\*frequencies, int \*data\_length, double gps\_time, std::string \*detectors, int Nmod, int \*bppe, std::string generation\_method, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)

Wrapper for the MCMC MH function, specifically for GW analysis.

void PTMCMC\_MH\_dynamic\_PT\_alloc\_GW (double \*\*\*output, int dimension, int N\_steps, int chain\_N, int max\_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swpc\_freq, int t0, int nu, std::string chain\_distribution\_scheme, double(\*log\_prior)(double \*param, int dimension, int chain\_id), int numThreads, bool pool, bool show\_prog, int num\_detectors, std::complex < double > \*\*data, double \*\*noise\_psd, double \*\*frequencies, int \*data\_length, double gps\_time, std::string \*detectors, int Nmod, int \*bppe, std::string generation\_method, std::string statistics\_filename, std::string chain\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_filename)

Takes in an MCMC checkpoint file and continues the chain.

void continue\_PTMCMC\_MH\_GW (std::string start\_checkpoint\_file, double \*\*\*output, int dimension, int N←
 \_steps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension, int chain\_id), int numThreads, bool
 pool, bool show\_prog, int num\_detectors, std::complex< double > \*\*data, double \*\*noise\_psd, double
 \*\*frequencies, int \*data\_length, double gps\_time, std::string \*detectors, int Nmod, int \*bppe, std::string
 generation\_method, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename,
 std::string likelihood\_log\_filename, std::string final\_checkpoint\_filename)

Takes in an MCMC checkpoint file and continues the chain.

void PTMCMC\_method\_specific\_prep (std::string generation\_method, int dimension, double \*seeding\_var, bool local seeding)

Unpacks MCMC parameters for method specific initiation.

• void MCMC\_fisher\_wrapper (double \*param, int dimension, double \*\*output, int chain\_id)

Fisher function for MCMC for GW.

double MCMC\_likelihood\_extrinisic (bool save\_waveform, gen\_params \*parameters, std::string generation\_method, int \*data\_length, double \*\*frequencies, std::complex< double > \*\*data, double \*\*psd, std::string \*detectors, fftw\_outline \*fftw\_plans, int num\_detectors, double RA, double DEC, double gps\_time)

double MCMC\_likelihood\_wrapper (double \*param, int dimension, int chain\_id)
 log likelihood function for MCMC for GW

#### 9.27.1 Detailed Description

Routines for implementation in MCMC algorithms specific to GW CBC analysis

#### 9.27.2 Function Documentation

#### 9.27.2.1 continue\_PTMCMC\_MH\_GW()

```
void continue_PTMCMC_MH_GW (
             std::string start_checkpoint_file,
             double *** output,
             int dimension,
             int N_steps,
             int swp_freq,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             int numThreads,
             bool pool,
             bool show_prog,
             int num_detectors,
             std::complex< double > ** data,
             double ** noise_psd,
             double ** frequencies,
             int * data_length,
             double gps_time,
             std::string * detectors,
             int Nmod,
             int * bppe,
             std::string generation_method,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string final_checkpoint_filename )
```

Takes in an MCMC checkpoint file and continues the chain.

Obviously, the user must be sure to correctly match the dimension, number of chains, the generation\_method, the prior function, the data, psds, freqs, and the detectors (number and name), and the gps\_time to the previous run, otherwise the behavior of the sampler is undefined.

numThreads and pool do not necessarily have to be the same

### 9.27.2.2 Log\_Likelihood()

Unmarginalized log of the likelihood.

#### 9.27.2.3 Log\_Likelihood\_internal()

```
double Log_Likelihood_internal (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    std::complex< double > * detector_response,
    int length,
    fftw_outline * plan )
```

Internal function for the unmarginalized log of the likelihood.

```
.5 * ((h|h)-2(D|h))
```

## 9.27.2.4 maximized\_coal\_Log\_Likelihood()

```
double maximized_coal_Log_Likelihood (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    size_t length,
    gen_params * params,
    std::string detector,
    std::string generation_method,
    fftw_outline * plan,
    double * tc,
    double * phic )
```

Function to maximize only over coalescence variables to and phic, returns the maximum values used.

#### 9.27.2.5 maximized\_coal\_log\_likelihood\_IMRPhenomD() [1/3]

Function to calculate the log Likelihood as defined by -1/2 (d-h|d-h) maximized over the extrinsic parameters phic and tc.

frequency array must be uniform spacing - this shouldn't be a problem when working with real data as DFT return uniform spacing

#### **Parameters**

chirpmass in solar masses

# $\textbf{9.27.2.6} \quad maximized\_coal\_log\_likelihood\_IMRPhenomD() \ \ \, \texttt{[2/3]}$

#### **Parameters**

chirpmass in solar masses

## 9.27.2.7 maximized\_coal\_log\_likelihood\_IMRPhenomD() [3/3]

```
size_t length,
double * real_data,
double * imag_data,
double * noise,
double SNR,
double chirpmass,
double symmetric_mass_ratio,
double spin1,
double spin2,
bool NSflag,
fftw_outline * plan )
```

#### **Parameters**

chirpmass in solar masses

## 9.27.2.8 maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param() [1/3]

## **Parameters**

chirpmass in solar masses

### 9.27.2.9 maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param() [2/3]

```
double spin2,
double Luminosity_Distance,
double theta,
double phi,
double iota,
bool NSflag )
```

#### **Parameters**

```
chirpmass in solar masses
```

#### 9.27.2.10 maximized\_coal\_log\_likelihood\_IMRPhenomD\_Full\_Param() [3/3]

```
double maximized_coal_log_likelihood_IMRPhenomD_Full_Param (
             double * frequencies,
             size_t length,
             double * real_data,
             double * imag_data,
             double * noise,
             double chirpmass,
             double symmetric_mass_ratio,
             double spin1,
             double spin2,
             double Luminosity_Distance,
             double theta,
             double phi,
             double iota,
             bool NSflag,
             fftw_outline * plan )
```

#### **Parameters**

```
chirpmass in solar masses
```

## 9.27.2.11 maximized\_Log\_Likelihood()

```
double maximized_Log_Likelihood (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    size_t length,
    gen_params * params,
    std::string detector,
    std::string generation_method,
    fftw_outline * plan )
```

routine to maximize over all extrinsic quantities and return the log likelihood

 $\label{local-phic} \begin{array}{l} \textbf{IMRPhenomD} - \textbf{maximizes over DL}, \textbf{phic, tc, } \textbf{hota}, \textbf{hota$ 

#### 9.27.2.12 maximized\_Log\_Likelihood\_aligned\_spin\_internal()

```
double maximized_Log_Likelihood_aligned_spin_internal (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    std::complex< double > * detector_response,
    size_t length,
    fftw_outline * plan )
```

Maximized match over coalescence variables - returns log likelihood NOT NORMALIZED for aligned spins.

Note: this function is not properly normalized for an absolute comparison. This is made for MCMC sampling, so to minimize time, constant terms like (Data|Data), which would cancel in the Metropolis-Hasting ratio, are left out for efficiency

### 9.27.2.13 maximized\_Log\_Likelihood\_unaligned\_spin\_internal()

```
double maximized_Log_Likelihood_unaligned_spin_internal (
    std::complex< double > * data,
    double * psd,
    double * frequencies,
    std::complex< double > * hplus,
    std::complex< double > * hcross,
    size_t length,
    fftw_outline * plan )
```

log likelihood function that maximizes over extrinsic parameters tc, phic, D, and phiRef, the reference frequency for unaligned spins

Ref: arXiv 1603.02444v2

#### 9.27.2.14 MCMC\_fisher\_wrapper()

Fisher function for MCMC for GW.

Wraps the fisher calculation in src/fisher.cpp and unpacks parameters correctly for common GW analysis

Supports all the method/parameter combinations found in MCMC MH GW

## 9.27.2.15 MCMC\_likelihood\_wrapper()

log likelihood function for MCMC for GW

Wraps the above likelihood functions and unpacks parameters correctly for common GW analysis

Supports all the method/parameter combinations found in MCMC\_MH\_GW

#### 9.27.2.16 PTMCMC\_method\_specific\_prep()

```
void PTMCMC_method_specific_prep (
          std::string generation_method,
          int dimension,
          double * seeding_var,
          bool local_seeding )
```

Unpacks MCMC parameters for method specific initiation.

Populates seeding vector if non supplied, populates mcmc\_Nmod, populates mcmc\_log\_beta, populates mcmc\_← intrinsic

## 9.27.2.17 PTMCMC\_MH\_dynamic\_PT\_alloc\_GW()

```
void PTMCMC_MH_dynamic_PT_alloc_GW (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             int t0,
             int nu.
             std::string chain_distribution_scheme,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             int numThreads,
             bool pool,
             bool show_prog,
             int num_detectors,
             std::complex< double > ** data,
             double ** noise_psd,
             double ** frequencies,
             int * data_length,
             double gps_time,
             std::string * detectors,
             int Nmod,
             int * bppe,
             std::string generation_method,
             std::string statistics_filename,
             std::string chain_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_filename )
```

Takes in an MCMC checkpoint file and continues the chain.

Obviously, the user must be sure to correctly match the dimension, number of chains, the generation\_method, the prior function, the data, psds, freqs, and the detectors (number and name), and the gps\_time to the previous run, otherwise the behavior of the sampler is undefined.

numThreads and pool do not necessarily have to be the same

#### 9.27.2.18 PTMCMC\_MH\_GW()

```
void PTMCMC_MH_GW (
             double *** output.
             int dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             int numThreads,
             bool pool,
             bool show_prog,
             int num_detectors,
             std::complex< double > ** data,
             double ** noise_psd,
             double ** frequencies,
             int * data_length,
             double gps_time,
             std::string * detectors,
             int Nmod,
             int * bppe,
             std::string generation_method,
             std::string statistics_filename,
             std::string chain filename.
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

Wrapper for the MCMC MH function, specifically for GW analysis.

Handles the details of setting up the MCMC sampler and wraps the fisher and log likelihood to conform to the format of the sampler

*NOTE* – This sampler is NOT thread safe. There is global memory declared for each call to MCMC\_MH\_GW, so separate samplers should not be run in the same process space

Supported parameter combinations:

```
IMRPhenomD - 4 dimensions - In chirpmass, eta, chi1, chi2
```

IMRPhenomD - 7 dimensions - In D\_L, tc, phic, In chirpmass, eta, chi1, chi2

IMRPhenomD - 8 dimensions - cos inclination, RA, DEC, In D L, In chirpmass, eta, chi1, chi2

 $\frac{dCS\_IMRPhenomD\_log}{dCS\_IMRPhenomD\_log} - 8 \ dimensions - cos \ inclination, \ RA, \ DEC, \ In \ D\_L, \ In \ chirpmass, \ eta, \ chi1, \ chi2, \ In \ \ lapha^2 \ (the \ coupling \ parameter)$ 

dCS\_IMRPhenomD- 8 dimensions – cos inclination, RA, DEC, In D\_L, In chirpmass, eta, chi1, chi2,  $\alpha^2$  (the coupling parameter)

dCS\_IMRPhenomD\_root\_alpha- 8 dimensions – cos inclination, RA, DEC, In D\_L, In chirpmass, eta, chi1, chi2, \sqrt \alpha (in km) (the coupling parameter)

IMRPhenomPv2 - 9 dimensions - cos J\_N, In chirpmass, eta, |chi1|, |chi1|, theta\_1, theta\_2, phi\_1, phi\_2

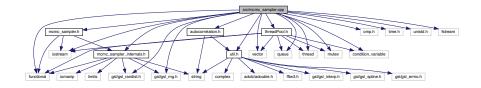
#### **Parameters**

statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.28 src/mcmc\_sampler.cpp File Reference

```
#include "mcmc_sampler.h"
#include "autocorrelation.h"
#include "util.h"
#include "mcmc_sampler_internals.h"
#include "threadPool.h"
#include <iostream>
#include <gsl/gsl_rng.h>
#include <gsl/gsl_randist.h>
#include <omp.h>
#include <time.h>
#include <condition_variable>
#include <mutex>
#include <thread>
#include <vector>
#include <queue>
#include <functional>
#include <unistd.h>
#include <fstream>
```

# Include dependency graph for mcmc\_sampler.cpp:



### **Classes**

- class Comparator
  - Class to facilitate the comparing of chains for priority.
- · class Comparatorswap
- class ThreadPool

# Macros

#define omp ignore

#### **Functions**

void RJPTMCMC\_MH\_internal (double \*\*\*output, int \*\*\*parameter\_status, int max\_dimension, int min← \_dimension, int N\_steps, int chain\_N, double \*initial\_pos, int \*initial\_status, double \*seeding\_var, double \*chain\_temps, int swp\_freq, std::function< double(double \*, int \*, int, int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double \*, int \*, int, double \*\*, int)>fisher, std::function< void(double \*, double \*, int \*, int \*, int, int)> RJ\_proposal, int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)

Generic reversable jump sampler, where the likelihood, prior, and reversable jump proposal are parameters supplied by the user.

void PTMCMC\_MH\_dynamic\_PT\_alloc\_internal (double \*\*\*output, int dimension, int N\_steps, int chain 
 \_N, int max\_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, int t0, int nu, std::string chain\_distribution\_scheme, std::function< double(double \*, int \*, int, int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double \*, int \*, int, double \*\*, int)>fisher, int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)

Dyanmically tunes an MCMC for optimal spacing. step width, and chain number.

void PTMCMC\_MH\_internal (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_
pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, std::function< double(double \*, int \*, int, int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double \*, int \*, int, double \*\*, int)>fisher, int numThreads, bool pool, bool show\_prog, std::string statistics\_
filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std
::string checkpoint file)

Generic sampler, where the likelihood, prior are parameters supplied by the user.

void continue\_PTMCMC\_MH\_internal (std::string start\_checkpoint\_file, double \*\*\*output, int N\_steps, int swp\_freq, std::function< double(double \*, int \*, int, int)> log\_prior, std::function< double(double \*, int \*, int, int)> log\_likelihood, std::function< void(double \*, int \*, int, double \*\*, int)>fisher, int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_cilename, std::string likelihood\_log\_filename, std::string end\_checkpoint\_file)

Routine to take a checkpoint file and begin a new chain at said checkpoint.

void PTMCMC\_MH\_step\_incremental (sampler \*sampler, int increment)

Internal function that runs the actual loop for the sampler – increment version.

void PTMCMC MH loop (sampler \*sampler)

Internal function that runs the actual loop for the sampler.

- void mcmc\_step\_threaded (int j)
- void mcmc\_swap\_threaded (int i, int j)
- void PTMCMC\_MH (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension), double(\*log\_likelihood)(double \*param, int dimension), void(\*fisher)(double \*param, int dimension, double \*fisher), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_
  filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)
- void PTMCMC\_MH (double \*\*\*output, int dimension, int N\_steps, int chain\_N, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_freq, double(\*log\_prior)(double \*param, int dimension, int chain\_id), void(\*fisher)(double \*param, int dimension, double \*fisher, int chain\_id), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_
  filename, std::string checkpoint\_file)
- void continue\_PTMCMC\_MH (std::string start\_checkpoint\_file, double \*\*\*output, int N\_steps, int swp\_← freq, double(\*log\_prior)(double \*param, int dimension, int chain\_id), double(\*log\_likelihood)(double \*param, int dimension, int chain\_id), void(\*fisher)(double \*param, int dimension, double \*\*fisher, int chain\_id), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string auto corr filename, std::string likelihood log filename, std::string end checkpoint file)
- void continue\_PTMCMC\_MH (std::string start\_checkpoint\_file, double \*\*\*output, int N\_steps, int swp\_
  freq, double(\*log\_prior)(double \*param, int dimension), double(\*log\_likelihood)(double \*param, int dimension), void(\*fisher)(double \*param, int dimension, double \*\*fisher), int numThreads, bool pool, bool show
  prog, std::string statistics\_filename, std::string chain\_filename, std::string auto\_corr\_filename, std::string likelihood\_log\_filename, std::string end\_checkpoint\_file)

void PTMCMC\_MH\_dynamic\_PT\_alloc (double \*\*\*output, int dimension, int N\_steps, int chain\_N, int max←
 \_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp\_←
 freq, int t0, int nu, std::string chain\_distribution\_scheme, double(\*log\_prior)(double \*param, int dimension),
 double(\*log\_likelihood)(double \*param, int dimension), void(\*fisher)(double \*param, int dimension, double
 \*\*fisher), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_←
 filename, std::string likelihood log filename, std::string checkpoint file)

- void PTMCMC\_MH\_dynamic\_PT\_alloc (double \*\*\*output, int dimension, int N\_steps, int chain\_N, int max
   \_chain\_N\_thermo\_ensemble, double \*initial\_pos, double \*seeding\_var, double \*chain\_temps, int swp
   \_freq, int t0, int nu, std::string chain\_distribution\_scheme, double(\*log\_prior)(double \*param, int dimension, int chain\_id), double(\*log\_likelihood)(double \*param, int dimension, int chain\_id), void(\*fisher)(double \*param, int dimension, double \*\*fisher, int chain\_id), int numThreads, bool pool, bool show\_prog, std::string statistics\_filename, std::string chain\_filename, std::string likelihood\_log\_filename, std::string checkpoint\_file)

#### **Variables**

- const gsl\_rng\_type \* T
- gsl rng \* r
- sampler \* samplerptr
- ThreadPool \* poolptr

#### 9.28.1 Detailed Description

Source file for the sampler foundation

Source file for generic MCMC sampler. Sub routines that are application agnostic are housed in mcmc\_sampler 
\_internals

## 9.28.2 Function Documentation

### 9.28.2.1 continue\_PTMCMC\_MH() [1/2]

```
bool show_prog,
std::string statistics_filename,
std::string chain_filename,
std::string auto_corr_filename,
std::string likelihood_log_filename,
std::string end_checkpoint_file )
```

#### **Parameters**

	start_checkpoint_file	File for starting checkpoint
out	output	output array, dimensions: output[chain_N][N_steps][dimension]
	N_steps	Number of new steps to take
	swp_freq	frequency of swap attempts between temperatures
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use
	pool	Boolean for whether to use deterministic'' vsstochastic" sampling
	show_prog	Boolean for whether to show progress or not (turn off for cluster runs
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	end_checkpoint_file	Filename to output data for checkpoint at the end of the continued run, if empty string, not saved

#### **9.28.2.2** continue\_PTMCMC\_MH() [2/2]

```
void continue_PTMCMC_MH (
            std::string start_checkpoint_file,
             double *** output,
             int N\_steps,
             int swp_freq,
             double(*)(double *param, int dimension) log_prior,
             double(*)(double *param, int dimension) log_likelihood,
             void(*)(double *param, int dimension, double **fisher) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string end_checkpoint_file )
```

	start_checkpoint_file	File for starting checkpoint
out	output	output array, dimensions: output[chain_N][N_steps][dimension]
	N_steps	Number of new steps to take
	swp_freq	frequency of swap attempts between temperatures
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use

#### **Parameters**

pool	Boolean for whether to use deterministic'' vsstochastic" sampling
show_prog	Boolean for whether to show progress or not (turn off for cluster runs
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
end_checkpoint_file	Filename to output data for checkpoint at the end of the continued run, if empty string, not saved

## 9.28.2.3 continue\_PTMCMC\_MH\_internal()

```
void continue_PTMCMC_MH_internal (
            std::string start_checkpoint_file,
             double *** output,
             int N_steps,
             int swp_freq,
             std::function< double(double *, int *, int, int) > log_prior,
             std::function< double(double *, int *, int, int) > log_likelihood,
             std::function< void(double *, int *, int, double **, int) > fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string end_checkpoint_file )
```

Routine to take a checkpoint file and begin a new chain at said checkpoint.

See MCMC\_MH\_internal for more details of parameters (pretty much all the same)

	start_checkpoint_file	File for starting checkpoint
out	output	output array, dimensions: output[chain_N][N_steps][dimension]
	N_steps	Number of new steps to take
	swp_freq	frequency of swap attempts between temperatures
	log_prior	std::function for the log_prior function – takes double *position, int dimension, int chain_id
	log_likelihood	std::function for the log_likelihood function – takes double *position, int dimension, int chain_id
	fisher	std::function for the fisher function – takes double *position, int dimension, double **output_fisher, int chain_id
	numThreads	Number of threads to use
	pool	Boolean for whether to use deterministic'' vsstochastic" sampling
	show_prog	Boolean for whether to show progress or not (turn off for cluster runs

#### **Parameters**

statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
end_checkpoint_file	Filename to output data for checkpoint at the end of the continued run, if empty string, not saved

# **9.28.2.4 PTMCMC\_MH()** [1/2]

```
void PTMCMC_MH (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             double(*)(double *param, int dimension) log_prior,
             double(*)(double *param, int dimension) log_likelihood,
             void(*)(double *param, int dimension, double **fisher) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have >2 threads)

#### **Parameters**

show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# **9.28.2.5 PTMCMC\_MH()** [2/2]

```
void PTMCMC_MH (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             \verb|double(*)| (\verb|double *param*, int dimension*, int chain_id) | log\_prior*,
             double(*)(double *param, int dimension, int chain_id) log_likelihood,
             void(*)(double *param, int dimension, double **fisher, int chain_id) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

urumeters		
out	output	Output chains, shape is double[chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	Funcion pointer for the log_prior
	log_likelihood	Function pointer for the log_likelihood
	fisher	Function pointer for the fisher - if NULL, fisher steps are not used
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have >2 threads)
		I .

### **Parameters**

show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.28.2.6 PTMCMC\_MH\_dynamic\_PT\_alloc() [1/2]

```
void PTMCMC_MH_dynamic_PT_alloc (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp\_freq,
             int t0,
             int nu,
             std::string chain_distribution_scheme,
             double(*)(double *param, int dimension) log_prior,
             double(*)(double *param, int dimension) log_likelihood,
             void(*)(double *param, int dimension, double **fisher) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

out	output	Output chains, shape is double[max_chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain AFTER chain allocation
	chain_N	Maximum number of chains to use
	max_chain_N_thermo_ensemble	Maximum number of chains to use in the thermodynamic ensemble (may use less)
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]

#### **Parameters**

chain_temps	Final chain temperatures used – should be shape double[chain_N]
swp_freq	the frequency with which chains are swapped
tO	Time constant of the decay of the chain dynamics (~1000)
nu	Initial amplitude of the dynamics (~100)
log_prior	Funcion pointer for the log_prior
log_likelihood	Function pointer for the log_likelihood
fisher	Function pointer for the fisher - if NULL, fisher steps are not used
numThreads	Number of threads to use (=1 is single threaded)
pool	boolean to use stochastic chain swapping (MUST have $>$ 2 threads)
show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.28.2.7 PTMCMC\_MH\_dynamic\_PT\_alloc() [2/2]

```
void PTMCMC_MH_dynamic_PT_alloc (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             int t0,
             int nu,
             std::string chain_distribution_scheme,
             double(*)(double *param, int dimension, int chain_id) log_prior,
             double(*)(double *param, int dimension, int chain_id) log_likelihood,
             void(*)(double *param, int dimension, double **fisher, int chain_id) fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

out	output	Output chains, shape is double[max_chain_N,
		N_steps,dimension]

### **Parameters**

dimension	dimension of the parameter space being explored
N_steps	Number of total steps to be taken, per chain AFTER chain allocation
chain_N	Maximum number of chains to use
max_chain_N_thermo_ensemble	Maximum number of chains to use in the thermodynamic ensemble (may use less)
initial_pos	Initial position in parameter space - shape double[dimension]
seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
chain_temps	Final chain temperatures used – should be shape double[chain_N]
swp_freq	the frequency with which chains are swapped
t0	Time constant of the decay of the chain dynamics ( $\sim$ 1000)
nu	Initial amplitude of the dynamics (~100)
log_prior	Funcion pointer for the log_prior
log_likelihood	Function pointer for the log_likelihood
fisher	Function pointer for the fisher - if NULL, fisher steps are not used
numThreads	Number of threads to use (=1 is single threaded)
pool	boolean to use stochastic chain swapping (MUST have $>$ 2 threads)
show_prog	boolean whether to print out progress (for example, should be se to `'false'' if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.28.2.8 PTMCMC\_MH\_dynamic\_PT\_alloc\_internal()

```
void PTMCMC_MH_dynamic_PT_alloc_internal (
             double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             int max_chain_N_thermo_ensemble,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             int t0,
             int nu,
             std::string chain_distribution_scheme,
             std::function< double(double *, int *, int, int) > log_prior,
             std::function< double(double *, int *, int, int) > log_likelihood,
             std::function< void(double *, int *, int, double **, int)> fisher,
             int numThreads,
             bool pool,
```

```
bool show_prog,
std::string statistics_filename,
std::string chain_filename,
std::string likelihood_log_filename,
std::string checkpoint_file )
```

Dyanmically tunes an MCMC for optimal spacing. step width, and chain number.

NOTE: nu, and t0 parameters determine the dynamics, so these are important quantities. nu is related to how many swap attempts it takes to substantially change the temperature ladder, why t0 determines the length of the total dyanimcally period. Moderate initial choices would be 10 and 1000, respectively.

Based on arXiv:1501.05823v3

Currently, Chain number is fixed

max\_chain\_N\_thermo\_ensemble sets the maximium number of chains to use to in successively hotter chains to cover the likelihood surface while targeting an optimal swap acceptance target\_swp\_acc.

max\_chain\_N determines the total number of chains to run once thermodynamic equilibrium has been reached. This results in chains being added after the initial PT dynamics have finished according to chain\_distribution\_\circ
scheme.

If no preference, set max\_chain\_N\_thermo\_ensemble = max\_chain\_N = numThreads = (number of cores (number of threads if hyperthreaded))— this will most likely be the most optimal configuration. If the number of cores on the system is low, you may want to use n\*numThreads for some integer n instead, depending on the system.

chain\_distribution\_scheme:

"cold": All chains are added at T=1 (untempered)

"refine": Chains are added between the optimal temps geometrically – this may be a good option as it will be a good approximation of the ideal distribution of chains, while keeping the initial dynamical time low

"double": Chains are added in order of rising temperature that mimic the distribution achieved by the earier PT dynamics

"half\_ensemble": For every cold chain added, half of the ensemble is added again. Effectively, two cold chains for every ensemble

out	output	Output chains, shape is double[max_chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain AFTER chain allocation
	chain_N	Maximum number of chains to use
	max_chain_N_thermo_ensemble	Maximum number of chains to use in the thermodynamic ensemble (may use less)
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
out	chain_temps	Final chain temperatures used – should be shape double[chain_N]
	swp_freq	the frequency with which chains are swapped
	t0	Time constant of the decay of the chain dynamics ( $\sim$ 1000)

#### **Parameters**

nu	Initial amplitude of the dynamics ( $\sim$ 100)
log_prior	std::function for the log_prior function – takes double *position, int dimension, int chain_id
log_likelihood	std::function for the log_likelihood function – takes double *position, int dimension, int chain_id
fisher	std::function for the fisher function – takes double *position, int dimension, double **output_fisher, int chain_id
numThreads	Number of threads to use (=1 is single threaded)
pool	boolean to use stochastic chain swapping (MUST have >2 threads)
show_prog	boolean whether to print out progress (for example, should be set to `'false" if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

#### 9.28.2.9 PTMCMC\_MH\_internal()

```
void PTMCMC_MH_internal (
            double *** output,
             int dimension,
             int N_steps,
             int chain_N,
             double * initial_pos,
             double * seeding_var,
             double * chain_temps,
             int swp_freq,
             std::function< double(double *, int *, int, int) > log_prior,
             std::function< double(double *, int *, int, int)> log_likelihood,
             std::function< void(double *, int *, int, double **, int)> fisher,
             int numThreads,
             bool pool,
             bool show_prog,
             std::string statistics_filename,
             std::string chain_filename,
             std::string auto_corr_filename,
             std::string likelihood_log_filename,
             std::string checkpoint_file )
```

Generic sampler, where the likelihood, prior are parameters supplied by the user.

Base of the sampler, generic, with user supplied quantities for most of the samplers properties

Uses the Metropolis-Hastings method, with the option for Fisher/MALA steps if the Fisher routine is supplied.

3 modes to use -

single threaded (numThreads = 1) runs single threaded

multi-threaded `'deterministic'' (numThreads>1; pool = false) progresses each chain in parallel for swp\_freq steps, then waits for all threads to complete before swapping temperatures in sequenctial order (j, j+1) then (j+1, j+2) etc (sequenctially)

multi-threaded `'stochastic" (numThreads>2; pool = true) progresses each chain in parallel by queueing each temperature and evaluating them in the order they were submitted. Once finished, the threads are queued to swap, where they swapped in the order they are submitted. This means the chains are swapped randomly, and the chains do NOT finish at the same time. The sampler runs until the the 0th chain reaches the step number

Note on limits: In the prior function, if a set of parameters should be disallowed, return -std::numeric\_ limits < double > ::infinity() - (this is in the < limits > file in std)

Format for the auto\_corr file (compatable with csv, dat, txt extensions): each row is a dimension of the cold chain, with the first row being the lengths used for the auto-corr calculation:

lengths: length1, length2...

dim1: length1, length2...

Format for the chain file (compatable with csv, dat, txt extensions): each row is a step, each column a dimension:

Step1: dim1, dim2, ...

Step2: dim1, dim2, ...

Statistics filename: should be txt extension

checkpoint\_file: This file saves the final position of all the chains, as well as other metadata, and can be loaded by the function <FUNCTION> to continue the chain from the point it left off. Not meant to be read by humans, the data order is custom to this software library. An empty string ("") means no checkpoint will be saved. For developers, the contents are:

dimension, # of chains

temps of chains

Stepping widths of all chains

Final position of all chains

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
	dimension	dimension of the parameter space being explored
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[dimension]
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	std::function for the log_prior function – takes double *position, int dimension, int chain_id
	log_likelihood	std::function for the log_likelihood function – takes double *position, int dimension, int chain_id
	fisher	std::function for the fisher function – takes double *position, int dimension, double **output_fisher, int chain_id

#### **Parameters**

numThreads	Number of threads to use (=1 is single threaded)
pool	boolean to use stochastic chain swapping (MUST have >2 threads)
show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
statistics_filename	Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

### 9.28.2.10 PTMCMC\_MH\_loop()

Internal function that runs the actual loop for the sampler.

# 9.28.2.11 PTMCMC\_MH\_step\_incremental()

Internal function that runs the actual loop for the sampler – increment version.

The regular loop function runs for the entire range, this increment version will only step `increment' steps – asynchronous: steps are measured by the cold chains NEEDS TO CHANGE

# 9.28.2.12 RJPTMCMC\_MH()

#### **Parameters**

out	output	Output chains, shape is double[chain_N, N_steps,dimension]
out	parameter_status	Parameter status for each step corresponding to the output chains, shape is double[chain_N, N_steps,dimension]
	max_dimension	maximum dimension of the parameter space being explored – only consideration is memory, as memory scales with dimension. Keep this reasonable, unless memory is REALLY not an issue
	min_dimension	minimum dimension of the parameter space being explored >=1
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	initial_status	Initial status of the parameters in the initial position in parameter space - shape int[max_dim]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[max_dimension] – initial seeding of zero corresponds to the dimension turned off initially
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	RJ_proposal	std::function for the log_likelihood function – takes double *position, int *param_status,int dimension, int chain_id
	numThreads	Number of threads to use (=1 is single threaded)
	pool	boolean to use stochastic chain swapping (MUST have >2 threads)
	show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
	statistics_filename	Filename to output sampling statistics, if empty string, not output
	chain_filename	Filename to output data (chain 0 only), if empty string, not output
	auto_corr_filename	Filename to output auto correlation in some interval, if empty string, not output
	likelihood_log_filename	Filename to write the log_likelihood and log_prior at each step – use empty string to skip
	checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.28.2.13 RJPTMCMC\_MH\_internal()

```
int *** parameter_status,
int max_dimension,
int min_dimension,
int N_steps,
int chain_N,
double * initial_pos,
int * initial status,
double * seeding_var,
double * chain_temps,
int swp_freq,
std::function< double(double *, int *, int, int) > log_prior,
std::function< double(double *, int *, int, int) > log_likelihood,
std::function< void(double *, int *, int, double **, int)> fisher,
std::function< void(double *, double *, int *, int *, int, int) > RJ_proposal,
int numThreads,
bool pool,
bool show_prog,
std::string statistics_filename,
std::string chain_filename,
std::string auto_corr_filename,
std::string likelihood_log_filename,
std::string checkpoint_file )
```

Generic reversable jump sampler, where the likelihood, prior, and reversable jump proposal are parameters supplied by the user.

Note: Using a min\_dimension tells the sampler that there is a base model'', and that the dimensions from min\_dim to max\_dim are small" corrections to that model. This helps inform some of the proposal algorithms and speeds up computation. If using discrete models with no overlap of variables (ie model A or model B), set min\_dim to 0. Even if reusing certain parameters, if the extra dimensions don't describe 'small' deviations, it's probably best to set min dim to 0.

Currently, no dynamic PT option, as it would be too many free parameters for the sampler to converge to a reasonable temperature distribution in a reasonable amount of time. Best use case, use the PTMCMC\_MH\_dyanmic\_PT for the `'base'' dimension space, and use that temperature ladder.

Base of the sampler, generic, with user supplied quantities for most of the samplers properties

Uses the Metropolis-Hastings method, with the option for Fisher/MALA steps if the Fisher routine is supplied.

3 modes to use -

single threaded (numThreads = 1) runs single threaded

multi-threaded `'deterministic'' (numThreads>1; pool = false) progresses each chain in parallel for swp\_freq steps, then waits for all threads to complete before swapping temperatures in sequenctial order (j, j+1) then (j+1, j+2) etc (sequenctially)

multi-threaded `'stochastic" (numThreads>2; pool = true) progresses each chain in parallel by queueing each temperature and evaluating them in the order they were submitted. Once finished, the threads are queued to swap, where they swapped in the order they are submitted. This means the chains are swapped randomly, and the chains do NOT finish at the same time. The sampler runs until the the 0th chain reaches the step number

Note on limits: In the prior function, if a set of parameters should be disallowed, return -std::numeric $\_\leftarrow$  limits<double>::infinity() - (this is in the limits> file in std)

The parameter array uses the dimensions [0,min\_dim] always, and [min\_dim, max\_dim] in RJPTMCMC fashion

Format for the auto\_corr file (compatable with csv, dat, txt extensions): each row is a dimension of the cold chain, with the first row being the lengths used for the auto-corr calculation:

lengths: length1, length2...

dim1: length1, length2...

Format for the chain file (compatable with csv, dat, txt extensions): each row is a step, each column a dimension:

Step1: dim1, dim2, ..., max\_dim, param\_status1, param\_status2, ...

 $Step 2: dim1 \ , dim2 \ , ..., max\_dim, param\_status 1, param\_status 2, ...$ 

Statistics\_filename : should be txt extension

checkpoint\_file: This file saves the final position of all the chains, as well as other metadata, and can be loaded by the function <FUNCTION> to continue the chain from the point it left off. Not meant to be read by humans, the data order is custom to this software library. An empty string ("") means no checkpoint will be saved. For developers, the contents are:

dimension, # of chains

temps of chains

Stepping widths of all chains

Final position of all chains

out	output	Output chains, shape is double[chain N, N steps,dimension]
out	parameter_status	Parameter status for each step corresponding to the output chains, shape is double[chain_N, N_steps,dimension]
	max_dimension	maximum dimension of the parameter space being explored – only consideration is memory, as memory scales with dimension. Keep this reasonable, unless memory is REALLY not an issue
	min_dimension	minimum dimension of the parameter space being explored >=1
	N_steps	Number of total steps to be taken, per chain
	chain_N	Number of chains
	initial_pos	Initial position in parameter space - shape double[dimension]
	initial_status	Initial status of the parameters in the initial position in parameter space - shape int[max_dim]
	seeding_var	Variance of the normal distribution used to seed each chain higher than 0 - shape double[max_dimension] – initial seeding of zero corresponds to the dimension turned off initially
	chain_temps	Double array of temperatures for the chains
	swp_freq	the frequency with which chains are swapped
	log_prior	std::function for the log_prior function – takes double *position, int *param_status, int dimension, int chain_id
	log_likelihood	std::function for the log_likelihood function – takes double *position, int *param_status,int dimension, int chain_id
	fisher	std::function for the fisher function – takes double *position, int *param_status,int dimension, double **output_fisher, int chain_id
	RJ_proposal	std::function for the log_likelihood function – takes double *position, int *param_status,int dimension, int chain_id
	numThreads	Number of threads to use (=1 is single threaded)

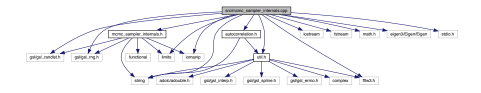
#### **Parameters**

pool	boolean to use stochastic chain swapping (MUST have >2 threads)
show_prog	boolean whether to print out progress (for example, should be set to `'false'' if submitting to a cluster)
statistics_filenan	ne Filename to output sampling statistics, if empty string, not output
chain_filename	Filename to output data (chain 0 only), if empty string, not output
auto_corr_filena	me Filename to output auto correlation in some interval, if empty string, not output
likelihood_log_fii	lename Filename to write the log_likelihood and log_prior at each step – use empty string to skip
checkpoint_file	Filename to output data for checkpoint, if empty string, not saved

# 9.29 src/mcmc\_sampler\_internals.cpp File Reference

```
#include "mcmc_sampler_internals.h"
#include "autocorrelation.h"
#include "util.h"
#include <iostream>
#include <fstream>
#include <string>
#include <math.h>
#include <gsl/gsl_randist.h>
#include <gsl/gsl_rng.h>
#include eigen3/Eigen/Eigen>
#include <iomanip>
#include <fftw3.h>
#include <stdio.h>
```

Include dependency graph for mcmc\_sampler\_internals.cpp:



## **Functions**

• int mcmc\_step (sampler \*sampler, double \*current\_param, double \*next\_param, int \*current\_status, int \*next\_status, int chain\_number)

interface function between the sampler and the internal step functions

void gaussian\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_

 status, int \*proposed\_status, int chain\_id)

Straight gaussian step.

• void fisher\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed\_status, int chain\_index)

Fisher informed gaussian step.

- void update\_fisher (sampler \*sampler, double \*current\_param, int \*param\_status, int chain\_index)
- void mmala\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed\_status)

MMALA informed step - Currently not supported.

void diff\_ev\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed\_status, int chain\_id)

differential evolution informed step

- void RJ\_smooth\_history (sampler \*sampler, int \*current\_param\_status, int base\_history\_id, double \*eff\_←
  history coord, int \*eff history status, int chain id)
- void RJ\_step (sampler \*sampler, double \*current\_param, double \*proposed\_param, int \*current\_status, int \*proposed status, int chain number)

RJ-proposal step for trans-dimensional MCMCs.

void chain\_swap (sampler \*sampler, double \*\*\*output, int \*\*\*param\_status, int step\_num, int \*swp\_

 accepted, int \*swp\_rejected)

subroutine to perform chain comparison for parallel tempering

int single\_chain\_swap (sampler \*sampler, double \*chain1, double \*chain2, int \*chain1\_status, int \*chain1\_status, int \*chain1\_status, int T1\_index, int T2\_index)

subroutine to actually swap two chains

void assign probabilities (sampler \*sampler, int chain index)

update and initiate probabilities for each variety of step

void transfer\_chain (sampler \*samplerptr\_dest, sampler \*samplerptr\_source, int id\_dest, int id\_source, bool transfer\_output)

Copies contents of one chain to another.

• bool check\_sampler\_status (sampler \*samplerptr)

Checks the status of a sampler for the stochastic sampling.

void update\_step\_widths (sampler \*samplerptr, int chain\_id)

Updates the step widths, shooting for 20% acceptance ratios for each type of step.

- void allocate sampler mem (sampler \*sampler)
- void deallocate\_sampler\_mem (sampler \*sampler)
- void update\_history (sampler \*sampler, double \*new\_params, int \*new\_param\_status, int chain\_index)
- void write\_stat\_file (sampler \*sampler, std::string filename)
- void write\_checkpoint\_file (sampler \*sampler, std::string filename)

Routine that writes metadata and final positions of a sampler to a checkpoint file.

• void load\_temps\_checkpoint\_file (std::string check\_file, double \*temps, int chain\_N)

load temperatures from checkpoint file

void load\_checkpoint\_file (std::string check\_file, sampler \*sampler)

load checkpoint file into sampler struct

- void assign\_ct\_p (sampler \*sampler, int step, int chain\_index)
- void assign ct m (sampler \*sampler, int step, int chain index)
- void assign\_initial\_pos (sampler \*samplerptr, double \*initial\_pos, int \*initial\_status, double \*seeding\_var)
- double PT dynamical timescale (int t0, int nu, int t)

Timescale of the PT dynamics.

• void update temperatures (sampler \*samplerptr, int t0, int nu, int t)

updates the temperatures for a sampler such that all acceptance rates are equal

• void initiate\_full\_sampler (sampler \*sampler\_new, sampler \*sampler\_old, int chain\_N\_thermo\_ensemble, int chain\_N, std::string chain\_allocation\_scheme)

For the dynamic PT sampler, this is the function that starts the full sampler with the max number of chains.

## 9.29.1 Detailed Description

File containing definitions for all the internal, generic mcmc subroutines

# 9.29.2 Function Documentation

### 9.29.2.1 assign\_probabilities()

update and initiate probabilities for each variety of step

Type 0: Gaussian step

Type 1: Differential Evolution step

Type 2: MMALA step (currently not supported)

Type 3: Fisher step

### 9.29.2.2 chain\_swap()

subroutine to perform chain comparison for parallel tempering

The total output file is passed, and the chains are swapped sequentially

This is the routine for `'Deterministic" sampling (parallel or sequential, but not pooled)

### **Parameters**

sampler	sampler struct
output	output vector containing chains
param_status	Parameter status
step_num	current step number

# 9.29.2.3 check\_sampler\_status()

```
bool check_sampler_status (
          sampler * samplerptr )
```

Checks the status of a sampler for the stochastic sampling.

Just loops through the ref\_chain\_status variables

### 9.29.2.4 diff\_ev\_step()

# differential evolution informed step

Differential evolution uses the past history of the chain to inform the proposed step:

Take the difference of two random, accepted previous steps and step along that with some step size, determined by a gaussian

# **Parameters**

	sampler	Sampler struct
	current_param	current position in parameter space
out	proposed_param	Proposed position in parameter space

# 9.29.2.5 fisher\_step()

Fisher informed gaussian step.

#### **Parameters**

	sampler	Sampler struct
	current_param	current position in parameter space
out	proposed_param	Proposed position in parameter space

### 9.29.2.6 gaussian\_step()

```
int * current_status,
int * proposed_status,
int chain_id )
```

Straight gaussian step.

### **Parameters**

	sampler	Sampler struct
	current_param	current position in parameter space
out	proposed_param	Proposed position in parameter space

# 9.29.2.7 initiate\_full\_sampler()

```
void initiate_full_sampler (
          sampler * sampler_new,
          sampler * sampler_old,
          int chain_N_thermo_ensemble,
          int chain_N,
          std::string chain_allocation_scheme )
```

For the dynamic PT sampler, this is the function that starts the full sampler with the max number of chains.

The output file will be reused, but the positions are set back to zero (copying the current position to position zero)

Assumes the output, chain\_temps have been allocated in memory for the final number of chains chain\_N and steps N steps

Allocates memory for the new sampler sampler\_new -> it's the user's responsibility to deallocate with deallocate -- \_sampler\_mem

#### **Parameters**

sampler_old	Dynamic sampler
chain_N_thermo_ensemble	Number of chains used in the thermodynamic ensemble
chain_N	Number of chains to use in the static sampler
chain_allocation_scheme	Scheme to use to allocate any remaining chains

# 9.29.2.8 load\_checkpoint\_file()

load checkpoint file into sampler struct

NOTE - allocate\_sampler called in function - MUST deallocate manually

NOTE - sampler->chain\_temps allocated internally - MUST free manually

### 9.29.2.9 load\_temps\_checkpoint\_file()

load temperatures from checkpoint file

Assumed the temps array is already allocated in memory for the correct number of chains

Just a utility routine to read temperatures from checkpoint file

It would be easy to read in the chain number and allocate memory in the function, but I prefer to leave allocation/deallocation up to the client

### 9.29.2.10 mmala\_step()

MMALA informed step - Currently not supported.

### **Parameters**

		sampler	Sampler struct
ſ		current_param	current position in parameter space
	out	proposed_param	Proposed position in parameter space

### 9.29.2.11 PT\_dynamical\_timescale()

Timescale of the PT dynamics.

kappa in the the language of arXiv:1501.05823v3

t0	Timescale of the dyanmics	
nu	Initial amplitude (number of steps to base dynamics on)	
t	current time	

### 9.29.2.12 RJ\_smooth\_history()

### **Parameters**

	sampler	Current sampler
	current_param_status	Current parameters to match
base_history_id		Original history element
out	eff_history_coord	Modified history coord
out	eff_history_status	Modified History status
	chain_id	Chain ID of the current chain

# 9.29.2.13 RJ\_step()

RJ-proposal step for trans-dimensional MCMCs.

This extra step may seem unnecessary, I'm just adding it in in case the extra flexibility is useful in the future for preprocessing of the chain before sending it to the user's RJ\_proposal

	sampler	sampler
	current_param	current coordinates in parameter space
out	proposed_param	Proposed coordinates in parameter space
	current_status	Current status of parameters
out	proposed_status	Proposed status of parameters
	chain_number	chain mumber

### 9.29.2.14 single\_chain\_swap()

subroutine to actually swap two chains

This is the more general subroutine, which just swaps the two chains passed to the function

#### **Parameters**

sampler	npler sampler structure	
chain1 parameter position of chain that could be changed		
chain2	chain that is not swapped, but provides parameters to be swapped by the other chain	
chain1_status	Parameter status array for chain1	
chain2_status Parameter status array for chain2		
T1_index	number of chain swappe in chain_temps	
T2_index number of chain swapper in chain_temps		

### 9.29.2.15 transfer\_chain()

Copies contents of one chain to another.

Transfers id\_source in samplerptr\_source to id\_dest samplerptr\_dest

NOTE: This copies the VALUE, not the reference. This could be expensive, so use with caution

id\_dest is ERASED

samplerptr\_dest and samplerptr\_source MUST have the same dimension, the same sampling details (like having or not having a fisher) etc

samplerptr\_dest must be previously allocated properly

As output is the largest transfer by far, the transfer\_output flag can be used to allow the user to handle that manually.

### 9.29.2.16 update\_temperatures()

updates the temperatures for a sampler such that all acceptance rates are equal

Follows the algorithm outlined in arXiv:1501.05823v3

Fixed temperatures for the first and last chain

```
used in MCMC_MH_dynamic_PT_alloc_internal
```

For defined results, this should be used while the sampler is using non-pooling methods

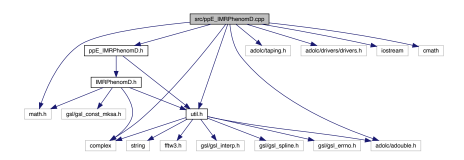
# 9.29.2.17 write\_checkpoint\_file()

Routine that writes metadata and final positions of a sampler to a checkpoint file.

# 9.30 src/ppE\_IMRPhenomD.cpp File Reference

```
#include "ppE_IMRPhenomD.h"
#include <math.h>
#include <adolc/adouble.h>
#include <adolc/taping.h>
#include <adolc/drivers/drivers.h>
#include <iostream>
#include <cmath>
#include <complex>
#include "util.h"
```

Include dependency graph for ppE\_IMRPhenomD.cpp:



# 9.30.1 Detailed Description

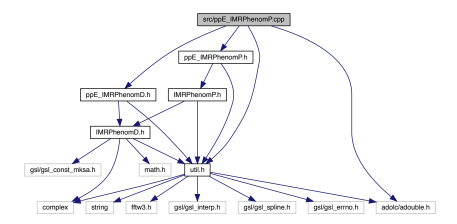
File for the implementation of the ppE formalism for testing GR

Extends the IMRPhenomD template to include non-GR phase terms

Supported waveforms: ppE Inspiral, ppE IMR, dCS, EdGB

# 9.31 src/ppE\_IMRPhenomP.cpp File Reference

```
#include "ppE_IMRPhenomP.h"
#include "ppE_IMRPhenomD.h"
#include "util.h"
#include <adolc/adouble.h>
Include dependency graph for ppE_IMRPhenomP.cpp:
```



# Macros

- #define ROTATEZ(angle, vx, vy, vz)
- #define **ROTATEY**(angle, vx, vy, vz)

# **Variables**

• const double **sqrt 6** = 2.44948974278317788

# 9.31.1 Detailed Description

Source code file for parameterized post Einsteinian Modifications to the precessing waveform model IMRPhenomP

# 9.31.2 Macro Definition Documentation

### 9.31.2.1 ROTATEY

```
#define ROTATEY(

angle,

vx,

vy,

vz)
```

#### Value:

```
tmp1 = vx*cos(angle) + vz*sin(angle);\
tmp2 = - vx*sin(angle) + vz*cos(angle);\
vx = tmp1;\
vz = tmp2
```

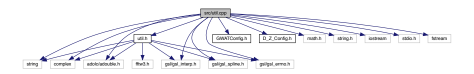
#### 9.31.2.2 ROTATEZ

#### Value:

```
tmp1 = vx*cos(angle) - vy*sin(angle);\
tmp2 = vx*sin(angle) + vy*cos(angle);\
vx = tmp1;\
vy = tmp2
```

# 9.32 src/util.cpp File Reference

```
#include "util.h"
#include "GWATConfig.h"
#include "D_Z_Config.h"
#include <math.h>
#include <string>
#include <string.h>
#include <complex>
#include <iostream>
#include <fstream>
#include <adolc/adouble.h>
#include <gsl/gsl_interp.h>
#include <gsl/gsl_spline.h>
#include <gsl/gsl_errno.h>
Include dependency graph for util.cpp:
```



#### **Functions**

void initiate\_LumD\_Z\_interp (gsl\_interp\_accel \*\*Z\_DL\_accel\_ptr, gsl\_spline \*\*Z\_DL\_spline\_ptr)

Function that uses the GSL libraries to interpolate pre-calculated Z-D\_L data.

void free\_LumD\_Z\_interp (gsl\_interp\_accel \*\*Z\_DL\_accel\_ptr, gsl\_spline \*\*Z\_DL\_spline\_ptr)

Frees the allocated interpolation function.

- adouble Z\_from\_DL\_interp (adouble DL, gsl\_interp\_accel \*Z\_DL\_accel\_ptr, gsl\_spline \*Z\_DL\_spline\_ptr)
- double Z\_from\_DL\_interp (double DL, gsl\_interp\_accel \*Z\_DL\_accel\_ptr, gsl\_spline \*Z\_DL\_spline\_ptr)
- double Z\_from\_DL (double DL, std::string cosmology)

Calculates the redshift given the luminosity distance.

adouble Z\_from\_DL (adouble DL, std::string cosmology)

Calculates the redshift given the luminosity distance adouble version for ADOL-C implementation.

double DL\_from\_Z (double Z, std::string cosmology)

Calculates the luminosity distance given the redshift.

adouble DL\_from\_Z (adouble Z, std::string cosmology)

Calculates the luminosity distance given the redshift adouble version for ADOL-C implementation.

double cosmology\_interpolation\_function (double x, double \*coeffs, int interp\_degree)

Custom interpolation function used in the cosmology calculations.

• adouble cosmology\_interpolation\_function (adouble x, double \*coeffs, int interp\_degree)

Custom interpolation function used in the cosmology calculations adouble version for ADOL-C.

double cosmology\_lookup (std::string cosmology)

Helper function for mapping cosmology name to an internal index.

void printProgress (double percentage)

routine to print the progress of a process to the terminal as a progress bar

void allocate\_FFTW\_mem\_forward (fftw\_outline \*plan, int length)

Allocate memory for FFTW3 methods used in a lot of inner products input is a locally defined structure that houses all the pertinent data.

void allocate\_FFTW\_mem\_reverse (fftw\_outline \*plan, int length)

Allocate memory for FFTW3 methods used in a lot of inner products –INVERSE input is a locally defined structure that houses all the pertinent data.

void deallocate\_FFTW\_mem (fftw\_outline \*plan)

deallocates the memory used for FFTW routines

double calculate chirpmass (double mass1, double mass2)

Calculates the chirp mass from the two component masses.

- adouble calculate\_chirpmass (adouble mass1, adouble mass2)
- double calculate\_eta (double mass1, double mass2)

Calculates the symmetric mass ration from the two component masses.

- adouble calculate\_eta (adouble mass1, adouble mass2)
- double calculate\_mass1 (double chirpmass, double eta)

Calculates the larger mass given a chirp mass and symmetric mass ratio.

- adouble calculate\_mass1 (adouble chirpmass, adouble eta)
- double calculate mass2 (double chirpmass, double eta)

Calculates the smaller mass given a chirp mass and symmetric mass ratio.

- adouble calculate mass2 (adouble chirpmass, adouble eta)
- long factorial (long num)

Local function to calculate a factorial.

double pow\_int (double base, int power)

Local power function, specifically for integer powers.

- adouble **pow\_int** (adouble base, int power)
- · double cbrt internal (double base)

Fucntion that just returns the cuberoot.

adouble cbrt\_internal (adouble base)

Fucntion that just returns the cuberoot ADOL-C doesn't have the cbrt function (which is faster), so have to use the power function.

double \*\* allocate\_2D\_array (int dim1, int dim2)

Utility to malloc 2D array.

- int \*\* allocate\_2D\_array\_int (int dim1, int dim2)
- void deallocate 2D array (double \*\*array, int dim1, int dim2)

Utility to free malloc'd 2D array.

- void deallocate\_2D\_array (int \*\*array, int dim1, int dim2)
- double \*\*\* allocate 3D array (int dim1, int dim2, int dim3)

Utility to malloc 3D array.

- int \*\*\* allocate\_3D\_array\_int (int dim1, int dim2, int dim3)
- void deallocate\_3D\_array (double \*\*\*array, int dim1, int dim2, int dim3)

Utility to free malloc'd 2D array.

void deallocate\_3D\_array (int \*\*\*array, int dim1, int dim2, int dim3)

Utility to free malloc'd 2D array.

void read\_file (std::string filename, double \*\*output, int rows, int cols)

Utility to read in data.

• void read\_file (std::string filename, double \*output)

Utility to read in data (single dimension vector)

• void read\_LOSC\_data\_file (std::string filename, double \*output, double \*data\_start\_time, double \*duration, double \*fs)

Read data file from LIGO Open Science Center.

void read\_LOSC\_PSD\_file (std::string filename, double \*\*output, int rows, int cols)

Read PSD file from LIGO Open Science Center.

• void allocate\_LOSC\_data (std::string \*data\_files, std::string psd\_file, int num\_detectors, int psd\_length, int data\_file\_length, double trigger\_time, std::complex< double > \*\*data, double \*\*psds, double \*\*freqs)

Prepare data for MCMC directly from LIGO Open Science Center.

- void free\_LOSC\_data (std::complex < double > \*\*data, double \*\*psds, double \*\*freqs, int num\_detectors, int length)
- void tukey\_window (double \*window, int length, double alpha)

Tukey window function for FFTs.

void write\_file (std::string filename, double \*\*input, int rows, int cols)

Utility to write 2D array to file.

void write\_file (std::string filename, double \*input, int length)

Utility to write 1D array to file.

• void celestial\_horizon\_transform (double RA, double DEC, double gps\_time, double LONG, double LAT, double \*phi, double \*theta)

Utility to transform from celestial coord RA and DEC to local horizon coord for detector response functions.

double gps\_to\_GMST (double gps\_time)

Utility to transform from gps time to GMST https://aa.usno.navy.mil/fag/docs/GAST.php.

double gps\_to\_JD (double gps\_time)

Utility to transform from gps to JD.

void transform\_cart\_sph (double \*cartvec, double \*sphvec)

utility to transform a vector from cartesian to spherical (radian)

void transform sph cart (double \*sphvec, double \*cartvec)

utility to transform a vector from spherical (radian) to cartesian

template < class T >

std::complex< T > cpolar (T mag, T phase)

template < class T >

std::complex < T > XLALSpinWeightedSphericalHarmonic (T theta, T phi, int s, int I, int m)

• template std::complex< double > XLALSpinWeightedSphericalHarmonic< double > (double, double, int, int, int)

- template std::complex< adouble > XLALSpinWeightedSphericalHarmonic< adouble > (adouble, adouble, int, int, int)
- template std::complex< double > cpolar< double > (double, double)
- template std::complex< adouble > cpolar< adouble > (adouble, adouble)

# 9.32.1 Detailed Description

General utilities that are not necessarily specific to any part of the project at large

#### 9.32.2 Function Documentation

#### 9.32.2.1 allocate\_2D\_array()

Utility to malloc 2D array.

# 9.32.2.2 allocate\_3D\_array()

Utility to malloc 3D array.

# 9.32.2.3 allocate\_LOSC\_data()

Prepare data for MCMC directly from LIGO Open Science Center.

Trims data for Tobs (determined by PSD file) 3/4\*Tobs in front of trigger, and 1/4\*Tobs behind

Currently, default to sampling frequency and observation time set by PSD – cannot be customized

Output is in order of PSD columns – string vector of detectos MUST match order of PSD cols

Output shapes—psds = [num\_detectors][psd\_length] data = [num\_detectors][psd\_length]

freqs = [num\_detectors][psd\_length]

Total observation time = 1/(freq[i] - freq[i-1]) (from PSD file)

Sampling frequency fs = max frequency from PSD file

ALLOCATES MEMORY - must be freed to prevent memory leak

#### **Parameters**

	data_files	Vector of strings for each detector file from LOSC
psd_file		String of psd file from LOSC
	num_detectors	Number of detectors to use
	psd_length	Length of the PSD file (number of rows of DATA)
	data_file_length	Length of the data file (number of rows of DATA)
	trigger_time	Time for the signal trigger (GPS)
out	data	Output array of data for each detector
out	psds	Output array of psds for each detector
out	freqs	Output array of freqs for each detector

## 9.32.2.4 calculate\_chirpmass()

Calculates the chirp mass from the two component masses.

The output units are whatever units the input masses are

#### 9.32.2.5 calculate\_mass1()

Calculates the larger mass given a chirp mass and symmetric mass ratio.

Units of the output match the units of the input chirp mass

# 9.32.2.6 calculate\_mass2()

Calculates the smaller mass given a chirp mass and symmetric mass ratio.

Units of the output match the units of the input chirp mass

### 9.32.2.7 celestial\_horizon\_transform()

Utility to transform from celestial coord RA and DEC to local horizon coord for detector response functions.

Outputs are the spherical polar angles defined by North as 0 degrees azimuth and the normal to the earth as 0 degree polar

# **Parameters**

	RA	Right acsension (rad)
	DEC	Declination (rad)
	gps_time	GPS time
	LONG	Longitude (rad)
	LAT	Latitude (rad)
out	phi	horizon azimuthal angle (rad)
out	theta	horizon polar angle (rad)

# 9.32.2.8 cosmology\_interpolation\_function()

Custom interpolation function used in the cosmology calculations.

Power series in half power increments of x, up to 11/2. powers of x

# 9.32.2.9 deallocate\_2D\_array()

Utility to free malloc'd 2D array.

# **9.32.2.10** deallocate\_3D\_array() [1/2]

Utility to free malloc'd 2D array.

#### 9.32.2.11 deallocate\_3D\_array() [2/2]

```
void deallocate_3D_array (
    int *** array,
    int dim1,
    int dim2,
    int dim3 )
```

Utility to free malloc'd 2D array.

# 9.32.2.12 DL\_from\_Z()

```
double DL_from_Z ( \label{eq:cosmology} \mbox{double $Z$,} \\ \mbox{std::string $cosmology$ )}
```

Calculates the luminosity distance given the redshift.

Based on Astropy.cosmology calculations – see python script in the ./data folder of the project – numerically calculated given astropy.cosmology's definitions ( http://docs.astropy.org/en/stable/cosmology/) and used scipy.optimize to fit to a power series, stepping in half powers of Z. These coefficients are then output to a header file ( $D_Z$ -config.h) which are used here to calculate distance. Custom cosmologies etc can easily be acheived by editing the python script  $D_Z$ -config.py, the c++ functions do not need modification. They use whatever data is available in the header file. If the functional form of the fitting function changes, these functions DO need to change.

5 cosmological models are available (this argument must be spelled exactly):

PLANCK15, PLANCK13, WMAP9, WMAP7, WMAP5

# 9.32.2.13 free\_LOSC\_data()

/brief Free data allocated by prep\_LOSC\_data function

# 9.32.2.14 initiate\_LumD\_Z\_interp()

Function that uses the GSL libraries to interpolate pre-calculated Z-D L data.

Initiates the requried functions - GSL interpolation requires allocating memory before hand

## 9.32.2.15 pow\_int()

Local power function, specifically for integer powers.

Much faster than the std version, because this is only for integer powers

# 9.32.2.16 printProgress()

routine to print the progress of a process to the terminal as a progress bar

Call everytime you want the progress printed

Utility to read in data.

Takes filename, and assigns to output[rows][cols]

File must be comma separated doubles

#### **Parameters**

	filename	input filename, relative to execution directory
out	output	array to store output, dimensions rowsXcols
	rows	first dimension
	cols	second dimension

```
9.32.2.18 read_file() [2/2]
```

Utility to read in data (single dimension vector)

Takes filename, and assigns to output[i\*rows + cols]

Output vector must be long enough, no check is done for the length

File must be comma separated doubles

### **Parameters**

	filename	input filename, relative to execution directory	
out	output	output array, assumed to have the proper length of total items	]

### 9.32.2.19 read\_LOSC\_data\_file()

```
void read_LOSC_data_file (
    std::string filename,
    double * output,
    double * data_start_time,
    double * duration,
    double * fs )
```

Read data file from LIGO Open Science Center.

Convenience function for cutting off the first few lines of text

#### **Parameters**

	filename	input filename
out	output	Output data
out	data_start_time	GPS start time of the data in file
out	duration	Duration of the signal
out	fs	Sampling frequency of the data

# 9.32.2.20 read\_LOSC\_PSD\_file()

```
void read_LOSC_PSD_file (
    std::string filename,
    double ** output,
    int rows,
    int cols )
```

Read PSD file from LIGO Open Science Center.

Convenience function for cutting off the first few lines of text

# 9.32.2.21 transform\_cart\_sph()

utility to transform a vector from cartesian to spherical (radian)

order:

cart: x, y, z

spherical: r, polar, azimuthal

## 9.32.2.22 transform\_sph\_cart()

utility to transform a vector from spherical (radian) to cartesian

order:

cart: x, y, z

spherical: r, polar, azimuthal

# 9.32.2.23 tukey\_window()

Tukey window function for FFTs.

As defined by https://en.wikipedia.org/wiki/Window\_function

Utility to write 2D array to file.

Grid of data, comma separated

Grid has rows rows and cols columns

filename	Filename of output file, relative to execution directory
input	Input 2D array pointer array[rows][cols]
rows	First dimension of array
cols	second dimension of array

```
double * input,
int length )
```

Utility to write 1D array to file.

Single column of data

#### **Parameters**

filename	Filename of output file, relative to execution directory
input	input 1D array pointer array[length]
length	length of array

# 9.32.2.26 XLALSpinWeightedSphericalHarmonic()

# Shamelessly stolen from LALsuite

### **Parameters**

theta	polar angle (rad)	
phi	azimuthal angle (rad)	
s	spin weight	
1	mode number l	
m	mode number m	

# 9.32.2.27 Z\_from\_DL()

Calculates the redshift given the luminosity distance.

Based on Astropy.cosmology calculations – see python script in the ./data folder of the project – numerically calculated given astropy.cosmology's definitions ( http://docs.astropy.org/en/stable/cosmology/) and used scipy.optimize to fit to a power series, stepping in half powers of DL. These coefficients are then output to a header file (D\_Z\_config.h) which are used here to calculate redshift. Custom cosmologies etc can easily be acheived by editing the python script D\_Z\_config.py, the c++ functions do not need modification. They use whatever data is available in the header file.

5 cosmological models are available (this argument must be spelled exactly, although case insensitive):

PLANCK15, PLANCK13, WMAP9, WMAP7, WMAP5

### **9.32.2.28 Z\_from\_DL\_interp()** [1/2]

Function that returns Z from a given luminosity Distance - only Planck15

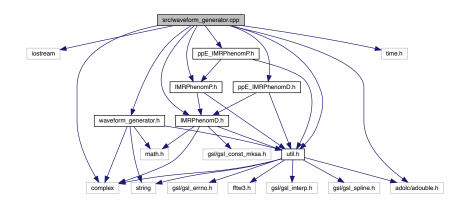
adouble version for ADOL-C calculations

Function that returns Z from a given luminosity Distance - only Planck15

# 9.33 src/waveform\_generator.cpp File Reference

```
#include <iostream>
#include "waveform_generator.h"
#include "IMRPhenomD.h"
#include "IMRPhenomP.h"
#include "ppE_IMRPhenomD.h"
#include "util.h"
#include <complex>
#include <time.h>
#include <adolc/adouble.h>
```

Include dependency graph for waveform\_generator.cpp:



### **Functions**

• int fourier\_waveform (double \*frequencies, int length, std::complex< double > \*waveform\_plus, std

::complex< double > \*waveform\_cross, string generation\_method, gen\_params \*parameters)

Function to produce the plus/cross polarizations of an quasi-circular binary.

- int fourier\_waveform (double \*frequencies, int length, double \*waveform\_plus\_real, double \*waveform
   —plus\_imag, double \*waveform\_cross\_real, double \*waveform\_cross\_imag, string generation\_method,
   gen\_params \*parameters)
- int fourier\_waveform (double \*frequencies, int length, std::complex< double > \*waveform, string generation\_method, gen\_params \*parameters)

Function to produce the (2,2) mode of an quasi-circular binary.

- int fourier\_waveform (double \*frequencies, int length, double \*waveform\_real, double \*waveform\_imag, string generation\_method, gen\_params \*parameters)
- int fourier\_amplitude (double \*frequencies, int length, double \*amplitude, string generation\_method, gen\_params \*parameters)

Function to produce the amplitude of the (2,2) mode of an quasi-circular binary.

• int fourier\_phase (double \*frequencies, int length, double \*phase, string generation\_method, gen\_params \*parameters)

Function to produce the phase of the (2,2) mode of an quasi-circular binary.

# 9.33.1 Detailed Description

File that handles the construction of the (2,2) waveform as described by IMRPhenomD by Khan et. al.

Builds a waveform for given DETECTOR FRAME parameters

### 9.33.2 Function Documentation

#### 9.33.2.1 fourier\_amplitude()

Function to produce the amplitude of the (2,2) mode of an quasi-circular binary.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

frequencies	double array of frequencies for the waveform to be evaluated at
length	integer length of all the arrays
amplitude	output array for the amplitude
generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY

#### 9.33.2.2 fourier phase()

Function to produce the phase of the (2,2) mode of an quasi-circular binary.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

#### **Parameters**

frequencies	double array of frequencies for the waveform to be evaluated at	
length	integer length of all the arrays	
phase	output array for the phase	
generation_method	generation_method   String that corresponds to the generation method - MUST BE SPELLED EXACTL	

# **9.33.2.3** fourier\_waveform() [1/4]

Function to produce the plus/cross polarizations of an quasi-circular binary.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

This puts the responsibility on the user to pass the necessary parameters

NEED TO OUTLINE OPTIONS FOR EACH METHOD IN DEPTH

NEW PHASE OPTIONS for

PHENOMD ONLY:

If phic is assigned, the reference frequency and reference phase are IGNORED.

If Phic is unassigned, a reference phase AND a reference frequency are looked for. If no options are found, both are set to 0.

If to is assigned, it is used.

If to is unassigned, the waveform is shifted so the merger happens at 0.

PhenomPv2:

PhiRef and f ref are required, phic is not an option.

tc, if specified, is used with the use of interpolation. If not, tc is set such that coalescence happens at t=0

#### **Parameters**

	frequencies double array of frequencies for the waveform to be evaluated at		
	length integer length of all the arrays		
out	waveform_plus	complex array for the output plus polarization waveform	
out	waveform_cross	_cross complex array for the output cross polarization waveform	
	generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY	
	parameters	structure containing all the source parameters	

# **9.33.2.4** fourier\_waveform() [2/4]

#### **Parameters**

frequencies	double array of frequencies for the waveform to be evaluated at	
length	integer length of all the arrays	
waveform_plus_real	complex array for the output waveform	
waveform_plus_imag	complex array for the output waveform	
waveform_cross_real	complex array for the output waveform	
waveform_cross_imag	complex array for the output waveform	
generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY	
parameters	structure containing all the source parameters	

# **9.33.2.5** fourier\_waveform() [3/4]

Function to produce the (2,2) mode of an quasi-circular binary.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

#### **Parameters**

frequencies	double array of frequencies for the waveform to be evaluated at	
length	integer length of all the arrays	
waveform	complex array for the output waveform	
generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY	
parameters	structure containing all the source parameters	

# **9.33.2.6** fourier\_waveform() [4/4]

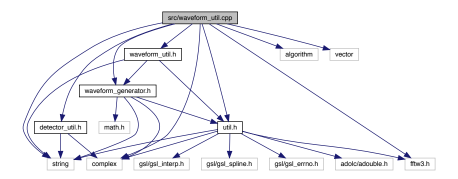
#### **Parameters**

frequencies	double array of frequencies for the waveform to be evaluated at	
length	integer length of all the arrays	
waveform_real	complex array for the output waveform	
waveform_imag	complex array for the output waveform	
generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY	
parameters	structure containing all the source parameters	

# 9.34 src/waveform\_util.cpp File Reference

```
#include "waveform_util.h"
#include "util.h"
#include "waveform_generator.h"
#include "detector_util.h"
#include <fftw3.h>
#include <algorithm>
#include <complex>
#include <vector>
#include <string>
```

Include dependency graph for waveform\_util.cpp:



#### **Functions**

• double data\_snr\_maximized\_extrinsic (double \*frequencies, int length, std::complex< double > \*data, double \*psd, std::string detector, std::string generation method, gen params \*param)

Utility to calculate the snr of a fourier transformed data stream while maximizing over the coalescence parameters phic and tc.

• double data\_snr\_maximized\_extrinsic (double \*frequencies, int length, double \*data\_real, double \*data\_← imag, double \*psd, std::string detector, std::string generation method, gen params \*param)

Light wrapper for the data\_snr\_maximized\_extrinsic method.

double calculate\_snr (std::string detector, std::complex< double > \*waveform, double \*frequencies, int length)

Caclulates the snr given a detector and waveform (complex) and frequencies.

- int fourier\_detector\_response (double \*frequencies, int length, std::complex< double > \*hplus, std
   ::complex< double > \*hcross, std::complex< double > \*detector\_response, double theta, double phi, std
   ::string detector)
- int fourier\_detector\_response (double \*frequencies, int length, std::complex< double > \*hplus, std
   ::complex< double > \*hcross, std::complex< double > \*detector\_response, double theta, double phi, double
   psi, std::string detector)
- int fourier\_detector\_response\_equatorial (double \*frequencies, int length, std::complex< double > \*hplus, std::complex< double > \*hcross, std::complex< double > \*detector\_response, double ra, double dec, double psi, double gmst, std::string detector)
- int fourier\_detector\_response (double \*frequencies, int length, std::complex< double > \*response, std 
  ::string detector, std::string generation\_method, gen\_params \*parameters)

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary.

• int fourier\_detector\_response\_equatorial (double \*frequencies, int length, std::complex< double > \*response, std::string detector, std::string generation method, gen params \*parameters)

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary for equatorial coordinates.

• int fourier\_detector\_amplitude\_phase (double \*frequencies, int length, double \*amplitude, double \*phase, std::string detector, std::string generation\_method, gen\_params \*parameters)

Calculates the amplitude (magnitude) and phase (argument) of the response of a given detector.

# 9.34.1 Detailed Description

Utilities for waveforms - SNR calculation and detector response

includes snr and detector response

# 9.34.2 Function Documentation

#### 9.34.2.1 calculate\_snr()

```
double calculate_snr (
          std::string detector,
          std::complex< double > * waveform,
          double * frequencies,
          int length )
```

Caclulates the snr given a detector and waveform (complex) and frequencies.

This function computes the un-normalized snr: \sqrt( ( H | H ) )

#### **Parameters**

detector	detector name - must match the string of populate_noise precisely	
waveform	complex waveform	
frequencies	double array of frequencies that the waveform is evaluated at	
length	length of the above two arrays	

## 9.34.2.2 data\_snr\_maximized\_extrinsic() [1/2]

Utility to calculate the snr of a fourier transformed data stream while maximizing over the coalescence parameters phic and tc.

The gen\_params structure holds the parameters for the template to be used (the maximimum likelihood parameters)

frequencies	Frequencies used by data	
length	length of the data	
data	input data in the fourier domain	
	·	
psd	PSD for the detector that created the data	
detector	Name of the detector –See noise_util for options	
generation_method	Generation method for the template – See waveform_generation.cpp for options	
param	gen_params structure for the template	

#### 9.34.2.3 data\_snr\_maximized\_extrinsic() [2/2]

Light wrapper for the data\_snr\_maximized\_extrinsic method.

Splits the data into real and imaginary, so all the arguments are C-safe

#### **Parameters**

frequencies	Frequencies used by data	
length	length of the data	
data_real	input data in the fourier domain – real part	
data_imag	input data in the fourier domain – imaginary part	
psd	PSD for the detector that created the data	
detector	Name of the detector –See noise_util for options	
generation_method	Generation method for the template – See waveform_generation.cpp for options	
param	gen_params structure for the template	

# 9.34.2.4 fourier\_detector\_amplitude\_phase()

Calculates the amplitude (magnitude) and phase (argument) of the response of a given detector.

This is for general waveforms, and will work for precessing waveforms

Not as fast as non-precessing, but that can't be helped. MUST include plus/cross polarizations

#### 9.34.2.5 fourier\_detector\_response() [1/3]

#### **Parameters**

	frequencies	array of frequencies corresponding to waveform
	length	length of frequency/waveform arrays
	hcross	precomputed cross polarization of the waveform
out	detector_response	detector response
	theta	polar angle (rad) theta in detector frame
	phi	azimuthal angle (rad) phi in detector frame
	detector	detector - list of supported detectors in noise_util

# 9.34.2.6 fourier\_detector\_response() [2/3]

	frequencies	array of frequencies corresponding to waveform
	length	length of frequency/waveform arrays
	hcross	precomputed cross polarization of the waveform
out	detector_response	detector response
	theta	polar angle (rad) theta in detector frame
	phi	azimuthal angle (rad) phi in detector frame
	psi	polarization angle (rad) phi in detector frame
	detector	detector - list of supported detectors in noise_util

#### 9.34.2.7 fourier\_detector\_response() [3/3]

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

This puts the responsibility on the user to pass the necessary parameters

Detector options include classic interferometers like LIGO/VIRGO (coming soon: ET and LISA)

This is a wrapper that combines generation with response functions: if producing mulitple responses for one waveform (ie stacking Hanford, Livingston, and VIRGO), it will be considerably more efficient to calculate the waveform once, then combine each response manually

#### **Parameters**

	frequencies	double array of frequencies for the waveform to be evaluated at	
	length integer length of all the arrays		
out	response	complex array for the output plus polarization waveform	
	generation_method   String that corresponds to the generation method - MUST BE SPELLED EXACTL		
	parameters	structure containing all the source parameters	

# 9.34.2.8 fourier\_detector\_response\_equatorial() [1/2]

	frequencies	array of frequencies corresponding to waveform
	length	length of frequency/waveform arrays
	hcross	precomputed cross polarization of the waveform
out	detector_response	detector response
	ra	Right Ascension in rad

#### **Parameters**

dec	Declination in rad
psi	polarization angle (rad)
gmst	greenwich mean sidereal time
detector	detector - list of supported detectors in noise_util

# 9.34.2.9 fourier\_detector\_response\_equatorial() [2/2]

Function to produce the detector response caused by impinging gravitational waves from a quasi-circular binary for equatorial coordinates.

By using the structure parameter, the function is allowed to be more flexible in using different method of waveform generation - not all methods use the same parameters

This puts the responsibility on the user to pass the necessary parameters

Detector options include classic interferometers like LIGO/VIRGO (coming soon: ET and LISA)

This is a wrapper that combines generation with response functions: if producing mulitple responses for one waveform (ie stacking Hanford, Livingston, and VIRGO), it will be considerably more efficient to calculate the waveform once, then combine each response manually

	frequencies	double array of frequencies for the waveform to be evaluated at
	length	integer length of all the arrays
out	response	complex array for the output plus polarization waveform
	generation_method	String that corresponds to the generation method - MUST BE SPELLED EXACTLY
	parameters	structure containing all the source parameters

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