# **MRCPP**

Release 1.3.0

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The MultiResolution Computation Program Package (MRCPP) is a general purpose numerical mathematics library based on multiresolution analysis and the multiwavelet basis which provide low-scaling algorithms as well as rigorous error control in numerical computations.

The code is being developed at the Hylleraas Centre for Quantum Molecular Sciences at UiT - The Arctic University of Norway.

The code can be found on GitHub.

INSTALLATION 1

2 INSTALLATION

## **ONE**

## **OBTAINING THE CODE**

The latest version of MRCPP is available on GitHub:

\$ git clone git@github.com:MRChemSoft/mrcpp.git

**TWO** 

## **BUILDING THE CODE**

## 2.1 Prerequisites

- g++-5.4 or later (std=c++14)
- CMake version 3.11 or higher.
- Eigen version 3.3 or higher.
- BLAS (optional)

## 2.2 Configuration

The configuration and build process is managed through CMake, and a setup script is provided for the configuration step. MRCPP's only dependency is Eigen3, which will be downloaded at configure time unless it is already available on the system. If you have a local version *not* under the system path, you can point to it before running setup:

```
$ export EIGEN3_ROOT=/path/to/eigen3
$ ./setup [options] [<builddir>]
```

The setup script will create a directory called *<builddir>* (default build) and run CMake. There are several options available for the setup, and the most important are:

- --cxx=<CXX> C++ compiler [default: g++]
- --omp Enable OpenMP parallelization [default: False]
- --mpi Enable MPI parallelization [default: False]
- --enable-tests Enable tests [default: True]
- --enable-examples Enable tests [default: False]
- --type=<TYPE> Set the CMake build type (debug, release, relwithdebinfo, minsizerel) [default: release]
- --prefix=<PATH> Set the install path for make install
- -h --help List all options

## 2.3 Compilation

After successful configuration, the code is compiled using the make command in the *<builddir>* directory:

```
$ cd <builddir>
```

<sup>\$</sup> make

**THREE** 

## **RUNNING TESTS**

A set of tests is provided with the code to verify that the code compiled properly. To compile the test suite, add the --enable-tests option to setup, then run the tests with ctest:

- \$ ./setup --enable-tests build
- \$ cd build
- \$ make
- \$ ctest

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## **RUNNING EXAMPLES**

In addition to the test suite, the code comes with a number of small code snippets that demonstrate the features and the API of the library. These are located in the examples directory. To compile the example codes, add the enable-examples option to setup, and the example executables can be found under <build-dir>/bin/. E.g. to compile and run the MW projection example:

```
$ ./setup --enable-examples build-serial
$ cd build-serial
$ make
$ bin/projection
```

The shared memory parallelization (OpenMP) is controlled by the environment variable OMP\_NUM\_THREADS (make sure you have compiled with the --omp option to setup). E.g. to compile and run the Poisson solver example using 10 CPU cores:

```
$ ./setup --enable-examples --omp build-omp
$ cd build-omp
$ make
$ OMP_NUM_THREADS=10 bin/poisson
```

To run in MPI parallel, use the mpirun (or equivalent) command (make sure you have compiled with the --mpi option to setup, and used MPI compatible compilers, e.g. --cxx=mpicxx). Only examples with an *mpi* prefix will be affected by running in MPI:

```
$ ./setup --cxx=mpicxx --enable-examples --mpi build-mpi
$ cd build-mpi
$ make
$ mpirun -np 4 bin/mpi_send_tree
```

To run in hybrid OpenMP/MPI parallel, simply combine the two above:

```
$ ./setup --cxx=mpicxx --enable-examples --omp --mpi build-hybrid
$ cd build-hybrid
$ make
$ export OMP_NUM_THREADS=5
$ mpirun -np 4 bin/mpi_send_tree
```

Note that the core of MRCPP is *only* OpenMP parallelized. All MPI data or work distribution must be done manually in the application program, using the tools provided by MRCPP (see the Parallel section of the API).

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## **PILOT CODE**

Finally, MRCPP comes with a personal sandbox where you can experiment and test new ideas, without messing around in the git repository. In the pilot/directory you will find a skeleton code called mrcpp.cpp.sample. To trigger a build, re-name (copy) this file to mrcpp.cpp:

```
$ cd pilot
$ cp mrcpp.cpp.sample mrcpp.cpp
```

Now a corresponding executable will be build in <builddir>/bin/mrcpp-pilot/. Feel free to do whatever you like in your own pilot code, but please don't add this file to git. Also, please don't commit any changes to the existing examples (unless you know what you're doing).

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## MRCPP AS A DEPENDENCY

Building MRCPP provides CMake configuration files exporting the libraries and headers as targets to be consumed by third-party projects also using CMake:

```
$ ./setup --prefix=$HOME/Software/mrcpp
$ cd build
$ make
$ ctest
$ make install
```

Now libraries, headers and CMake configuration files can be found under the given prefix:

```
mrcpp/
— include/
— MRCPP/
— lib64/
— libmrcpp.a
— libmrcpp.so -> libmrcpp.so.1*
— libmrcpp.so.1*
— share/
— cmake/
```

As an example, the pilot sample can be built with the following CMakeLists.txt:

```
cmake_minimum_required(VERSION 3.11 FATAL_ERROR)
project(UseMRCPP LANGUAGES CXX)

set(CMAKE_CXX_STANDARD 14)
set(CMAKE_CXX_EXTENSIONS OFF)
set(CMAKE_CXX_EXTENSIONS OFF)
set(CMAKE_CXX_STANDARD_REQUIRED ON)
include(GNUInstallDirs)

set(CMAKE_ARCHIVE_OUTPUT_DIRECTORY ${PROJECT_BINARY_DIR}/${CMAKE_INSTALL_LIBDIR})
set(CMAKE_LIBRARY_OUTPUT_DIRECTORY ${PROJECT_BINARY_DIR}/${CMAKE_INSTALL_LIBDIR})
set(CMAKE_RUNTIME_OUTPUT_DIRECTORY ${PROJECT_BINARY_DIR}/${CMAKE_INSTALL_BINDIR})

find_package(MRCPP CONFIG REQUIRED)
get_property(_loc TARGET MRCPP::mrcpp PROPERTY LOCATION)
message(STATUS "Found MRCPP: ${_loc} (found version ${MRCPP_VERSION})")

add_executable(mrcpp mrcpp.cpp)
target_link_libraries(mrcpp
PUBLIC
```

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```
MRCPP::mrcpp
)

set_target_properties(mrcpp
    PROPERTIES
    MACOSX_RPATH ON
    SKIP_BUILD_RPATH OFF
    BUILD_WITH_INSTALL_RPATH OFF
    INSTALL_RPATH "$ORIGIN/../${CMAKE_INSTALL_LIBDIR}"
    INSTALL_RPATH_USE_LINK_PATH ON
)
```

This will set up the include paths and library paths correctly. During configuration you will have to specify *where* the CMake configuration file for MRCPP is located:

```
$ cmake -H. -Bbuild -DMRCPP_DIR=$HOME/Software/share/cmake/MRCPP
```

## **SEVEN**

## INTRODUCTION

The main features of MRCPP are the numerical multiwavelet (MW) representations of functions and operators. Two integral convolution operators are implemented (the Poisson and Helmholtz operators), as well as the partial derivative and arithmetic operators. In addition to the numerical representations there are a limited number of analytic functions that are usually used as starting point for the numerical computations. Also, MRCPP provides three convenience classes (Timer, Printer and Plotter) that can be made available to the application program.

The API consists of seven include files which will be discussed in more detail below:



- **MRCPP/MWFunctions** Provides features for representation and manipulation of real-valued scalar functions in a MW basis, including projection of analytic function, numerical integration and scalar products, as well as arithmetic operations and function mappings.
- **MRCPP/MWOperators** Provides features for representation and application of MW operators. Currently there are three operators available: Poisson, Helmholtz and Cartesian derivative.
- **MRCPP/Gaussians** Provides some simple features for analytical Gaussian functions, useful e.g. to generate initial guesses for MW computations.
- **MRCPP/Parallel** Provides some simple MPI features for MRCPP, in particular the possibility to send complete MW function representations between MPI processes.
- **MRCPP/Printer** Provides simple (parallel safe) printing options. All MRCPP internal printing is done with this class, and the printer must be initialized in order to get any printed output, otherwise MRCPP will run silently.
- **MRCPP/Plotter** Provides options to generate data files for plotting of MW function representations. These include line plots, surface plots and cube plots, as well as grid visualization using geomyiew.
- MRCPP/Timer Provides an accurate timer for the wall clock in parallel computations.

## 7.1 Analytic functions

The general way of defining an analytic function in MRCPP is to use lambdas (or std::function), which provide lightweight functions that can be used on the fly. However, some analytic functions, like Gaussians, are of special importance and have been explicitly implemented with additional functionality (see Gaussian chapter).

In order to be accepted by the MW projector (see MWFunctions chapter), the lambda must have the following signature:

```
auto f = [] (const mrcpp::Coord<D> &r) -> double;
```

e.i. it must take a D-dimensional Cartesian coordinate (mrcpp::Coord<D> is simply an alias for std::array<double, D>), and return a double. For instance, the electrostatic potential from a point nuclear charge Z (in atomic units) is

$$f(r) = \frac{Z}{r}$$

which can be written as the lambda function

```
auto Z = 1.0; // Hydrogen nuclear charge
auto f = [Z] (const mrcpp::Coord<3> &r) -> double {
   auto R = std::sqrt(r[0]*r[0] + r[1]*r[1] + r[2]*r[2]);
   return Z/R;
};
```

Note that the function signature must be exactly as given above, which means that any additional arguments (such as Z in this case) must be given in the capture list (square brackets), see e.g. cppreference.com for more details on lambda functions and how to use the capture list.

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## **MWFUNCTIONS**

Everything that is discussed in the following chapter is available to the application program by including:

#include "MRCPP/MWFunctions"

Multiwavelet (MW) representations of real-valued scalar functions are in MRCPP called FunctionTrees. These are in principle available in any dimension using the template parameter D (in practice D=1,2,3). There are several different ways of constructing MW functions (computing the expansion coefficients in the MW basis):

- · Projection of analytic function
- Arithmetic operations
- · Application of MW operator

The first two will be discribed in this chapter, while the last one regarding operators will be the topic of the next chapter.

The interface for constructing MW representations in MRCPP has a dual focus: on the one hand we want a simple, intuitive way of producing adaptive numerical approximations with guaranteed precision that does not require detailed knowledge of the internals of the MW code and with a minimal number of parameters that have to be set. On the other hand we want the possibility for more detailed control of the construction and refinement of the numerical grid where such control is possible and even necessary. In the latter case it is important to be able to reuse the existing grids in e.g. iterative algorithms without excessive allocation/deallocation of memory.

## 8.1 MultiResolution Analysis (MRA)

template<int D>

class mrcpp::MultiResolutionAnalysis

Class collecting computational domain and MW basis.

In order to combine different functions and operators in mathematical operations, they need to be compatible. That is, they must be defined on the same computational domain and constructed using the same polynomial basis (order and type). This information constitutes an MRA, which needs to be defined and passed as argument to all function and operator constructors, and only functions and operators with compatible MRAs can be combined in subsequent calculations.

### **Public Functions**

MultiResolutionAnalysis (const BoundingBox<D> &bb, const ScalingBasis &sb, int depth = MaxDepth)

Return New MultiResolutionAnalysis object

#### **Parameters**

- [in] bb: Computational domain
- [in] sb: Polynomial basis
- [in] depth: Maximum allowed resolution depth, relative to root scale

## template<int D>

## class mrcpp::BoundingBox

Class defining the computational domain.

The computational domain is made up of a collection of D-dimensional boxes on a particular length scale n. The size of each box is then  $[2^{-n}]^D$ , i.e. higher scale means smaller boxes, and the scale may be negative. The number of boxes can be different in each dimension  $[n_x, n_y, \ldots]$ , but they must all be on the same scale (size). Box translations relative to the world origin *must* be an integer multiple of the given scale size  $2^{-n}$ .

Subclassed by mrcpp::NodeBox< D >

### **Public Functions**

```
BoundingBox (int n = 0, const std::array<int, D > \&l = \{\}, const std::array<int, D > \&nb = \{\}, const std::array<double, D > \&sf = \{\})
```

Return New BoundingBox object

### **Parameters**

- [in] n: Length scale, default 0
- [in] 1: Corner translation, default [0, 0, ...]
- [in] nb: Number of boxes, default [1, 1, ...]
- [in] sf: Scaling factor, default [1.0, 1.0, ...]

## $\textbf{class} \ \texttt{mrcpp::LegendreBasis:public} \ mrcpp::ScalingBasis$

Legendre scaling functions as defined by Alpert, SIAM J Math Anal 24 (1), 246 (1993).

#### **Public Functions**

## LegendreBasis (int k)

Return New LegendreBasis object

## **Parameters**

• [in] k: Polynomial order of basis, 1 < k < 40

## class mrcpp::InterpolatingBasis: public mrcpp::ScalingBasis

Interpolating scaling functions as defined by Alpert et al, J Comp Phys 182, 149-190 (2002).

## **Public Functions**

InterpolatingBasis (int k)

Return New Interpolating Basis object

**Parameters** 

• [in] k: Polynomial order of basis, 1 < k < 40

## 8.2 FunctionTree

template<int D>

**class** mrcpp::**FunctionTree**: **public** mrcpp::MWTree<*D*>, **public** mrcpp::RepresentableFunction<*D*> Function representation in MW basis.

Constructing a full grown *FunctionTree* involves a number of steps, including setting up a memory allocator, constructing root nodes according to the given MRA, building an adaptive tree structure and computing MW coefficients. The *FunctionTree* constructor does only half of these steps: It takes an MRA argument, which defines the computational domain and scaling basis (these are fixed parameters that cannot be changed after construction). The tree is initialized with a memory allocator and a set of root nodes, but it does not compute any coefficients and the function is initially *undefined*. An undefined *FunctionTree* will have a well defined tree structure (at the very least the root nodes of the given MRA, but possibly with additional refinement) and its MW coefficient will be allocated but uninitialized, and its square norm will be negative (minus one).

#### **Public Functions**

FunctionTree (const MultiResolutionAnalysis<D> &mra, SharedMemory \*sh\_mem = nullptr)

Constructs an uninitialized tree, containing only empty root nodes. If a shared memory pointer is provided the tree will be allocated in this shared memory window, otherwise it will be local to each MPI process.

Return New FunctionTree object

## **Parameters**

- [in] mra: Which MRA the function is defined
- [in] sh mem: Pointer to MPI shared memory block

## 8.2.1 Creating defined FunctionTrees

The following functions will *define* MW coefficients where there are none, and thus *require* that the output FunctionTree is in an *undefined* state. All functions marked with 'adaptive grid' will use the same building algorithm:

- 1. Start with an initial guess for the grid
- 2. Compute the MW coefficients for the output function on the current grid
- 3. Refine the grid where necessary based on the local wavelet norm
- 4. Iterate points 2 and 3 until the grid is converged

With a *negative* precision argument, the grid will be *fixed*, e.i. it will not be refined beyond the initial grid. There is also an argument to limit the number of *extra* refinement levels beyond the initial grid, in which the adaptive refinement will stop, even if the local precision requirement is not met.

```
void mrcpp::MWTree::setZero()
```

Set the MW coefficients to zero, fixed grid.

Keeps the node structure of the tree, even though the zero function is representable at depth zero. Use cropTree to remove unnecessary nodes.

## template<int D>

```
void mrcpp::project (double prec, FunctionTree<D> &out, RepresentableFunction<D> &inp, int maxIter,
bool absPrec)
```

Project an analytic function onto the MW basis, adaptive grid.

The output function will be computed using the general algorithm:

- · Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

## **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] inp: Input function
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

### template<int D>

```
void mrcpp::copy func (FunctionTree<D> &out, FunctionTree<D> &inp)
```

Copy function from one tree onto the grid of another tree, fixed grid.

The output function will be computed using the general algorithm:

- · Loop through current leaf nodes of the output tree
- Copy MW coefs from the corresponding input node

## **Parameters**

- [out] out: Output function
- [in] inp: Input function

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called and will overwrite any existing coefs.

## template<int D>

Addition of several MW function representations, adaptive grid.

The output function will be computed as the sum of all input functions in the vector (including their numerical coefficients), using the general algorithm:

- Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

#### **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] inp: Vector of input function
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

## template<int D>

```
void mrcpp::add (double prec, FunctionTree<D> &out, double a, FunctionTree<D> &inp_a, double b,
FunctionTree<D> &inp_b, int maxIter, bool absPrec)
```

Addition of two MW function representations, adaptive grid.

The output function will be computed as the sum of the two input functions (including the numerical coefficient), using the general algorithm:

- · Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

## **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] a: Numerical coefficient of function a
- [in] inp\_a: Input function a
- [in] b: Numerical coefficient of function b
- [in] inp\_b: Input function b
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

template<int D>

void mrcpp::multiply (double prec, FunctionTree<D> &out, FunctionTreeVector<D> &inp, int maxIter,
bool absPrec, bool useMaxNorms)

Multiplication of several MW function representations, adaptive grid.

The output function will be computed as the product of all input functions in the vector (including their numerical coefficients), using the general algorithm:

- · Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

### **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] inp: Vector of input function
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision
- [in] useMaxNorms: Build output tree based on norm estimates from input

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

## template<int D>

void mrcpp::multiply (double prec, FunctionTree<D> &out, double c, FunctionTree<D> &inp\_a, FunctionTree<D> &inp\_b, int maxIter, bool absPrec, bool useMaxNorms)

Multiplication of two MW function representations, adaptive grid.

The output function will be computed as the product of the two input functions (including the numerical coefficient), using the general algorithm:

- · Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

## **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] c: Numerical coefficient
- [in] inp\_a: Input function a
- [in] inp\_b: Input function b
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

• [in] useMaxNorms: Build output tree based on norm estimates from input

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

## template<int D>

void mrcpp::square (double prec, FunctionTree<D> &out, FunctionTree<D> &inp, int maxIter, bool absPrec)

Out-of-place square of MW function representations, adaptive grid.

The output function will be computed as the square of the input function, using the general algorithm:

- Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

### **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] inp: Input function to square
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

#### template<int D>

void mrcpp::power (double prec, FunctionTree<D> &out, FunctionTree<D> &inp, double p, int maxIter,
bool absPrec)

Out-of-place power of MW function representations, adaptive grid.

The output function will be computed as the input function raised to the given power, using the general algorithm:

- Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

### **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] inp: Input function to square
- [in] p: Numerical power

- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

## template<int D>

void mrcpp: :dot (double prec, FunctionTree<D> &out, FunctionTreeVector<D> &inp\_a, FunctionTreeVector<D> &inp\_b, int maxIter, bool absPrec)

Dot product of two MW function vectors, adaptive grid.

The output function will be computed as the dot product of the two input vectors (including their numerical coefficients). The precision parameter is used only in the multiplication part, the final addition will be on the fixed union grid of the components.

#### **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] inp\_a: Input function vector
- [in] inp\_b: Input function vector
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

**Note** The length of the input vectors must be the same.

### template<int D>

void mrcpp::map (double prec, FunctionTree<D> &out, FunctionTree<D> &inp, FMap fmap, int maxIter,
bool absPrec)

map a MW function onto another representations, adaptive grid

The output function tree will be computed by mapping the input tree values through the fmap function, using the general algorithm:

- Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

## **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] inp: Input function
- [in] fmap: mapping function
- [in] maxIter: Maximum number of refinement iterations in output tree
- [in] absPrec: Build output tree based on absolute precision

No assumption is made for how the mapping function looks. It is left to the end-user to guarantee that the mapping function does not lead to numerically unstable/inaccurate situations (e.g. divide by zero, overflow, etc...)

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

## 8.2.2 Creating undefined FunctionTrees

The grid of a FunctionTree can also be constructed without computing any MW coefficients:

## template<int D>

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Build empty grid based on info from analytic function.

The grid of the output function will be EXTENDED using the general algorithm:

- Loop through current leaf nodes of the output tree
- Refine node based on custom split check from the function
- Repeat until convergence or maxIter is reached
- maxIter < 0 means no bound

### **Parameters**

- [out] out: Output tree to be built
- [in] inp: Input function
- [in] maxIter: Maximum number of refinement iterations in output tree

Note This algorithm will start at whatever grid is present in the out tree when the function is called. It requires that the functions is Visible AtScale() and is ZeroOnInterval() is implemented in the particular Representable Function.

### template<int D>

void mrcpp::build grid(FunctionTree<D> &out, const GaussExp<D> &inp, int maxIter)

Build empty grid based on info from Gaussian expansion.

The grid of the output function will be EXTENDED using the general algorithm:

- Loop through current leaf nodes of the output tree
- Refine node based on custom split check from the function
- Repeat until convergence or maxIter is reached
- maxIter < 0 means no bound

## **Parameters**

- [out] out: Output tree to be built
- [in] inp: Input Gaussian expansion
- [in] maxIter: Maximum number of refinement iterations in output tree

Note This algorithm will start at whatever grid is present in the out tree when the function is called. It will loop through the Gaussians in the expansion and extend the grid based on the position and exponent of each term. Higher exponent means finer resolution.

## template<int D>

```
void mrcpp::build grid(FunctionTree<D> &out, FunctionTree<D> &inp, int maxIter)
```

Build empty grid based on another MW function representation.

The grid of the output function will be EXTENDED with all existing nodes in corresponding input function, using the general algorithm:

- · Loop through current leaf nodes of the output tree
- Refine node if the corresponding node in the input has children
- Repeat until all input nodes are covered or maxIter is reached
- maxIter < 0 means no bound

### **Parameters**

- [out] out: Output tree to be built
- [in] inp: Input tree
- [in] maxIter: Maximum number of refinement iterations in output tree

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called. This means that all nodes on the input tree will also be in the final output tree (unless maxIter is reached, but NOT vice versa.

## template<int D>

```
void mrcpp::build_grid(FunctionTree<D> &out, FunctionTree Vector<D> &inp, int maxIter)
     Build empty grid based on several MW function representation.
```

The grid of the output function will be EXTENDED with all existing nodes in all corresponding input functions, using the general algorithm:

- · Loop through current leaf nodes of the output tree
- Refine node if the corresponding node in one of the inputs has children
- Repeat until all input nodes are covered or maxIter is reached
- maxIter < 0 means no bound

#### **Parameters**

- [out] out: Output tree to be built
- [in] inp: Input tree vector
- [in] maxIter: Maximum number of refinement iterations in output tree

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called. This means that the final output grid will contain (at least) the union of the nodes of all input trees (unless maxIter is reached).

## template<int D>

```
void mrcpp::copy_grid (FunctionTree<D> &out, FunctionTree<D> &inp)
     Build empty grid that is identical to another MW grid.
```

**Note** The difference from the corresponding build\_grid function is that this will first clear the grid of the out function, while build\_grid will *extend* the existing grid.

## **Parameters**

- [out] out: Output tree to be built
- [in] inp: Input tree

#### template<int D>

```
void mrcpp::clear_grid(FunctionTree<D> &out)
```

Clear the MW coefficients of a function representation.

**Note** This will only clear the MW coefs in the existing nodes, it will not change the grid of the function. Use FunctionTree::clear() to remove all grid refinement as well.

#### **Parameters**

• [inout] out: Output function to be cleared

```
void mrcpp::FunctionTree::clear()
```

Remove all nodes in the tree.

Leaves the tree inn the same state as after construction, i.e. undefined function containing only root nodes without coefficients. The assigned memory (nodeChunks in SerialTree) is NOT released, but is immediately available to the new function.

## 8.2.3 Changing FunctionTrees

There are also a number of in-place operations that *change* the MW coefficients of a given defined FunctionTree. All changing operations *require* that the FunctionTree is in a *defined* state.

```
void mrcpp::FunctionTree::rescale (double c)
```

In-place multiplication by a scalar, fixed grid.

The leaf node point values of the function will be in-place multiplied by the given coefficient, no grid refinement.

#### **Parameters**

• [in] c: Scalar coefficient

```
void mrcpp::FunctionTree::normalize()
```

In-place rescaling by a function norm  $||f||^{-1}$ , fixed grid.

```
void mrcpp::FunctionTree::add (double c, FunctionTree<D> &inp)
```

In-place addition with MW function representations, fixed grid.

The input function will be added in-place on the current grid of the function, i.e. no further grid refinement.

#### **Parameters**

- [in] c: Numerical coefficient of input function
- [in] inp: Input function to add

```
void mrcpp::FunctionTree::multiply(double c, FunctionTree<D> &inp)
```

In-place multiplication with MW function representations, fixed grid.

The input function will be multiplied in-place on the current grid of the function, i.e. no further grid refinement.

## **Parameters**

• [in] c: Numerical coefficient of input function

• [in] inp: Input function to multiply

```
void mrcpp::FunctionTree::square()
```

In-place square of MW function representations, fixed grid.

The leaf node point values of the function will be in-place squared, no grid refinement.

```
void mrcpp::FunctionTree::power(double p)
```

In-place power of MW function representations, fixed grid.

The leaf node point values of the function will be in-place raised to the given power, no grid refinement.

### **Parameters**

• [in] p: Numerical power

```
void mrcpp::FunctionTree::map(FMapfmap)
```

In-place mapping with a predefined function f(x), fixed grid.

The input function will be mapped in-place on the current grid of the function, i.e. no further grid refinement.

### **Parameters**

• [in] fmap: mapping function

int mrcpp::FunctionTree::crop (double prec, double splitFac = 1.0, bool absPrec = true)

Reduce the precision of the tree by deleting nodes.

This will run the tree building algorithm in "reverse", starting from the leaf nodes, and perform split checks on each node based on the given precision and the local wavelet norm.

#### **Parameters**

- prec: New precision criterion
- splitFac: Splitting factor: 1, 2 or 3
- absPrec: Use absolute precision

**Note** The splitting factor appears in the threshold for the wavelet norm as  $||w|| < 2^{-sn/2}||f||\epsilon$ . In principal, s should be equal to the dimension; in practice, it is set to s=1.

### template<int D>

```
int mrcpp::refine_grid(FunctionTree<D> &out, int scales)
```

Refine the grid of a MW function representation.

This will split ALL leaf nodes in the tree the given number of times, then it will compute scaling coefs of the new nodes, thus leaving the function representation unchanged, but on a larger grid.

**Return** The number of nodes that were split

## **Parameters**

- [inout] out: Output tree to be refined
- [in] scales: Number of refinement levels

#### template<int D>

```
int mrcpp::refine_grid(FunctionTree<D> &out, double prec, bool absPrec)
```

Refine the grid of a MW function representation.

This will first perform a split check on the existing leaf nodes in the tree based on the provided precision parameter, then it will compute scaling coefs of the new nodes, thus leaving the function representation unchanged, but (possibly) on a larger grid.

Return The number of nodes that were split

### **Parameters**

- [inout] out: Output tree to be refined
- [in] prec: Precision for initial split check
- [in] absPrec: Build output tree based on absolute precision

## template<int D>

```
int mrcpp::refine_grid(FunctionTree<D> &out, FunctionTree<D> &inp)
```

Refine the grid of a MW function representation.

This will first perform a split check on the existing leaf nodes in the output tree based on the structure of the input tree (same as build\_grid), then it will compute scaling coefs of the new nodes, thus leaving the function representation unchanged, but on a larger grid.

**Return** The number of nodes that were split

## **Parameters**

- [inout] out: Output tree to be refined
- [in] inp: Input tree that defines the new grid

## 8.2.4 File I/O

```
void mrcpp::FunctionTree::saveTree(const std::string &file) override
Write the tree structure to disk, for later use.
```

#### **Parameters**

• [in] file: File name, will get ".tree" extension

```
void mrcpp::FunctionTree::loadTree(const std::string &file) override
Read a previously stored tree structure from disk.
```

**Note** This tree must have the exact same MRA the one that was saved

#### **Parameters**

• [in] file: File name, will get ".tree" extension

## 8.2.5 Extracting data

Given a FunctionTree that is a well defined function representation, the following data can be extracted:

```
double mrcpp::FunctionTree::integrate() const
```

Return Integral of the function over the entire computational domain

```
double \ \texttt{mrcpp::} \textit{FunctionTree::} \textbf{evalf} \ (\textbf{const Coord} < D > \& r) \ \textbf{const override}
```

Return Function value in a point, out of bounds returns zero

**Note** This will only evaluate the *scaling* part of the leaf nodes in the tree, which means that the function values will not be fully accurate. If you want to include also the *final* wavelet part you'll have to manually extend the MW grid by one level before evaluating, using mrcpp::refine\_grid(tree, 1) This is done to allow a fast and const function evaluation that can be done in OMP parallel.

### **Parameters**

• [in] r: Cartesian coordinate

template<int D>

double mrcpp::dot (FunctionTree<D> &bra, FunctionTree<D> &ket)

The dot product is computed with the trees in compressed form, i.e. scaling coefs only on root nodes, wavelet coefs on all nodes. Since wavelet functions are orthonormal through ALL scales and the root scaling functions are orthonormal to all finer level wavelet functions, this becomes a rather efficient procedure as you only need to compute the dot product where the grids overlap.

**Return** Dot product <br/>
bralket> of two MW function representations

### **Parameters**

- [in] bra: Bra side input function
- [in] ket: Ket side input function

## 8.3 FunctionTreeVector

The FunctionTreeVector is simply an alias for a std::vector of std::tuple containing a numerical coefficient and a FunctionTree pointer.

```
template<int D>
```

void mrcpp::clear (FunctionTreeVector<*D*> &fs, bool dealloc = false)
Remove all entries in the vector.

### **Parameters**

- [in] fs: Vector to clear
- [in] dealloc: Option to free *FunctionTree* pointer before clearing

## template<int D>

```
double mrcpp::get_coef (const FunctionTreeVectorD> &fs, int i)
```

**Return** Numerical coefficient at given position in vector

## **Parameters**

- [in] fs: Vector to fetch from
- [in] i: Position in vector

## template<int D>

```
FunctionTree < D > \&mrcpp::get_func (FunctionTree Vector < D > \&fs, int i)
```

**Return** FunctionTree at given position in vector

## **Parameters**

```
• [in] fs: Vector to fetch from
```

• [in] i: Position in vector

## template<int D>

```
int mrcpp::get_n_nodes(const FunctionTreeVector<D>&fs)
```

**Return** Total number of nodes of all trees in the vector

#### **Parameters**

• [in] fs: Vector to fetch from

## template<int D>

```
int mrcpp::get_size_nodes (const FunctionTreeVector<D> &fs)
```

**Return** Total size of all trees in the vector, in kB

#### **Parameters**

• [in] fs: Vector to fetch from

## 8.4 Examples

## 8.4.1 Constructing an MRA

An MRA is defined in two steps, first the computational domain is given by a BoundingBox (D is the dimension), e.g. for a total domain of  $[-32, 32]^3$  in three dimensions (eight root boxes of size  $[16]^3$  each):

which is combined with a ScalingBasis to give an MRA, e.g. interpolating scaling functions of order k = 9:

Two types of ScalingBasis are supported (LegendreBasis and InterpolatingBasis), and they are both available at orders k = 1, 2, ..., 40 (note that some operators are constructed using intermediates of order 2k, so in that case the maximum order available is k = 20).

## 8.4.2 Working withFunctionTreeVectors

Elements can be appended to the vector using the std::make\_tuple, elements are obtained with the get\_func and get\_coef functions:

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## 8.4.3 Building empty grids

Sometimes it is useful to construct an empty grid based on some available information of the function that is about to be represented. This can be e.g. that you want to copy the grid of an existing FunctionTree or that an analytic function has more or less known grid requirements (like Gaussians). Sometimes it is even necessary to force the grid refinement beyond the coarsest scales in order for the adaptive refining algorithm to detect a wavelet "signal" that allows it to do its job properly (this happens for narrow Gaussians where none of the initial quadrature points hits a function value significantly different from zero).

The simplest way to build an empty grid is to copy the grid from an existing tree (assume that f\_tree has been properly built so that it contains more than just root nodes)

```
mrcpp::FunctionTree<D> f_tree(MRA);  // Input tree
mrcpp::FunctionTree<D> g_tree(MRA);  // Output tree

mrcpp::project(prec, f_tree, f_func); // Build adaptive grid for f_tree
mrcpp::copy_grid(g_tree, f_tree);  // Copy grid from f_tree to g_tree
```

Passing an analytic function as argument to the generator will build a grid based on some predefined information of the function (if there is any, otherwise it will do nothing)

```
mrcpp::RepresentableFunction<D> func; // Analytic function
mrcpp::FunctionTree<D> tree(MRA); // Output tree
mrcpp::build_grid(tree, func); // Build grid based on f_func
```

The lambda analytic functions do *not* provide such information, this must be explicitly implemented as a RepresentableFunction sub-class (see MRCPP programmer's guide for details).

Actually, the effect of the build\_grid is to *extend* the existing grid with any missing nodes relative to the input. There is also a version of build\_grid taking a FunctionTree argument. Its effect is very similar to the copy\_grid above, with the only difference that now the output grid is *extended* with the missing nodes (e.i. the nodes that are already there are *not* removed first). This means that we can build the union of two grids by successive applications of build\_grid

In contrast, doing the same with <code>copy\_grid</code> would clear the <code>f\_tree</code> grid in between, and you would <code>only</code> get a (identical) copy of the last <code>h\_tree</code> grid, with no memory of the <code>g\_tree</code> grid that was once there. One can also make the grids of two functions equal to their union

The union grid of several trees can be constructed in one go using a FunctionTreeVector

```
mrcpp::FunctionTreeVector<D> inp_vec;
inp_vec.push_back(std::make_tuple(1.0, tree_1));
inp_vec.push_back(std::make_tuple(1.0, tree_2));
```

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# 8.4.4 Projection

The project function takes an analytic D-dimensional scalar function (which can be defined as a lambda function or one of the explicitly implemented sub-classes of the RepresentableFunction base class in MRCPP) and projects it with the given precision onto the MRA defined by the FunctionTree. E.g. a unit charge Gaussian is projected in the following way (the MRA must be initialized as above)

```
// Defining an analytic function
double beta = 10.0;
double alpha = std::pow(beta/pi, 3.0/2.0);
auto func = [alpha, beta] (const mrcpp::Coord<3> &r) -> double {
    double R = std::sqrt(r[0]*r[0] + r[1]*r[1] + r[2]*r[2]);
    return alpha*std::exp(-beta*R*R);
};

double prec = 1.0e-5;
mrcpp::FunctionTree<3> tree(MRA);
mrcpp::project(prec, tree, func);
```

This projection will start at the default initial grid (only the root nodes of the given MRA), and adaptively build the full grid. Alternatively, the grid can be estimated *a priori* if the analytical function has some known features, such as for Gaussians:

This will first produce an empty grid suited for representing the analytic function func (this is meant as a way to make sure that the projection starts on a grid where the function is actually visible, as for very narrow Gaussians, it's not meant to be a good approximation of the final grid) and then perform the projection on the given numerical grid. With a negative prec (or max\_iter = 0) the projection will be performed strictly on the given initial grid, with no further refinements.

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## 8.4.5 Addition

Arithmetic operations in the MW representation are performed using the FunctionTreeVector, and the general sum  $f = \sum_i c_i f_i(x)$  is done in the following way

```
double a, b, c;
                                                   // Addition parameters
                                                   // Input function
mrcpp::FunctionTree<D> a_tree(MRA);
mrcpp::FunctionTree<D> b_tree(MRA);
                                                   // Input function
mrcpp::FunctionTree<D> c_tree(MRA);
                                                   // Input function
mrcpp::FunctionTreeVector<D> inp vec;
                                                  // Vector to hold input functions
                                                  // Append to vector
inp_vec.push_back(std::make_tuple(a, &a_tree));
                                                  // Append to vector
inp_vec.push_back(std::make_tuple(b, &b_tree));
inp_vec.push_back(std::make_tuple(c, &c_tree));
                                                  // Append to vector
mrcpp::FunctionTree<D> f_tree(MRA);
                                                  // Output function
                                                   // Adaptive addition
mrcpp::add(prec, f_tree, inp_vec);
```

The default initial grid is again only the root nodes, and a positive prec is required to build an adaptive tree structure for the result. The special case of adding two functions can be done directly without initializing a FunctionTreeVector

```
mrcpp::FunctionTree<D> f_tree(MRA);
mrcpp::add(prec, f_tree, a, a_tree, b, b_tree);
```

Addition of two functions is usually done on their (fixed) union grid

Note that in the case of addition there is no extra information to be gained by going beyond the finest refinement levels of the input functions, so the union grid summation is simply the best you can do, and adding a positive prec will not make a difference. There are situations where you want to use a *smaller* grid, though, e.g. when performing a unitary transformation among a set of FunctionTrees. In this case you usually don't want to construct *all* the output functions on the union grid of *all* the input functions, and this can be done by adding the functions adaptively starting from root nodes.

If you have a summation over several functions but want to perform the addition on the grid given by the *first* input function, you first copy the wanted grid and then perform the operation on that grid

Here you can of course also add a positive prec to the addition and the resulting function will be built adaptively starting from the given initial grid.

## 8.4.6 Multiplication

The multiplication follows the exact same syntax as the addition, where the product  $f = \prod_i c_i f_i(x)$  is done in the following way

```
double a, b, c;
                                                   // Multiplication parameters
                                                  // Input function
mrcpp::FunctionTree<D> a_tree(MRA);
mrcpp::FunctionTree<D> b_tree(MRA);
                                                   // Input function
mrcpp::FunctionTree<D> c_tree (MRA);
                                                   // Input function
mrcpp::FunctionTreeVector<D> inp vec;
                                                  // Vector to hold input functions
inp_vec.push_back(std::make_tuple(a, &a_tree));
                                                  // Append to vector
inp_vec.push_back(std::make_tuple(b, &b_tree));
                                                  // Append to vector
inp_vec.push_back(std::make_tuple(c, &c_tree));
                                                  // Append to vector
mrcpp::FunctionTree<D> f_tree (MRA);
                                                  // Output function
mrcpp::multipy(prec, f_tree, inp_vec);
                                                  // Adaptive multiplication
```

In the special case of multiplying two functions the coefficients are collected into one argument

```
mrcpp::FunctionTree<D> f_tree(MRA);
mrcpp::multiply(prec, f_tree, a*b, a_tree, b_tree);
```

For multiplications, there might be a loss of accuracy if the product is restricted to the union grid. The reason for this is that the product will contain signals of higher frequency than each of the input functions, which require a higher grid refinement for accurate representation. By specifying a positive prec you will allow the grid to adapt to the higher frequencies, but it is usually a good idea to restrict to one extra refinement level beyond the union grid (by setting max\_iter=1) as the grids are not guaranteed to converge for such local operations (like arithmetics, derivatives and function mappings)

# 8.4.7 Re-using grids

Given a FunctionTree that is a valid function representation, we can clear its MW expansion coefficients as well as its grid refinement

This action will leave the FunctionTree in the same state as after construction (undefined function, only root nodes), and its coefficients can now be re-computed.

In certain situations it might be desireable to separate the actions of computing MW coefficients and refining the grid. For this we can use the refine\_grid, which will adaptively refine the grid one level (based on the wavelet norm and the given precision) and project the existing function representation onto the new finer grid

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```
mrcpp::refine_grid(tree, prec);
```

E.i., this will *not* change the function that is represented in tree, but it *might* increase its grid size. The same effect can be made using another FunctionTree argument instead of the precision parameter

```
mrcpp::refine_grid(tree_out, tree_in);
```

which will extend the grid of tree\_out in the same way as build\_grid as shown above, but it will keep the function representation in tree\_out.

This functionality can be combined with <code>clear\_grid</code> to make a "manual" adaptive building algorithm. One example where this might be useful is in iterative algorithms where you want to fix the grid size for all calculations within one cycle and then relax the grid in the end in preparation for the next iteration. The following is equivalent to the adaptive projection above (<code>refine\_grid</code> returns the number of new nodes that were created in the process)

**CHAPTER** 

NINE

## **MWOPERATORS**

The MW operators discussed in this chapter is available to the application program by including:

#include "MRCPP/MWOperators"

# 9.1 ConvolutionOperator

**Note:** The convolution operators have separate precision parameters for their construction and application. The build\_prec argument to the operator constructors will affect e.g. the number of terms in the separated representations of the Poisson/Helmholtz approximations, as well as the operator bandwidth. The apply\_prec argument to the apply function relates only to the adaptive construction of the output function, based on a wavelet norm error estimate.

## template<int D>

The identity kernel (Dirac's delta function) is approximated by a narrow Gaussian function:  $I(r-r') = \delta(r-r') \approx \alpha e^{-\beta(r-r')^2}$ 

### **Public Functions**

**IdentityConvolution** (const MultiResolutionAnalysis < D > & mra, double pr)

This will project a kernel of a single gaussian with exponent sqrt(10/build\_prec).

Return New IdentityConvolution object

### **Parameters**

- [in] mra: Which MRA the operator is defined
- [in] pr: Build precision, closeness to delta function

## template<int **D**>

class mrcpp::DerivativeConvolution : public mrcpp::ConvolutionOperator<D>
 Convolution with a derivative kernel.

The derivative kernel (derivative of Dirac's delta function) is approximated by the derivative of a narrow Gaussian function:  $D^x(r-r') = \frac{d}{dx}\delta(r-r') \approx \frac{d}{dx}\alpha e^{-\beta(r-r')^2}$ 

### **Public Functions**

**DerivativeConvolution** (const *MultiResolutionAnalysis<D> &mra*, double *pr*)

This will project a kernel of a single differentiated gaussian with exponent sqrt(10/build\_prec).

Return New Derivative Convolution object

#### **Parameters**

- [in] mra: Which MRA the operator is defined
- [in] pr: Build precision, closeness to delta function

class mrcpp::PoissonOperator: public mrcpp::ConvolutionOperator<3>

Convolution with the Poisson Green's function kernel.

The Poisson kernel is approximated as a sum of Gaussian functions in order to allow for separated application of the operator in the Cartesian directions:  $P(r-r') = \frac{1}{|r-r'|} \approx \sum_{m}^{M} \alpha_m e^{-\beta_m (r-r')^2}$ 

### **Public Functions**

PoissonOperator (const MultiResolutionAnalysis < 3 > &mra, double pr = -1.0)

This will construct a gaussian expansion to approximate 1/r, and project each term into a one-dimensional MW operator. Subsequent application of this operator will apply each of the terms to the input function in all Cartesian directions.

Return New PoissonOperator object

### **Parameters**

- [in] mra: Which MRA the operator is defined
- [in] pr: Build precision, closeness to 1/r

class mrcpp::HelmholtzOperator: public mrcpp::ConvolutionOperator<3>

Convolution with the Helmholtz Green's function kernel.

The Helmholtz kernel is approximated as a sum of gaussian functions in order to allow for separated application of the operator in the Cartesian directions:  $H(r-r')=\frac{e^{-\mu|r-r'|}}{|r-r'|}\approx \sum_m^M \alpha_m e^{-\beta_m(r-r')^2}$ 

### **Public Functions**

HelmholtzOperator (const MultiResolutionAnalysis < 3 > & mra, double m, double pr = -1.0)

This will construct a gaussian expansion to approximate exp(-mu\*r)/r, and project each term into a one-dimensional MW operator. Subsequent application of this operator will apply each of the terms to the input function in all Cartesian directions.

Return New HelmholtzOperator object

### **Parameters**

- [in] mra: Which MRA the operator is defined
- [in] m: Exponential parameter of the operator
- [in] pr: Build precision, closeness to exp(-mu\*r)/r

template<int D>

void mrcpp::apply (double prec, FunctionTree<D> &out, ConvolutionOperator<D> &oper, FunctionTree<D> &inp, int maxIter, bool absPrec)

Application of MW integral convolution operator.

The output function will be computed using the general algorithm:

- Compute MW coefs on current grid
- Refine grid where necessary based on prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

### **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] oper: Convolution operator to apply
- [in] inp: Input function
- [in] maxIter: Maximum number of refinement iterations in output tree, default -1
- [in] absPrec: Build output tree based on absolute precision, default false

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

### template<int D>

void mrcpp::apply (double prec, FunctionTree<D> &out, ConvolutionOperator<D> &oper, FunctionTree<D> &inp, FunctionTree Vector<D> &precTrees, int maxIter, bool absPrec)
Application of MW integral convolution operator.

The output function will be computed using the general algorithm:

- Compute MW coefs on current grid
- Refine grid where necessary based on scaled prec
- Repeat until convergence or maxIter is reached
- prec < 0 or maxIter = 0 means NO refinement
- maxIter < 0 means no bound

### **Parameters**

- [in] prec: Build precision of output function
- [out] out: Output function to be built
- [in] oper: Convolution operator to apply
- [in] inp: Input function
- [in] precTrees: Precision trees
- [in] maxIter: Maximum number of refinement iterations in output tree, default -1
- [in] absPrec: Build output tree based on absolute precision, default false

The precision will be scaled locally by the maxNorms of the precTrees input vector.

**Note** This algorithm will start at whatever grid is present in the out tree when the function is called (this grid should however be EMPTY, e.i. no coefs).

# 9.2 DerivativeOperators

**Note:** The derivative operators have clearly defined requirements on the output grid structure, based on the grid of the input function. This means that there is no real grid adaptivity, and thus no precision parameter is needed for the application of such an operator.

### template<int D>

**class** mrcpp::ABGVOperator: public mrcpp::DerivativeOperator<*D*> Derivative operator as defined by Alpert, Beylkin, Ginez and Vozovoi, J Comp Phys 182, 149-190 (2002).

#### **Public Functions**

**ABGVOperator** (const *MultiResolutionAnalysis<D> &mra*, double a, double b)

Boundary parameters correspond to:

- a=0.0 b=0.0: Strictly local "center" difference
- a=0.5 b=0.5: Semi-local central difference
- a=1.0 b=0.0: Semi-local forward difference
- a=0.0 b=1.0: Semi-local backward difference

Return New ABGVOperator object

### **Parameters**

- [in] mra: Which MRA the operator is defined
- [in] a: Left boundary condition
- [in] b: Right boundary condition

## template<int D>

**class** mrcpp::PHOperator: public mrcpp::DerivativeOperator<*D*>

Derivative operator based on the smoothing derivative of Pavel Holoborodko.

### **Public Functions**

PHOperator (const MultiResolutionAnalysis<D> &mra, int order)

Return New PHOperator object

### **Parameters**

- [in] mra: Which MRA the operator is defined
- [in] order: Derivative order, defined for 1 and 2

## template<int D>

```
class mrcpp::BSOperator:public mrcpp::DerivativeOperator<D>
```

B-spline derivative operator as defined by Anderson et al, J Comp Phys X 4, 100033 (2019).

### **Public Functions**

**BSOperator** (const *MultiResolutionAnalysis<D> &mra*, int *order*)

**Return** New *BSOperator* object

### **Parameters**

- [in] mra: Which MRA the operator is defined
- [in] order: Derivative order, defined for 1, 2 and 3

### template<int D>

```
void mrcpp::apply (FunctionTree<D> &out, DerivativeOperator<D> &oper, FunctionTree<D> &inp, int
      \frac{dir}{dr} Application of MW derivative operator.
```

The output function will be computed on a FIXED grid that is predetermined by the type of derivative operator. For a strictly local operator (ABGV 00), the grid is an exact copy of the input function. For operators that involve also neighboring nodes (ABGV\_55, PH, BS) the base grid will be WIDENED by one node in the direction of application (on each side).

### **Parameters**

- [out] out: Output function to be built
- [in] oper: Derivative operator to apply
- [in] inp: Input function
- [in] dir: Direction of derivative

**Note** The output function should contain only empty root nodes at entry.

### template<int D>

```
void mrcpp::divergence (FunctionTree<D> &out, DerivativeOperator<D> &oper, FunctionTreeVec-
                           tor < D > \&inp)
```

Calculation of divergence of a function vector.

The derivative operator is applied in each Cartesian direction to the corresponding components of the input vector and added up to the final output. The grid of the output is fixed as the union of the component grids (including any derivative widening, see derivative apply).

### **Parameters**

- [out] out: Output function
- [in] oper: Derivative operator to apply
- [in] inp: Input function vector

## Note

- The length of the input vector must be the same as the template dimension D.
- The output function should contain only empty root nodes at entry.

### template<int D>

```
FunctionTreeVector<D> mrcpp::gradient (DerivativeOperator<D> & oper, FunctionTree<D> & inp)
     Calculation of gradient vector of a function.
```

The derivative operator is applied in each Cartesian direction to the input function and appended to the output vector.

**Return** FunctionTreeVector containing the gradient

#### **Parameters**

- [in] oper: Derivative operator to apply
- [in] inp: Input function

**Note** The length of the output vector will be the template dimension D.

# 9.3 Examples

## 9.3.1 PoissonOperator

The electrostatic potential g arising from a charge distribution f are related through the Poisson equation

$$-\nabla^2 g(r) = f(r)$$

This equation can be solved with respect to the potential by inverting the differential operator into the Green's function integral convolution operator

$$g(r) = \int \frac{1}{4\pi \|r - r'\|} f(r') dr'$$

This operator is available in the MW representation, and can be solved with arbitrary (finite) precision in linear complexity with respect to system size. Given an arbitrary charge dirtribution f\_tree in the MW representation, the potential is computed in the following way:

```
double apply_prec;
double build_prec;

mrcpp::PoissonOperator P(MRA, build_prec);

→ operator

mrcpp::FunctionTree<3> f_tree(MRA);

mrcpp::FunctionTree<3> g_tree(MRA);

mrcpp::apply(apply_prec, g_tree, P, f_tree);

// Precision for operator application

// MW representation of Poisson_

// Input function

// Output function

// Apply operator adaptively
```

The Coulomb self-interaction energy can now be computed as the dot product:

```
double E = mrcpp::dot(g_tree, f_tree);
```

# 9.3.2 HelmholtzOperator

The Helmholtz operator is a generalization of the Poisson operator and is given as the integral convolution

$$g(r) = \int \frac{e^{-\mu \|r - r'\|}}{4\pi \|r - r'\|} f(r') dr'$$

The operator is the inverse of the shifted Laplacian

$$\left[ -\nabla^2 + \mu^2 \right] g(r) = f(r)$$

and appears e.g. when solving the SCF equations. The construction and application is similar to the Poisson operator, with an extra argument for the  $\mu$  parameter

## 9.3.3 ABGVOperator

The ABGV (Alpert, Beylkin, Gines, Vozovoi) derivative operator is initialized with two parameters a and b accounting for the boundary conditions between adjacent nodes, see Alpert et al.

```
double a = 0.0, b = 0.0;
                                                // Boundary conditions for operator
mrcpp::ABGVOperator<3> D(MRA, a, b);
                                                // MW derivative operator
mrcpp::FunctionTree<3> f(MRA);
                                                // Input function
mrcpp::FunctionTree<3> f_x (MRA);
                                               // Output function
mrcpp::FunctionTree<3> f_y (MRA);
                                               // Output function
mrcpp::FunctionTree<3> f_z(MRA);
                                               // Output function
                                               // Operator application in x direction
mrcpp::apply(f_x, D, f, 0);
mrcpp::apply(f_y, D, f, 1);
                                                // Operator application in y direction
mrcpp::apply(f_z, D, f, 2);
                                                // Operator application in z direction
```

The tree structure of the output function will depend on the choice of parameters a and b: if both are zero, the output grid will be identical to the input grid; otherwise the grid will be widened by one node (on each side) in the direction of application.

# 9.3.4 PHOperator

The PH derivative operator is based on the noise reducing derivative of Pavel Holoborodko. This operator is also available as a direct second derivative.

Special thanks to Prof. Robert J. Harrison (Stony Brook University) for sharing the operator coefficients.

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## 9.3.5 BSOperator

The BS derivative operator is based on a pre-projection onto B-splines in order to remove the discontinuities in the MW basis, see Anderson et al. This operator is also available as a direct second and third derivative.

```
mrcpp::BSOperator<3> D1(MRA, 1);
                                                // MW 1st derivative operator
mrcpp::BSOperator<3> D2(MRA, 2);
                                                // MW 2nd derivative operator
mrcpp::BSOperator<3> D3(MRA, 3);
                                                // MW 3nd derivative operator
                                                // Input function
mrcpp::FunctionTree<3> f(MRA);
mrcpp::FunctionTree<3> f_x(MRA);
                                                // Output function
mrcpp::FunctionTree<3> f_yy (MRA);
                                                // Output function
mrcpp::FunctionTree<3> f_zzz(MRA);
                                                // Output function
                                                // Operator application in x direction
mrcpp::apply(f_x, D1, f, 0);
                                                // Operator application in x direction
mrcpp::apply(f_yy, D2, f, 1);
mrcpp::apply(f_zzz, D3, f, 2);
                                                // Operator application in x direction
```

**CHAPTER** 

**TEN** 

# **GAUSSIANS**

MRCPP provides some simple features for analytic Gaussian functions. These are meant to be used as a starting point for MW computations, they are *not* meant for heavy analytical computation, like GTO basis sets. The Gaussian features are available by including:

#include "MRCPP/Gaussians"

## template<int D>

class mrcpp::GaussFunc:public mrcpp::Gaussian<D>

Gaussian function in D dimensions with a simple monomial in front.

• Monodimensional Gaussian (GaussFunc<1>):

$$g(x) = \alpha(x - x_0)^a e^{-\beta(x - x_0)^2}$$

• Multidimensional Gaussian (GaussFunc<D>):

$$G(x) = \prod_{d=1}^{D} g^d(x^d)$$

## **Public Functions**

**GaussFunc** (double *beta*, double *alpha*, **const** CoordD> &pos = {}, **const** std::arrayint, D> &pow = {})

Return New GaussFunc object

### **Parameters**

- [in] beta: Exponent,  $e^{-\beta r^2}$
- [in] alpha: Coefficient,  $\alpha e^{-r^2}$
- [in] pos: Position  $(x pos[0]), (y pos[1]), \dots$
- [in] pow: Monomial power,  $x^{pow[0]}, y^{pow[1]}, \dots$

double calcCoulombEnergy (GaussFunc<D> &rhs)

Compute Coulomb repulsion energy between two GaussFuncs.

Return Coulomb energy

**Note** Both Gaussians must be normalized to unit charge  $\alpha = (\beta/\pi)^{D/2}$  for this to be correct!

### **Parameters**

- [in] this: Left hand GaussFunc
- [in] rhs: Right hand GaussFunc

# double calcOverlap (GaussFunc<D> &rhs) override

Compute analytic overlap between two Gaussians.

## Return Overlap integral

### **Parameters**

- [in] this: Left hand Gaussian
- [in] rhs: Right hand Gaussian

## double calcOverlap (GaussPoly<D> &rhs) override

Compute analytic overlap between two Gaussians.

## Return Overlap integral

### **Parameters**

- [in] this: Left hand Gaussian
- [in] rhs: Right hand Gaussian

### GaussPoly<D> differentiate (int dir) override

Compute analytic derivative of Gaussian.

## Return New GaussPoly

### **Parameters**

• [in] dir: Cartesian direction of derivative

## GaussPoly<D> mult (const GaussFunc<D> &rhs)

Multiply two GaussFuncs.

### Return New GaussPoly

### **Parameters**

- [in] this: Left hand side of multiply
- [in] rhs: Right hand side of multiply

## GaussFunc<D> mult (double c)

Multiply GaussFunc by scalar.

### Return New GaussFunc

### **Parameters**

• [in] c: Scalar to multiply

# double evalf(const Coord < D > & r) const

**Return** Function value in a point

### **Parameters**

• [in] r: Cartesian coordinate

## double getSquareNorm()

**Return** Squared L2 norm of function  $||f||^2$ 

### void normalize()

Rescale function by its norm  $||f||^{-1}$ .

### template<int D>

## class mrcpp::GaussPoly:public mrcpp::Gaussian<D>

Gaussian function in D dimensions with a general polynomial in front.

• Monodimensional Gaussian (GaussPoly<1>):

$$g(x) = \alpha P(x - x_0)e^{-\beta(x - x_0)^2}$$

• Multidimensional Gaussian (GaussFunc<D>):

$$G(x) = \prod_{d=1}^{D} g^d(x^d)$$

### **Public Functions**

GaussPoly(double alpha = 0.0, double coef = 1.0, const Coord<D> &pos = {}, const
std::array<int, D> &power = {})

Return New GaussPoly object

### **Parameters**

- [in] beta: Exponent,  $e^{-\beta r^2}$
- [in] alpha: Coefficient,  $\alpha e^{-r^2}$
- [in] pos: Position (x pos[0]), (y pos[1]), ...
- [in] pow: Max polynomial degree,  $P_0(x), P_1(y), ...$

## double calcOverlap (GaussFunc < D > &b) override

Compute analytic overlap between two Gaussians.

**Return** Overlap integral

### **Parameters**

- [in] this: Left hand Gaussian
- [in] rhs: Right hand Gaussian

## double calcOverlap (GaussPoly < D > &b) override

Compute analytic overlap between two Gaussians.

**Return** Overlap integral

### **Parameters**

- [in] this: Left hand Gaussian
- [in] rhs: Right hand Gaussian

### GaussPoly differentiate (int dir) override

Compute analytic derivative of Gaussian.

Return New GaussPoly

#### **Parameters**

• [in] dir: Cartesian direction of derivative

GaussPoly<D> mult (double c)

Multiply GaussPoly by scalar.

Return New GaussPoly

### **Parameters**

• [in] c: Scalar to multiply

void **setPoly** (int *d*, Polynomial &poly)

Set polynomial in given dimension.

### **Parameters**

- [in] d: Cartesian direction
- [in] poly: Polynomial to set

double evalf (const Coord<*D*> &r) const

**Return** Function value in a point

#### **Parameters**

• [in] r: Cartesian coordinate

double getSquareNorm()

**Return** Squared L2 norm of function  $||f||^2$ 

void normalize()

Rescale function by its norm  $||f||^{-1}$ .

template<int D>

class mrcpp::GaussExp:public mrcpp::RepresentableFunction<D>

Gaussian expansion in D dimensions.

• Monodimensional Gaussian expansion:

$$g(x) = \sum_{m=1}^{M} g_m(x) = \sum_{m=1}^{M} \alpha_m e^{-\beta(x-x^0)^2}$$

• Multidimensional Gaussian expansion:

$$G(x) = \sum_{m=1}^{M} G_m(x) = \sum_{m=1}^{M} \prod_{d=1}^{D} g_m^d(x^d)$$

### **Public Functions**

double calcCoulombEnergy()

**Return** Coulomb repulsion energy between all pairs in *GaussExp*, including self-interaction

**Note** Each Gaussian must be normalized to unit charge  $c = (\alpha/\pi)^{D/2}$  for this to be correct!

double evalf (const Coord<D> &r) const override

**Return** Function value in a point

### **Parameters**

• [in] r: Cartesian coordinate

void append (const Gaussian D > &g)

Append Gaussian to expansion.

void append (const GaussExp<D>&g)
Append GaussExp to expansion.

# 10.1 Examples

A GaussFunc is a simple D-dimensional Gaussian function with a Cartesian monomial in front, e.g. in 3D:

$$f(r) = \alpha (x - x_0)^a (y - y_0)^b (z - z_0)^c e^{-\beta \|r - r_0\|^2}$$

```
double alpha, beta;
std::array<int, 3> pow = {a, b, c};
mrcpp::Coord<3> pos = {x_0, y_0, z_0};
mrcpp::GaussFunc<3> gauss(beta, alpha, pos, pow);

double E = gauss.calcCoulombEnergy(gauss); // Analytical energy
```

This Gaussian function can be used to build an empty grid based on the position and exponent. The grid will then be refined close to the center of the Gaussian, with deeper refinement for higher exponents (steeper function):

GaussPoly is a generalization of the GaussFunc, where there is an arbitrary polynomial in front of the exponential

$$f(r) = \alpha P(r - r_0)e^{-\beta \|r - r_0\|^2}$$

For instance, the following function can be constructed

$$f(r) = \alpha(a_x + b_x x + c_x x^2)(a_y + b_y y + c_y y^2)(a_z + b_z z + c_z z^2)e^{-\beta \|r - r_0\|^2}$$

```
auto gauss_poly = GaussPoly<D>(beta, alpha, pos, pow);

// Create polynomial in x, y and z direction
auto pol_x = Polynomial(2); // 2 is the degree of the polynomial
pol_x.getCoefs() << a_x, b_x, c_x;
auto pol_y = Polynomial(2);
pol_y.getCoefs() << a_y, b_y, c_y;
auto pol_z = Polynomial(2);
pol_z.getCoefs() << a_z, b_z, c_z;

// Add polynomials to gauss_poly
guass_poly.setPoly(0, pol_x);
guass_poly.setPoly(1, pol_y);
guass_poly.setPoly(2, pol_z);</pre>
```

A Gaussexp is a collection of Gaussians in the form

$$G(r) = \sum_{i} c_i g_i(r)$$

where  $g_i$  can be either GaussFunc or GaussPoly

$$g_i(r) = \alpha_i P_i(r - r_i) e^{-\beta_i \|r - r_i\|^2}$$

Individual Gaussian functions can be appended to the GaussExp and treated as a single function:

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**CHAPTER** 

## **ELEVEN**

## PARALLEL

The core features of MRCPP are parallelized using a shared memory model *only* (OpenMP). This means that there is *no intrinsic* MPI parallelization (e.i. no data distribution across machines) *within* the library routines. However, the code comes with a small set of features that facilitate MPI work and data distribution in the host program, in the sense that *entire* FunctionTree objects can be located on different machines and communicated between them. Also, a FunctionTree can be *shared* between several MPI processes that are located on the *same* machine. This means that several processes have read access to the same FunctionTree, thus reducing the memory footprint, as well as the need for communication.

The MPI features are available by including:

```
#include "MRCPP/Parallel"
```

# 11.1 The host program

In order to utilize the MPI features of MRCPP, the MPI instance must be initialized (and finalized) by the host program, as usual:

```
MPI_Init(&argc, &argv);
int size, rank;
MPI_Comm_size(MPI_COMM_WORLD, &size); // Get MPI world size
MPI_Comm_rank(MPI_COMM_WORLD, &rank); // Get MPI world rank
MPI_Finalize();
```

For the shared memory features we must make sure that the ranks within a communicator is actually located on the same machine. When running on distributed architectures this can be achieved by creating separate communicators for each physical machine, e.g. to split MPI\_COMM\_WORLD into a new communicator group called MPI\_COMM\_SHARED that share the same physical memory space:

```
// Initialize a new communicator called MPI_COMM_SHARE
MPI_Comm MPI_COMM_SHARE;

// Split MPI_COMM_WORLD into sub groups and assign to MPI_COMM_SHARE
MPI_Comm_split_type(MPI_COMM_WORLD, MPI_COMM_TYPE_SHARED, 0, MPI_INFO_NULL, &MPI_COMM_
SHARE);
```

Note that the main purpose of the shared memory feature of MRCPP is to avoid memory duplication and reduce the memory footprint, it will **not** automatically provide any work sharing parallelization for the construction of the shared FunctionTree.

# 11.2 Blocking communication

### template<int D>

void mrcpp::send\_tree (FunctionTree < D > &tree, int dst, int tag, mrcpp::mpi\_comm comm, int nChunks)
Send FunctionTree to a given MPI rank using blocking communication.

The number of memory chunks must be known before we can send the tree. This can be specified in the last argument if known a priori, in order to speed up communication, otherwise it will be communicated in a separate step before the main communication.

#### **Parameters**

- [in] tree: FunctionTree to send
- [in] dst: MPI rank to send to
- [in] tag: unique identifier
- [in] comm: Communicator that defines ranks
- [in] nChunks: Number of memory chunks to send

### template<int D>

void mrcpp::recv\_tree (FunctionTree<D> &tree, int src, int tag, mrcpp::mpi\_comm comm, int nChunks)
Receive FunctionTree from a given MPI rank using blocking communication.

The number of memory chunks must be known before we can receive the tree. This can be specified in the last argument if known a priori, in order to speed up communication, otherwise it will be communicated in a separate step before the main communication.

### **Parameters**

- [in] tree: FunctionTree to write into
- [in] src: MPI rank to receive from
- [in] tag: unique identifier
- [in] comm: Communicator that defines ranks
- [in] nChunks: Number of memory chunks to receive

## **11.2.1 Example**

A blocking send/receive means that the function call does not return until the communication is completed. This is a simple and safe option, but can lead to significant overhead if the communicating MPI processes are not synchronized.

```
mrcpp::FunctionTree<3> tree(MRA);

// At this point tree is uninitialized on both rank 0 and 1

// Only rank 0 projects the function
if (rank == 0) mrcpp::project(prec, tree, func);

// At this point tree is projected on rank 0 but still uninitialized on rank 1
```

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```
// Sending tree from rank 0 to rank 1
int tag = 111111; // Unique tag for each communication
int src=0, dst=1; // Source and destination ranks
if (rank == src) mrcpp::send_tree(tree, dst, tag, MPI_COMM_WORLD);
if (rank == dst) mrcpp::revc_tree(tree, src, tag, MPI_COMM_WORLD);

// At this point tree is projected on both rank 0 and 1

// Rank 0 clear the tree
if (rank == 0) mrcpp::clear(tree);

// At this point tree is uninitialized on rank 0 but still projected on rank 1
```

# 11.3 Shared memory

### class mrcpp::SharedMemory

Shared memory block within a compute node.

This class defines a shared memory window in a shared MPI communicator. In order to allocate a *FunctionTree* in shared memory, simply pass a *SharedMemory* object to the *FunctionTree* constructor.

### **Public Functions**

**SharedMemory** (mrcpp::mpi\_comm *comm*, int *sh\_size*) *SharedMemory* constructor.

### **Parameters**

- [in] comm: Communicator sharing resources
- [in] sh\_size: Memory size, in MB

## template<int D>

void mrcpp::share\_tree (FunctionTree < D > & tree, int src, int tag, mrcpp::mpi\_comm comm)

Share a FunctionTree among MPI processes that share the same physical memory.

This function should be called every time a shared function is updated, in order to update the local memory of each MPI process.

### **Parameters**

- [in] tree: FunctionTree to write into
- [in] src: MPI rank that last updated the function
- [in] tag: unique identifier
- [in] comm: Communicator that defines ranks

## **11.3.1 Example**

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The sharing of a FunctionTree happens in three steps: first a SharedMemory object is initialized with the appropriate shared memory communicator; then this object is used in the FunctionTree constructor; finally, after the FunctionTree has been properly computed, a call must be made to the share\_tree function. The reason for the last function call is that the internal memory pointers needs to be updated *locally* on each MPI process whenever the shared memory window has been updated.

```
// Get rank within the shared group
int rank;
MPI_Comm_rank (MPI_COMM_SHARE, &rank);
// Define master and worker ranks
int master = 0;
int worker = 1;
// The tree will be shared within the given communicator
int mem_size = 1000; //MB
mrcpp::SharedMemory shared_mem(MPI_COMM_SHARE, mem_size);
mrcpp::FunctionTree<3> tree(MRA, shared_mem);
// Master rank projects the tree
if (rank == master) mrcpp::project(prec, tree, func);
// When a shared function is updated, it must be re-shared
int tag = 333333; // Unique tag for each communication
mrcpp::share_tree(tree, master, tag, MPI_COMM_SHARE);
// Other ranks within the shared group can update the tree
if (rank == worker) tree.rescale(2.0);
// When a shared function is updated, it must be re-shared
mrcpp::share_tree(tree, worker, tag, MPI_COMM_SHARE);
```

**CHAPTER** 

## **TWELVE**

## **PRINTER**

MRCPP comes with a printer class to handle standard output:

```
#include "MRCPP/Printer"
```

The main purpose of this class is to provide (or suppress) any internal printing in MRCPP routines that might be useful for debugging. Also, it provides a sane printing environment for parallel computations that can also be used by the host program. By using the printing routines of this class, as opposed to the standard std::cout, only the master thread in a OpenMP region will provide any output while all other threads remain silent. Similarly, when running a host program in MPI parallel, the mrcpp::Printer provides three different options for handling printed output (see examples below):

- Only master rank prints to screen (stdout)
- All ranks prints to screen (stdout)
- All ranks prints to individual files

If you want only the master rank to print to an output file, this can be achieved by redirecting the output from the first option to a file (./program >file.out).

### class mrcpp::Printer

Convenience class to handle printed output.

The Printer singleton class holds the current state of the print environment. All mrcpp::print functions, as well as the println and printout macros, take an integer print level as first argument. When the global mrcpp::Printer is initialized with a given print level, only print statements with a lower print level will be displayed. All internal printing in MRCPP is at print level 10 or higher, so there is some flexibility left (levels 0 through 9) for adjusting the print volume within the host program.

### **Public Static Functions**

```
void init (int level = 0, int rank = 0, int size = 1, const char *file = nullptr)
Initialize print environment.
```

Only print statements with lower printlevel than level will be displayed. If a file name is given, each process will print to a separate file called {file}-{rank}.out. If no file name is given, only processes which initialize the printer with rank=0 will print to screen. By default, all ranks initialize with rank=0, i.e. all ranks print to screen by default.

#### **Parameters**

- [in] level: Desired print level of output
- [in] rank: MPI rank of current process
- [in] size: Total number of MPI processes

```
• [in] file: File name for printed output, will get "-{rank}.out" extension
void setScientific()
     Use scientific floating point notation, e.g. 1.0e-2.
void setFixed()
     Use fixed floating point notation, e.g. 0.01.
int setWidth (int i)
     Set new line width for printed output.
     Return Old width (number of characters)
     Parameters
           • [in] i: New width (number of characters)
int setPrecision (int i)
     Set new precision for floating point output.
     Return Old precision (digits after comma)
     Parameters
           • [in] i: New precision (digits after comma)
int setPrintLevel (int i)
     Set new print level.
     Return Old print level
     Parameters
           • [in] i: New print level
int getWidth()
     Return Current line width (number of characters)
int getPrecision()
     Return Current precision for floating point output (digits after comma)
int getPrintLevel()
     Return Current print level
```

## 12.1 Functions

Some convenience functions for printing output is provided within the mrcpp::print namespace. These functions use the data of the mrcpp::Printer class to provide pretty output of a few standard data types.

```
void mrcpp::print::environment (int level)
```

Print information about MRCPP version and build configuration.

### **Parameters**

• [in] level: Activation level for print statement

```
void mrcpp::print::separator(int level, const char &c, int newlines = 0)
Print a full line of a single character.
```

### **Parameters**

- [in] level: Activation level for print statement
- [in] c: Character to fill the line
- [in] newlines: Number of extra newlines

void mrcpp::print::header (int level, const std::string &txt, int newlines = 0, const char &c = '=') Print a text header.

### **Parameters**

- [in] level: Activation level for print statement
- [in] txt: Header text
- [in] newlines: Number of extra newlines
- [in] c: Character to fill the first line

void mrcpp::print::footer (int level, const Timer & timer, int newlines = 0, const char &c = '=')

Print a footer with elapsed wall time.

### **Parameters**

- [in] level: Activation level for print statement
- [in] t: *Timer* to be evaluated
- [in] newlines: Number of extra newlines
- [in] c: Character to fill the last line

### template<int D>

Print tree parameters (nodes, memory) and wall time.

### **Parameters**

- [in] level: Activation level for print statement
- [in] txt: Text string
- [in] tree: Tree to be printed
- [in] timer: Timer to be evaluated

void mrcpp::print::tree (int level, const std::string &txt, int n, int m, double t)
Print tree parameters (nodes, memory) and wall time.

### **Parameters**

- [in] level: Activation level for print statement
- [in] txt: Text string
- [in] n: Number of tree nodes
- [in] m: Memory usage (kB)

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• [in] t: Wall time (sec)

void mrcpp::print::time (int level, const std::string &txt, const Timer &timer)

Print elapsed time from *Timer*.

### **Parameters**

- [in] level: Activation level for print statement
- [in] txt: Text string
- [in] timer: Timer to be evaluated

void mrcpp::print::memory (int level, const std::string &txt)

Print the current memory usage of this process, obtained from system.

### **Parameters**

- [in] level: Activation level for print statement
- [in] txt: Text string

# 12.2 Macros

The following macros should replace the regular calls to std::cout:

### println (level, STR)

Print text at the given print level, with newline.

### printout (level, STR)

Print text at the given print level, without newline.

The following macros will print a message along with information on where you are in the code (file name, line number and function name). Only macros that end with \_ABORT will kill the program, all other will continue to run after the message is printed:

### MSG INFO (STR)

Print info message.

## MSG\_WARN (STR)

Print warning message.

#### MSG ERROR (STR)

Print error message, no abort.

### MSG ABORT (STR)

Print error message and abort.

### INVALID\_ARG\_ABORT

You have passed an invalid argument to a function.

### NOT IMPLEMENTED ABORT

You have reached a point in the code that is not yet implemented.

## NOT\_REACHED\_ABORT

You have reached a point that should not be reached, bug or inconsistency.

### NEEDS TESTING

You have reached an experimental part of the code, results cannot be trusted.

```
NEEDS FIX (STR)
```

You have hit a known bug that is yet to be fixed, results cannot be trusted.

# 12.3 Examples

Using the print level to adjust the amount of output:

Using headers and footers to get pretty output:

This will produce the following output:

```
Projecting analytic function

Projected function 520 nds 16 MB 0.09 sec
Integrated function (au) 9.99999999992e-01

Wall time: 9.32703e-02 sec
```

As mentioned above, when running in MPI parallel there are three different ways of handling printed output (master to stdout, all to stdout or all to files). These can be chosen by adding appropriate arguments to init. The default setting will in a parallel environment have all MPI ranks printing to screen, but by adding MPI info to the printer, we can separate the output of the different ranks:

```
int level = 10;
int wrank, wsize;
MPI_Comm_rank(MPI_COMM_WORLD, &wrank); // Get my rank
MPI_Comm_size(MPI_COMM_WORLD, &wsize); // Get total number of ranks

// All ranks will print to screen
mrcpp::Printer::init(level);

// Only master rank will print to screen
mrcpp::Printer::init(level, wrank, wsize);
```

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// All ranks will print to separate files called filename-<rank>.out
mrcpp::Printer::init(level, wrank, wsize, "filename");

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**CHAPTER** 

## **THIRTEEN**

# **PLOTTER**

MRCPP comes with its own plotter class which can be used by the host program to generate data files for visualization using e.g. gnuplot, paraview, blob and geomyiew. These features are available by including:

```
#include "MRCPP/Plotter"
```

### template<int D>

## class mrcpp::Plotter

Class for plotting multivariate functions.

This class will generate an equidistant grid in one (line), two (surf) or three (cube) dimensions, and subsequently evaluate the function on this grid.

The grid is generated from the vectors A, B and C, relative to the origin O:

- a linePlot will plot the line spanned by A, starting from O
- a surfPlot will plot the area spanned by A and B, starting from O
- a cubePlot will plot the volume spanned by A, B and C, starting from O

The vectors A, B and C do not necessarily have to be orthogonal.

The parameter D refers to the dimension of the *function*, not the dimension of the plot.

### **Public Functions**

```
Plotter (const Coord<D> &o = \{\})
```

Return New Plotter object

### **Parameters**

• [in] o: Plot origin, default (0, 0, ..., 0)

void **setSuffix** (int *t*, **const** std::string &s)

Set file extension for output file.

The file name you decide for the output will get a predefined suffix that differentiates between different types of plot.

### **Parameters**

- [in] t: Plot type (Plotter < D > :: Line, :: Surface, :: Cube, :: Grid)
- [in] s: Extension string, default .line, .surf, .cube, .grid

## void setOrigin (const Coord<D>&o)

Set the point of origin for the plot.

#### **Parameters**

```
• [in] o: Plot origin, default (0, 0, ..., 0)
```

void setRange (const CoordD> &a, const CoordD> &b = {}, const CoordD> &c = {}) Set boundary vectors A, B and C for the plot.

#### **Parameters**

- [in] a: A vector
- [in] b: B vector
- [in] c: C vector

### void gridPlot (const MWTree<D> &tree, const std::string &fname)

Grid plot of a MWTree.

Writes a file named fname + file extension (".grid" as default) to be read by geomview to visualize the grid (of endNodes) where the multiresolution function is defined. In MPI, each process will write a separate file, and will print only nodes owned by itself (pluss the rootNodes).

#### **Parameters**

- [in] tree: MWTree to plot
- [in] fname: File name for output, without extension

Parametric plot of a function.

Plots the function func parametrically with npts[0] along the vector A starting from the origin O to a file named fname + file extension (".line" as default).

### **Parameters**

- [in] npts: Number of points along A
- [in] func: Function to plot
- [in] fname: File name for output, without extension

Surface plot of a function.

Plots the function func in 2D on the area spanned by the two vectors A (npts[0] points) and B (npts[1] points), starting from the origin O, to a file named fname + file extension (".surf" as default).

### **Parameters**

- [in] npts: Number of points along  $\boldsymbol{A}$  and  $\boldsymbol{B}$
- [in] func: Function to plot
- [in] fname: File name for output, without extension

Cubic plot of a function.

Plots the function func in 3D in the volume spanned by the three vectors A (npts[0] points), B (npts[1] points) and C (npts[2] points), starting from the origin O, to a file named fname + file extension (".cube" as default).

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### **Parameters**

- [in] npts: Number of points along A, B and C
- [in] func: Function to plot
- [in] fname: File name for output, without extension

**Note:** When plotting a FunctionTree, only the *scaling* part of the leaf nodes will be evaluated, which means that the function values will not be fully accurate. This is done to allow a fast and const function evaluation that can be done in OMP parallel. If you want to include also the *final* wavelet corrections to your function values, you'll have to manually extend the MW grid by one level before plotting using mrcpp::refine\_grid(tree, 1).

# 13.1 Examples

A parametric line plot of a three-dimensional function along the z axis [-1, 1]:

A surface plot of a three-dimensional function in the x=[-2,2], y=[-1,1], z=0 plane:

```
int aPts = 2000;
                                                     // Number of points in a
int bPts = 1000;
                                                     // Number of points in b
mrcpp::Coord<3> o{-2.0, -1.0, 0.0};
                                                     // Origin vector
mrcpp::Coord<3> a{ 4.0, 0.0, 0.0};
                                                     // Boundary vector A
mrcpp::Coord<3> b{ 0.0, 2.0, 0.0};
                                                     // Boundary vector B
mrcpp::Plotter<3> plot(o);
                                                     // Plotter of 3D functions
                                                     // Set plot range
plot.setRange(a, b);
                                                    // Write to file f_tree.surf
plot.surfPlot({aPts, bPts}, f_tree, "f_tree");
```

A cube plot of a three-dimensional function in the volume x=[-2,2], y=[-1,1], z=[0,2]:

```
int aPts = 200;
                                                      // Number of points in a
                                                      // Number of points in b
int bPts = 100;
int cPts = 100;
                                                      // Number of points in c
mrcpp::Coord<3> o\{-2.0, -1.0, 0.0\};
                                                     // Origin vector
mrcpp::Coord<3> a{ 4.0, 0.0, 0.0};
                                                     // Boundary vector A
mrcpp::Coord<3> b{ 0.0, 2.0, 0.0};
                                                     // Boundary vector B
mrcpp::Coord<3> b{ 0.0, 0.0, 2.0};
                                                     // Boundary vector C
mrcpp::Plotter<3> plot(o);
                                                      // Plotter of 3D functions
plot.setRange(a, b, c);
                                                     // Set plot range
plot.cubePlot({aPts, bPts, cPts}, f_tree, "f_tree"); // Write to file f_tree.cube
```

A grid plot of a three-dimensional FunctionTree:

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## **CHAPTER**

# **FOURTEEN**

# **TIMER**

MRCPP comes with a timer class which can be used by the host program:

#include "MRCPP/Timer"

```
class mrcpp::Timer
     Records wall time between the execution of two lines of source code.
     Public Functions
     Timer (bool start_timer = true)
          Return New Timer object
          Parameters
                • [in] start_timer: option to start timer immediately
     Timer (const Timer &timer)
          Return Copy of Timer object, including its current state
          Parameters
                • [in] timer: Object to copy
     Timer &operator=(const Timer &timer)
          Return Copy of Timer object, including its current state
          Parameters
                • [in] timer: Object to copy
     void start()
          Start timer from zero.
     void resume()
          Resume timer from previous time.
     void stop()
          Stop timer.
     double elapsed() const
```

**Return** Current elapsed time, in seconds

# 14.1 Examples

The timer records wall (human) time, not CPU user time. The clock will by default start immediately after construction, and will keep running until explicitly stopped. The elapsed time can be evaluated while clock is running:

The timer can also be started explicitly at a later stage *after* construction, as well as explicitly stopped after the work is done. Then the *elapsed()* function will return the time spent between *start()* and *stop()*:

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**CHAPTER** 

**FIFTEEN** 

## PROGRAMMERS MANUAL

# 15.1 Clang-tidy

To ensure modern coding conventions are followed developers are encouraged to run clang-tidy on the code. Ensure clang-tidy is installed. Then to display available checkers run:

```
$ clang-tidy --list-checks -checks='*' | grep "modernize"
```

### This will generate a list looking like this:

```
$ modernize-avoid-bind
$ modernize-deprecated-headers
$ modernize-loop-convert
$ modernize-make-shared
$ modernize-make-unique
$ modernize-pass-by-value
$ modernize-raw-string-literal
$ modernize-redundant-void-arg
$ modernize-replace-auto-ptr
$ modernize-replace-random-shuffle
$ modernize-return-braced-init-list
$ modernize-shrink-to-fit
$ modernize-unary-static-assert
$ modernize-use-auto
$ modernize-use-bool-literals
$ modernize-use-default-member-init
$ modernize-use-emplace
$ modernize-use-equals-default
$ modernize-use-equals-delete
$ modernize-use-noexcept
$ modernize-use-nullptr
$ modernize-use-override
$ modernize-use-transparent-functors
$ modernize-use-using
```

To run any of these modernization's on the code, go to your build directory. From there run the command:

```
$ run-clang-tidy -header-filter='.*' -checks='-*, modernize-your-modernization' -fix
```

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