QUEST User's Guide

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0 Preface

This version of the QUEST User's Guide intends to eventually replace previous renditions of similar user's guides for QUEST. As of this writing, there are three of such renditions:

```
https://www.cs.ucdavis.edu/~bai/QUEST/manual.pdf (dated April 2012)
manual_old.pdf (available in this repository, dated November 2013)
https://code.google.com/archive/p/quest-qmc/ (wikis section has similar content, undated)
```

This document is made from a copy of the original .tex file that created manual_old.pdf, though it might change to a brand new .tex file or website in the future. Keep in mind that some of the information presented in this guide might be incorrect: this being an artefact of the old user's guide. It is intended for these errors to be fixed in this user's guide in the future, and for more content and detailed explanations to be provided.

Comments or suggestions (for this document or anything in the QUEST repository) are welcome to be emailed to jwwalker@ucdavis.edu or opened as an issue or discussion on the repository page.

1 Introduction

Quantum Electron Simulation Toolbox (QUEST) is a Fortran 90/95 package for performing determinant quantum Monte Carlo (DQMC) methods for strongly correlated electron systems. Its development was motived by a FORTRAN 77 DQMC code¹ written and maintained by Richard Scalettar at the University of California, Davis (UCD). The intention for creating QUEST was to modernize the the legacy code by incorporating three principles:

- 1. Improved simulation performance: QUEST has improved the performance of the legacy code simulations by using new algorithms, like delayed update, and by integrating modern numerical kernels, BLAS/LAPACK. A six to eight times speedup had been observed for medium sized simulations.
- 2. To integrate existing programs: QUEST has integrated many legacy codes by modularizing their computational components, which makes QUEST not only a single program, but a toolbox. The advantages of modularization also include the ease of maintenance and the convenience of program interfacing.
- 3. To assist new simulations development: QUEST has several desired properties for developing new simulations, such as the ability of creating new lattice geometries. Several novel simulations had been done by using QUEST.

Currently, QUEST is still under development and debugging. The latest version can be downloaded from the following link:

```
https://github.com/Meromorphics/quest
```

Additional information about QUEST (including files for older versions of QUEST) can be found at:

```
https://code.google.com/archive/p/quest-qmc/
https://www.cs.ucdavis.edu/~bai/QUEST/index.html
```

¹Known as the legacy code.

1.1 Problems that QUEST can solve

If no geometry is specified ², QUEST uses a two-dimensional periodic rectangular lattice as the default geometry in a simulation ³. For more information about running simulations in QUEST, see section 1.4 for the most essential information about running simulations (enough for a general end-user) and section SECTION for more advanced simulation configuration.

QUEST is stable for interaction strength from U=0 to U=16t and inverse temperatures $\beta=0$ to 20/t. Here U is the onsite repulsion and t is the coefficient of the kinetic energy term, see Section 2.1 for more information about these parameters. However, away from half-filling, $\mu=0$, error bars are large for large β (sign problem). The current implementation allows hundreds of sites to be simulated on commodity computers within a reasonable time. Appendix A provides performance benchmark for some example problems.

1.2 Organization of files

The top level files and directories of QUEST include

- README.md: general information about QUEST.
- Makefile: computer instructions for compiling QUEST.
- /SRC: directory for source code
- /EXAMPLE: directory for executable programs.
- /doc: directory for documentation.
- /makefiles: unused makefiles
- /geometries: example geometry files
- /OpenBLAS: copy of OpenBLAS source code
- /ford_documentation: directory for documentation generated by FORD ⁴.
- ford.md: instructions for generating FORD documentation

Under the EXAMPLE directory, several example programs are contained⁵.

- test: A simple test program for 2 dimensional square lattice.
- verify: A program verifying the correctness of QUEST by examining its results against theoretical values of two special cases, U = 0 and t = 0.
- tdm: A test program for time dependent measurements.
- parallel: A test program for MPI type parallelization.

²See sections 2.1 and 3 for more information about specifying geometries

³At the time of writing, this default geometry feature is bugged. Input files must be provided (this will be fixed in a future update).

⁴https://forddocs.readthedocs.io/en/latest/

⁵Only verify.F90 and ggeom.F90 are present from this list. This remainder of this list is kept here as a reminder for potential programs to include in a future update.

- wrap: Programs for different input and output formats.
- sheet: Programs for multilayer geometry.
- DCA: Programs for using DQMC as DCA solver.
- geom: Programs for general geometry and their simulations

The details for each program will be illustrated in Appendix A^6 .

1.3 Installation

Installation can be a pain for many programs, so this section attempts to be overly complete to try and make the process as easy as possible. For a more advanced user, the following concise instructions should be enough: first clone the QUEST repository (link just below), then enter it and run make. QUEST is developed and tested to run on a Linux (Ubuntu) system with Intel's oneAPI HPC toolkit installed in its default location (change MKLPATH in the Makefile if it is installed elsewhere). Potential support for the gnu compiler is included in the Makefile, but compilation using this compiler is untested. Edit the root directories Makefile as necessary.

The main (and currently being updated) repository for QUEST can be found at:

https://github.com/Meromorphics/quest

As of this writing, QUEST is only configured to run on Linux (specifically, it is developed on a system running Ubuntu 22.04.3 LTS). Support for Mac and Windows is planned for a future release, but users of these systems might be able to get QUEST to work ⁷. The easiest method installation is through the use of git. Assuming git is installed and the command git may be used on a terminal, QUEST may be installed by entering the following on the terminal:

git clone https://github.com/Meromorphics/quest.git

Which should create a directory named quest in the current working directory. After git installation, the new directory should be entered by entering on the terminal:

cd quest

Compilation of QUEST requires gnumake. To test if gnumake is installed, in a directory where no file named Makefile is present (eg an empty directory) entering the following in a terminal:

make

should return as output:

⁶verify.F90 is explained in section SECTION and ggeom.F90 in section SECTION. Appendix A is being kept as a reminder for potential programs to include in a future update.

⁷Before a bug with Intel's ifx compiler running QUEST was found, QUEST was confirmed to be running on Windows Subsystem for Linux (WSL). If a user wishes to run QUEST on a Mac or Windows system, the issue is most likely found within the Makefile of QUEST.

make: *** No targets specified and no makefile found. Stop

or some similar statement. Assuming gnumake is installed and the current working directory is quest, QUEST may be compiled by entering the command make on the terminal (identical statement to testing the gnumake installation). For many users, an error is likely to occur at this step. This is primarily due to how QUEST is developed. QUEST is developed on Ubuntu using compilers installed from Intel's oneAPI:

https://www.intel.com/content/www/us/en/developer/tools/oneapi/toolkits.html#gs.3kfmxb

In its default configuration, QUEST requires the oneAPI Base and HPC toolkits to be installed (these give the required compilers and LAPACK library, among other things). To change the installation configuration of QUEST, the Makefile in the root directory (directory just cd'd into) of QUEST must be configured ⁸. By default, QUEST assumes the Intel oneAPI ifort is being used ⁹ and this is installed in the default directory. If oneAPI is not installed in its default directory, the line:

```
MKLPATH = $(MKLROOT)/include/intel64/inp64
```

in Makefile may have to be changed to the Intel MKL library path. By default, QUEST is designed to run on one CPU thread. To run on multiple CPU threads by use of MPI, change the following line in Makefile:

```
PRG_FLAGS = $(FLAG_BSOFI) $(FLAG_ASQRD) $(FLAG_CKB) $(FLAG_CUDA)
```

to:

```
PRG_FLAGS = $(FLAG_BSOFI) $(FLAG_ASQRD) $(FLAG_CKB) $(FLAG_CUDA) -D_QMC_MPI
```

Note that at the time of writing, the MPI implementation of QUEST has a bug, so running on a single thread is required until this is fixed. QUEST has what older documentation says is "experimental GPU support." This GPU support has not been tested at all recently (no idea if it works), but there are remnants in the Makefile that may possibly get it to work.

Furthermore, before using the command make, the user must have oneAPI's environment variables set in their active terminal. To test if these environment variables are set, the user may enter on the terminal:

ifort

If the environment variables are not set, the output should have an appearance similar to:

⁸Quest contains many makefiles within its directories. Only the makefile located in the root directory should be ran using make, attempting to run make on the others will result in an error (but these files are still required for QUEST to compile).

⁹Attempting to use Intel oneAPI's ifx compiler results in an error when running ./ggeom. This bug is intended to be fixed in a later release. For now, using ifort will result in compiler warnings but no errors.

```
Command 'ifort' not found, did you mean:
command 'fort' from deb fort-validator (1.5.3-1build1)
command 'isort' from deb isort (5.6.4-1)
Try: sudo apt install <deb name>
```

If environment variables are set correctly, the output should have an appearance similar to:

```
ifort: remark #10448: Intel(R) Fortran Compiler Classic (ifort) is now deprecated and will be discontinued late 2024.

Intel recommends that customers transition now to using the LLVM-based Intel(R) Fortran Compiler (ifx) for continued Windows* and Linux* support, new language support, new language features, and optimizations. Use '-diag-disable=10448' to disable this message.

ifort: command line error: no files specified; for help type "ifort -help"
```

On a fresh installation of Linux, it is known that libomp must be installed for compilation to work correctly, which may be installed by running in a terminal:

```
sudo apt-get install libomp-dev
```

It is unknown if any other similar dependencies are required to install QUEST. User research may be required for such dependencies may be required ¹⁰.

Assuming compilation of QUEST succeeded with no errors (many warnings are expected and will be fixed in a future update), the user should verify that QUEST runs as expected as follows. Enter the directory EXAMPLE/verify (eg by entering cd EXAMPLE/verify in the root directory of QUEST). Then run the file verify by entering on the terminal:

```
./verify
```

Unless output was directed to a file (eg by using > output_file in the above command), text should begin to fill the terminal. An example of what output should be running down the terminal screen is:

```
Parameters: t = 0.00, mu = -0.50, U = 0.00, beta = 1.50,

Theoretical | Computed (avg +- error) | |T-C|: error

Density: 0.641643 | 0.641643 +- 0.000000 | 0.00: 0.00

Energy: 0.320821 | 0.320821 +- 0.000000 | 0.00: 0.00

Double occupancy: 0.000000 | 0.000000 +- 0.000000 | 0.00: 0.00
```

 $^{^{10}}$ If any other dependencies are found to be required, please create an issue or discussion on the Github repository or email <code>jwwalker@ucdavis.edu</code>

The most important part is what is printed at the very end:

```
81.94% within 1 error bar (Expected 63.2%).
95.83% within 2 error bar (Expected 86.5%).
Running time: 6.723399 (second)
```

The two percentages in the left columns should be around or greater than the two percentages in the right column. If this is not the case, QUEST ran incorrectly and further analysis of simulations should not be performed until either QUEST is bugfixed or installed correctly. The reason being is that this code tests out QUESTS predictions for two Hubbard models that have an analytic solution: the Hubbard model with 1 site, and a 4×4 Hubbard model with no Coulomb interaction (U = 0). See EXAMPLE/verify/verify.F90 for more details. These two tests are particularly important if you wish to make your own changes to QUEST (to make sure QUEST runs correctly).

As long as make ran without issue and verify gives good sounding results, QUEST should in theory be installed correctly.

1.4 Basic simulation usage

In essence, QUEST runs user defined "geometry" files and returns results (measurements from a simulation defined by an input geometry). Advanced usage of QUEST is just fine tuning what is returned and how (see section 3 for more details). In this section we detail how to get QUEST to give you results from input geometry files (details on how to make geometry files are in section SECTION). For a typical end-user, running simulations using the instructions provided here is enough.

To run a simulation in the most basic form, three files are required. QUEST provides one of these files:

```
/EXAMPLE/geom/ggeom.F90
```

It is strange and confusing that such an important file is located in a directory of the example directory, but such is life. This file performs simulations (it is the "simulator"). The second required file is an input file. The input file contains the information that the simulator uses to perform a simulation. To run an input file (assuming the terminal has as its working directory EXAMPLE/geom), enter:

```
./ggeom <input_file>
```

where **<input_file>** is the input file to **ggeom**. For example, upon installation, QUEST has one input file named **in** ready to run (for examples sake, this file may be deleted without harm):

```
./ggeom in
```

An input file contains parameters defining how the simulation should be run (such as how many warmup steps should be done for Monte Carlo, and the imaginary time step for the Suzuki-Trotter

expansion). If some parameters are skipped in the input file that are used in the simulator, these parameters are given "good" default values ¹¹ (see section 3 for more details about default parameter values). Precise details about input files are shown in section SECTION, but here we will detail two lines in an input file that are of particular importance. Any input file should ¹² contain the following two lines:

```
ofile = <output>
gfile = <geometry>.geom
```

ofile stands for output file, since QUEST will create (or overwrite) three output files with names:

```
<output>.geometry
<output>.out
<output>.tdm.out
```

So <output> should be a name (no file extension). For example, the file in mentioned earlier has ofile = test so the output would be three files names test.geometry, test.out, and test.tdm.out. See section 2.5 for details of the contents of these files (basically measurement results and a restatement of the input geometry). <geometry>.geom is the third and final user defined file required to run a simulation in its most basic form. As the name suggests, this file defines the geometry of the simulation. There are a few methods in which a .geom file may be formatted, see section 3 for more details. A sample geometry file is in the same directory as ./ggeom, and more can be found in /geometries.

2 Input/output basics

In this section we detail the basics of input and output in QUEST. In section 1.4 we mentioned how the program geom requires 2 user-input files and returns 3 output files. This section details the basics of those types of files, referred to as input and output files. For many, the QUEST output described in this section should suffice for most purposes. If a user wishes for QUEST to give other output than what is described here, they should consult section 3 for advanced output usage. Compared to output, a typical user would be more likely to find basic input insufficient for their desires (specifically lattice geometry). For this, we again refer the user to section 3 for advanced input configuration, but remind the user that there are many geometry template files located in the directory /geometries that might be of use for this issue.

2.1 Input files

QUEST requires two files for input. We will call these files the in file and the geometry file. The in file specifies how the simulation should be run (Metropolis algorithm configuration, measurement bin amount, ...), while the geometry file specifies the lattice to be simulated (hoppings, interaction energy, site locations, ...). The geometry file may be one of two types:

¹¹It is a known bug that some parameters are not actually taking default values when running the simulator says that they are. This will be fixed in a future release

¹²Again, with no bugs, QUEST should give "good" default values, but for practical purposes these two parameters should always be defined.

- free form (.def file extension)
- general geometry (.geom file extension)

Internally, QUEST stores lattice geometry information in a derived data type named Struct. When a geometry file is input, QUEST translates the contents into the Struct datatype so it may be used in its code. More details about the inner workings of Struct are provided in section 3.

In free form input, the user has complete control over how the lattice geometry is setup. This comes at a cost: the user must provide all information about the lattice (the hopping values between all sites on the lattice, information about the neighbors of all sites, ...). If the user desires to use this form of input, they might want to create programs designed for making these files.

In practice, general geometry input is used. In general geometry input, the user describes how the lattice geometry is constructed in terms of primitive cells. Basis vectors describing where to put cells are input, along with the Cartesian coordinates relative to cell centers of the sites within that cell (this is a potentially unintuitive part of QUEST: first a set of vectors describe where to put cells, then another set of vectors describes where to place sites in that cell; sites hold electrons, cells hold sites) and hoppings between the different types of sites defined in a cell; along with other parameters that will be detailed in this section. General geometry input allows geometries to be defined in ~ 50 lines of text, compared to free form, which can easily take thousands of lines for the same geometry. The drawback is a loss of control: the lattice is uniformly constructed.

2.2 in input files

As mentioned in section 2.1, in input files contain three things:

- a name for output files
- the name of the input geometry file
- simulation parameter configuration

A template for creating in files is located at geometries/templates/input_template. in files have the following syntax rules:

- # declares a comment for the remainder of the line
- parameter = value assigns value to parameter
- commands must be input on one line
- spaces only matter only in the specification of a token (eg 1.2 is not equivalent to 1. 2)

So an in file can be interpreted as a sequence of assignment statement separated by lines with optional comments (along with the usual rules for spaces). Somewhere in the in file (generally at the very top), two parameters must be assigned a value¹³:

• ofile (string)

¹³ Technically only the gfile, but good practice is to specify a name for ofile

• gfile (.geom or .def)

We will call the above two parameters file parameters. Technically only file parameters are the only parameters required in an **in** file. The other parameters that can be specified in an input can be classified into the following groups: Hubbard model, Metropolis algorithm, measurement, and numerical parameters. The Hubbard model parameters are as follows:

- mu_up (real): chemical potential for up-spin electrons
- mu_dn (real): chemical potential for spin-down electrons
- L (integer): number of imaginary time slices $(L = \beta/\Delta\tau)$
- HSF (integer)¹⁴: indicator of how the Hubbard Stratonovich field (HSF) is input
 - − -1: randomly generated HSF
 - − 0: use HSF in memory
 - 1: HSF is read from a file
- bcond (real, real, real): boundary conditions

The Metropolis algorithm parameters are as follows:

- nwarm (integer): number of warmup sweeps
- npass (integer): number of measurement sweeps
- ntry (integer): number of sites flipped in a global sweep
- tausk (integer): specifies frequency of performing physical measurements
- tdm (integer): whether or not to perform time-dependent measurements
 - 0: no time-dependent measurements are performed
 - 1: time-dependent measurements are performed

The measurement parameters are as follows:

- nbin (integer): determines how the computed data is divided into bins
- nhist (integer): unknown currently

The numerical parameters are as follows:

- north (integer): frequency of performing orthogonalization in matrix products¹⁵
- nwrap (integer): frequency of of performing Green's function calculation. QUEST dynamically adjusts this parameter according to the errors of updating. Numerical errors accumulates quickly for larger values

 $^{^{14}\}mathrm{At}$ the moment it is unknown what memory or input HSF means in implementation; but this option exists somehow

¹⁵Used when using a stabilization algorithm in the calculation of the Green's function

- fixwrap (integer): unknown currently 16
- errate (real): tolerable error rate of recomputing
- difflim (real): tolerable difference of the matrices computed from different methods

2.3 General geometry input files

As stated in section 2.1, general geometry input files describe lattice geometry by describing how the lattice is generated by primitive cells and how sites interact. Also mentioned there is why general geometry files are more practical to use compared to free form input files: they can be handwritten in much fewer lines.

General geometry input files have a very specific and restrictive syntax and must end in a .geom file extension. The syntax of a general geometry input file is as follows:

```
#FLAG
<FLAG DEPENDENT SYNTAX>
```

Blank lines are allowed. No comments are allowed. A #FLAG must appear alone on its own line. The #FLAG command indicates that the following lines of code up until the next #FLAG command are to be read by that particular flag. By flag, we mean parameter flag. For example, in #NDIM, NDIM is the flag (alias) for the number of dimensions parameter. There are 11 total parameters that may be specified in a general geometry file, but not all are required to be used. The directories /geometries and /geometries/templates have example general geometry files and templates for general geometry files (the user changes a few values to suite their need, eg for the template square_template the user should input how many x and y units the lattice should span). We will now detail the 11 different parameters that may be included in a general geometry file.

#NDIM: the number of dimensions of the lattice (1, 2, or 3). The value input here changes how #SUPER, #K-POINT, and #PHASE may be input since these require dimensional input. The syntax of #NDIM is as follows:

```
#NDIM
```

where n is either 1, 2, or 3.

#PRIM: the primitive vectors defining the basis for the lattice in Cartesian coordinates. A 3×3 array of real numbers. The syntax of #PRIM is as follows:

```
#PRIM a1x a1y a1z a2x a2y a2z a3x a3y a3z
```

¹⁶From https://code.google.com/archive/p/quest-qmc/wikis/input_parameters.wiki: NO IDEA. BUT! When ASQRD is enabled and fix wrap is set to 1, north and nwrap cannot be equal.

The primitive vectors are read row-wise. If NDIM is equal to 2, then the last row may be omitted or set to 0 0 1. Similarly if NDIM is equal to 1.

#SUPER: defines the supercell of the lattice. An NDIM x NDIM integer array. Defines the 'size' of the lattice: entries are weights of the primitive vectors. The syntax of #SUPER for an NDIM = 3 lattice is as follows:

#SUPER					
S1x	S1y	S1z			
S2x	S2y	S2z			
S3x	S3y	S3z			

Lower dimension lattices have similar syntax. For example, for an $\mathtt{NDIM} = 2$ lattice with the first row of $\#\mathtt{SUPER}$ being $\mathtt{x} \ 0$ and second $\mathtt{0} \ \mathtt{y}$ means that the lattice is obtained by using translation in the 1st primitive cell vector direction up to \mathtt{x} times and similarly the 2nd primitive cell vector up to \mathtt{y} times.

#K-POINT: NDIM long real vector. Defines the original k-point. Currently, the only supported value of this parameter is (0,0,0) (the default value of this parameter) so this flag may be omitted in geometry files without worry.

#ORB: rows listing out the sites within a primitive cell. The amount of rows is arbitrary (allowing an arbitrary amount of sites to be made in each primitive cell). The syntax of #ORB is as follows:

where the rows following #ORB may be defined at an arbitrary length. label is a string that serves as a label for the particular type of site, x y z describes the location of the site relative to the origin of the cell in Cartesian coordinates, #N is the number of the site being defined (beginning with 0 and incrementing by 1 for successive rows).

#HAMILT: defines the Hamiltonian of the system. The syntax of #HAMILT is as follows:

```
#HAMILT label2 x y z tup tdn U
```

where the rows repeat until the Hamiltonian is sufficiently described. The meaning of a row is as follows. label1 and label2 indicate that the row describes how those labeled site-types interact. label1 and label2 are the #N integers defined in #ORB (eg particles 1 and 3 interact in such a specified way in the Hamiltonian). Multiple rows starting with the same label1 and label2 may be required for a full description for reasons soon to be explained. x y z describes the vector connecting label1 to label2. Describing label2 to label1 is not needed (QUEST handles considering going from site A to B and B to A when considering only A to B or B to A). Multiple rows for the same set of labels may be required so electrons may hop to all adjacent primitive cells and not

just to a specific primitive cell. For example, if a lattice is filled out with primitive cells describing a square and each primitive cell has sites A and B in it, then just describing how an electron in an A site might get to a B site by moving 'to the right' might not be sufficient since there is a B cell right above it also. tup and tdn describe the hopping value for an electron hopping between sites with labels label1 and label2, where tup is the hopping value for up-spin electrons and tdn for down-spin electrons (this may be different values for up and down spin electrons). U describes interaction energy. For a single site indexed by label to have an interaction energy, the following row should be present:

label label 0.0 0.0 0.0 0.0 0.0 U

where U is the interaction energy. For differing labels and nonzero x y z, specifying a U value allows for the implementation of nearest neighbor interaction (or not-near, if x y z is made large).

#SYMM: QUEST allows for the input of 3 types of point group symmetry operations: rotation about an axis, mirror plane, and inversion. The main purpose of declaring symmetries is for simulation speedup. The syntax of declaring point group symmetries is as follows:

#SYM	M					
CN	Х	У	Z	x1	у1	z1
D	X	У	Z	x1	y1	z1
I	х	У	z			

The first row describes rotation about an axis symmetry. This first row states the following: the lattice is symmetric under rotations $2\pi/N$ (where N is the integer defined in CN) about the axis defined by containing the point (x,y,z) in the direction (x1,y1,z1). The second row describes mirror plane symmetry. This second row states the following: the lattice is symmetric under reflections about the plane defined by containing the point (x,y,z) with normal (x1,y1,z1). The third row describes inversion symmetry: $\mathbf{r} \mapsto -\mathbf{r}$ symmetry with inversion point (x,y,z).

#PHASE: an NDIM x NDIM integer array. Unknown usage¹⁷.

#BONDS: specifies the types of bondings between orbital types. The syntax of #BONDS is as follows:

label1 label2 x y z #bond_label

where each row is repeated as required to fully specify bonds. label1 and label2 are the labels of two types of orbitals (which can be the same). x y z is the vector connecting label1 to label2. bond_label labels the bond type, and should be incrementing down rows starting from 1.

#PAIR: specifies the types of pairings to be measured. Uses linear combinations of the entries in #BOND in its definitions of pairings; if not included QUEST will attempt to determine the pairings on its own. It is unknown how to fully implement this.

#END: declares the end of the geometry file. Syntax:

¹⁷Some information about the functionality is available at https://code.google.com/archive/p/quest-qmc/wikis/Lieb_lattice_tutorial.wiki

2.4 Free form input files

To be implemented in a future version. For now the old version text about input files is being kept. An input file of QUEST is consisted of a list of parameter assignments,

```
parameter = value
```

and single line comments, which are of the form (comments can only be declared one line at a time):

#<comment>

Each parameter is associated with one of the following types: *integer*, *real*, *real array*, and *string*. The basic parameters include

- 1. ofile (string): file name for output files (<name>.geometry, <name>.out, <name>.tdm.out).
- 2. gfile (string.geom): geometry file for lattice
- 3. n (integer): total number of sites.
- 4. nx (integer): number of sites in the x-direction.
- 5. ny (integer): number of sites in the y-direction.
- 6. nz (integer): number of sites in the z-direction.
- 7. U (real array): Hubbard parameters.
- 8. t (real array): Hubbard parameters.
- 9. mu (real array): Hubbard parameters.
- 10. L (integer): number of time slice.
- 11. dtau (real): discretization parameter.
- 12. HSF (integer): indicator of how Hubbard Stratonovich field is input.
- 13. HSFin (string): file name of input file of HSF.
- 14. HSFout (string): file name of output file of HSF.
- 15. nwarm (integer): number of warmup sweeps.
- 16. npass (integer): number of measurement (sample) sweeps.
- 17. nmeas (integer): frequency of performing equal time measurements.
- 18. tausk (integer): frequency of performing unequal time measurements.

- 19. nbin (integer): number of bins for measurement results.
- 20. seed (integer): random number seed.
- 21. north (integer): frequency of performing orthogonalization in matrix product.
- 22. nwrap (integer): frequency of performing recomputing in Green's function calculation.
- 23. difflim (real): tolerable difference of the matrices computed from different methods.
- 24. errrate (real): tolerable error rate of recomputing.
- 25. ntry (integer): number of sites to be flipped in the global sweep. UPD: This parameter controls the frequency of a type of Monte Carlo move which is needed for $U \ge 8$.

The parameters can be roughly divided into three groups:

1. Parameters for Hubbard model:

The parameters nx and ny specify the x-dimension and y-dimension of the 2D rectangular lattice to be simulated. Note that parameter n, the number of total sites, is equivalent to $nx \times ny$. The parameters t, mu, and U are used in the Hubbard Hamiltonian

$$\mathcal{H} = -t \sum_{\langle i,j \rangle, \sigma} \left(c_{i\sigma}^{\dagger} c_{j\sigma} + c_{j\sigma}^{\dagger} c_{i\sigma} \right) - \mu \sum_{i,\sigma} n_{i\sigma} + U \sum_{i} \left(n_{i\uparrow} - \frac{1}{2} \right) \left(n_{i\downarrow} - \frac{1}{2} \right),$$

for kinetic energy, chemical energy and potential energy respectively. Parameter L and dtau are for inverse temperature β .

$$\beta = L\Delta \tau.$$

Parameter HSF indicates how the Hubbard-Stratonovich Field (HSF) is initialized.

$$HSF = \begin{cases} -1, & \text{randomly generating HSF}; \\ 0, & \text{use HSF in the memory}; \\ 1, & \text{read HSF from a file.} \end{cases}$$

If HSF=1, then HSFin is the input file name. The generated HSF can be also output to a file by specifying the file name in HSFout.

- 2. Parameters for Monte Carlo simulation and physical measurements: Parameter nWarm and nPass in the second group decide how many Monte Carlo loops need be executed for warm up and measurement sweep. Parameter nMeas and tausk specify the frequency of performing physical measurements. Parameter nBin determine how the computed data is divided into bins. Parameter ntry is used to specify how many global flipping should be performed per sweep for the large U.
- 3. Parameters regarding numerical concerns.

Parameter seed is used for random number generator. If it is 0, a new seed will be generated from system time. Parameter nOrth specifies how often the stabilization algorithm should be performed in the calculation of Green's function. Parameter nWrap provides the initial frequency of recomputing Green's function. In QUEST, nWrap will be dynamically adjusted according to the errors of updating. Parameters difflim and errrate are used for the adjusting algorithm, which specify the tolerable matrix difference and the acceptable error rate.

In its current version, QUEST assumes all simulations are done for the Hubbard model. The source directory contains files that can be used to simulate the Holstein model, but these have not been tested (and implementation must manually be done if these were to be tested out, the Holstein model does not have a ggeom.F90 file).

2.5 Output Results

The output of QUEST varies in different programs. The goal for this section is to introduce what QUEST can output, and their formats. The output of QUEST can be classified into three types

- Input/condituration parameters.
- Equal time measurements.
- Unequal time measurements.

The input/configuration parameters are as introduced in the previous section. Some of them may be created or changed during the simulation, like **seed** and **nWrap**. Those configuration parameters can help identifying the output results.

In terms of formats, concerned only with measurements, there are three kinds

- Single real number.
- Array of real numbers.
- Array of complex numbers.

If a measurement is a single real number, it will be shown with three terms: name, average, and error. For example,

```
Density: 1.000000 +- 0.000000
```

The measurement formatted in an array of real numbers (or complex numbers) is a function, whose arguments can be anything, like the distances between sites. The output of this type enumerates all its function arguments and values. For example, the equal time Green's function of a 4×4 periodic lattice is output like

Equal time Green's function:

```
dx = 0, dy =
                        0.500000 +-
                                         0.00000
dx = 1, dy =
                       -0.119358 +-
                                         0.000754
dx =
     2, dy =
                        0.000000 + -
                                         0.000000
dx = 0, dy =
                       -0.118936 +-
                                         0.000414
dx = 1, dy =
                        0.000000 + -
                                         0.000000
dx =
     2, dy =
                        0.020050 +-
                                         0.000307
dx =
     0, dv =
                        0.000000 + -
                                         0.00000
     1, dy =
dx =
                        0.019829 + -
                                         0.000184
dx = 2, dy =
                        0.000000 +-
                                         0.000000
```

The first line is the name of measurements. Below that, the first column is the arguments of the function. The second and the third columns are the averages and errors.

Complex results will be defined similarly with separated error bars for the real part and imaginary part.

For the detail formula of each measurements, please referee the working notes.

2.5.1 Equal time measurements

There are three groups of equal time measurements. The first group is the measurements that aggregate values from entire lattice. The second group is the autocorrelation functions that average pair of sites within the same distance class. The third group measures the pair susceptibilities.

- 1. Up spin occupancy
- 2. Down spin occupancy
- 3. Potential energy
- 4. Kinetic energy
- 5. Total energy
- 6. Density
- 7. XX Ferromagnetic structure factor
- 8. ZZ Ferromagnetic structure factor
- 9. XX Antiferromagnetic structure factor
- 10. ZZ Antiferromagnetic structure factor
- 11. Equal time Green's function
- 12. Density-density correlation function (up-up)
- 13. Density-density correlation function (up-dn)
- 14. XX Spin correlation function
- 15. ZZ Spin correlation function
- 16. S-wave pair structure factor
- 17. SX-wave pair structure factor
- 18. D-wave pair structure factor
- 19. SXX-wave pair structure factor
- 20. DXX-wave pair structure factor
- 21. PX-wave pair structure factor
- 22. PY-wave pair structure factor
- 23. PXY-wave pair structure factor
- 24. PYX-wave pair structure factor

2.5.2 Unequal time measurements

Every unequal time measurement is a function of imaginary time. The format of array typed measurement with two arguments, space and time, is like

```
G(nx,ny,ti)
dx = 0, dy =
               0
              0.50000 +-
          0
                           0.00000
          1
              0.36076 + -
                           0.00311
              0.27942 +-
          2
                           0.00328
dx = 1, dy = 0
          0 -0.12042 +-
                           0.00162
          1 -0.07183 +-
                           0.00097
            -0.04248 +-
                           0.00133
dx = 2, dy = 0
```

where dx = 0, dy = 0 are labels of space and followed by a list of time label and corresponding values

Some measurements are from Fourier transformation, which are complex. Their format will be like

```
0(
     0.05179 +-
                   0.00270)
                              +i(
                                    0.00000 +-
                                                  0.00000)
1(
     0.05085 + -
                   0.00248)
                              +i(
                                   -0.00061 +-
                                                  0.00055)
     0.04343 +-
                   0.00190)
                                   -0.00067 +-
2(
                              +i(
                                                  0.00078)
3(
     0.03755 +-
                   0.00144)
                                   -0.00080 +-
                                                  0.00051)
                              +i(
     0.03004 +-
                   0.00099)
                                   -0.00090 +-
                                                  0.00076)
4(
                              +i(
5(
     0.02512 +-
                   0.00092)
                              +i(
                                   -0.00100 +-
                                                  0.00048)
6(
     0.02152 +-
                   0.00074
                              +i(
                                   -0.00134 +-
                                                  0.00031)
7(
     0.01829 +-
                   0.00072)
                              +i(
                                   -0.00087 +-
                                                  0.00055)
8(
     0.01499 + -
                   0.00066)
                              +i(
                                   -0.00051 +-
                                                  0.00055)
9(
     0.01332 +-
                   0.00065)
                              +i(
                                   -0.00042 +-
                                                  0.00035)
```

The numbers inside the first parenthesis are the average and error of the real part and the numbers in the second parenthesis are for imaginary part.

- 1. Unequal time Green's function: G(dx, dy, t)
- 2. Discrete cosine transformed G(dx, dy, t): G(qx, qy, t)
- 3. Fourier Transformed G(dx, dy, t): G(dx, dy, w)
- 4. Fourier Transformed G(qx, qy, t): G(qx, qy, w)
- 5. chi function: $\chi(dx, dy, t)$
- 6. Discrete cosine transformed $\chi(dx, dy, t)$: $\chi(qx, qy, t)$
- 7. Fourier Transformed $\chi(dx, dy, t)$: $\chi(dx, dy, w)$

- 8. Fourier Transformed $\chi(qx,qy,t)$: $\chi(qx,qy,w)$
- 9. S-Wave pair structure factor, vertex and nonvertex
- 10. SX-Wave pair structure factor, vertex and nonvertex
- 11. D-Wave pair structure factor, vertex and nonvertex
- 12. SXX-Wave pair structure factor, vertex and nonvertex
- 13. DXX-Wave pair structure factor, vertex and nonvertex
- 14. PX-Wave pair structure factor, vertex and nonvertex
- 15. PY-Wave pair structure factor, vertex and nonvertex
- 16. PXY-Wave pair structure factor, vertex and nonvertex
- 17. PYX-Wave pair structure factor, vertex and nonvertex
- 18. FTed S-Wave pair structure factor, vertex and nonvertex
- 19. FTed SX-Wave pair structure factor, vertex and nonvertex
- 20. FTed D-Wave pair structure factor, vertex and nonvertex
- 21. FTed SXX-Wave pair structure factor, vertex and nonvertex
- 22. FTed DXX-Wave pair structure factor, vertex and nonvertex
- 23. FTed PX-Wave pair structure factor, vertex and nonvertex
- 24. FTed PY-Wave pair structure factor, vertex and nonvertex
- 25. FTed PXY-Wave pair structure factor, vertex and nonvertex
- 26. FTed PYX-Wave pair structure factor, vertex and nonvertex

3 Advanced usages

To assist in creating new simulations, ¹⁸ QUEST provides several simple mechanisms to

- 1. Add new configuration parameters,
- 2. Create new lattice geometries,
- 3. Add new measurements.

This section will introduce how to apply them.

¹⁸The new simulations do not include creating new models. Currently, QUEST is pretty much limited in Hubbard's model.

3.1 Add new configuration parameters

Section 2.1 lists the basic input parameters, which are enough for current programs. However, for new simulations, additional parameters may be required. QUEST allows new parameters to be specified through current configuration system. For example, in the diluted lattice, in which some sites are randomly removed, the percentage of removal sites can be specified in the input file

```
rmv_ratio = 0.1
```

just like the other parameters.

To add new configuration parameters, user/developer need to edit a file called config.def. When the subroutine DQMC_Config_Read is called, it will first searches config.def for the definition of parameters. If the file exists, then the program will use the parameter defined in the file. Otherwise, it uses default parameter set, as described in section 2.1.

A parameter in config.def is defined by a quintuple:

```
{name, type, isArray, printed, default}
```

where

name: a string, maximum characters 30, specifying the name of the parameter.

type: an integer specifying the data type of the parameter.

isArray: a boolean {T,F} specifying whether the parameter is an array. Note, when type is a string, isArray cannot be T.

printed: a boolean {T,F} specifying whether the parameter will be printed in the output.

default: a string, maximum characters 30, specifying the default value of the parameter.¹⁹ QUEST will convert it to a proper data type based on the "type" tuple.

Each tuple is separated by spaces. For example, the parameter rmv_ratio can be defined in the config.def as

```
rmv_ratio 1 F T 0.1
```

The file config.def should be placed in the directory where the executable is.

3.2 Square lattice geometry file with comments.

#NDIM

2

Dimension of the lattice. 3 for cubic, 2 for square, triangular, etc

¹⁹Note, even a parameter is an array, its default value can be only set by one number.

#PRIM

1.0 0.0

0.0 1.0

Primitive lattice vectors. One can give a full 3×3 matrix for #PRIM even if #NDIM; 3. If you do that, QUEST only reads the upper left #NDIM \times #NDIM block.

```
#ORB
```

s0 0.0 0.0 0.0 #0

This is what allows QUEST to do geometries like the honeycomb lattice which requires a basis. #ORB has one line for each atom in the basis. The first entry is a string which serves as a label for the atom. The next three entries in the line are the position of the atom in the unit cell. QUEST authomatically assignes a number to each atom (ie to each line in the #ORB section of the .geom file) beginning with zero. In the #ORB section QUEST demands three dimensional objects.

#HAMILT block of the code defines the hoppings, on-site energies, and interaction strenghts. The convebtion for the lines in #HAMILT is following: Each line begins with two entries which are the (automatically assigned) atom numbers from #ORB. For problems without a basis these will just be '0.0'. To include a hopping, the next three entries in the line should be the direction to the neighboring site. QUEST authomatically makes \hat{H} Hermitian, so each pair of connected sites requires only one line. The next two entries are the hopping values for the up and down electrons, which are allowed to be different in QUEST. The final entry is U, which should be set to zero for lines defining hopping. To include a U value, use '0.0 0.0 0.0' for the neighboring site inputs and insert a value in the last (eighth) entry.

If you have a geometry with a basis (more than one line in #ORB) the different atoms can be assigned different site energies and interactions.

Here is example for the rectangular geometry:

#HAMILT

0 0 1.0 0.0 0.0 1.0 1.0 0.0 0 0 0 0.0 1.0 0.0 2.2 2.2 0.0 0.0 0.0 0.0 0.0 0.5 0.5 4.0

The first two lines are the hoppings in the x and y directions. The third line is an on-site energy 0.5, and is the same for the eup and down species. (the sign convebtion for the site energies is $+\epsilon n_i$, so that 0,5 in #HAMILT corresponds to -0.5 in the Chemical potential. The third line also sets U=4. (If you want, you can separate this into two distinct lines)). We again note that in #HAMILT it is required to supply a three component vector as the pointer to the site, even if #NDIM is two.

Note: The chemical potential defined in input is a **global** chemical potential which applies to all sites in the lattice. The on-site energies in #HAMILT allow for different chemical potentials on different atoms. Thus there is a slight redundancy in the code. Rather than having a global chemical potential one could shift all the site energies. If you do not specify chemical potentials in input, they are set, by default, equal to zero.

There is some redundancy in the way QUEST knows the geometry is rectangular. If the defined supercell in #SUPER has different diagonal entries, then QUEST knows not to assume x and y directions are equivalent. Likewise, a different entry for t_x and t_y will automatically be flagged by QUEST (Finally, the strange string label in #ORB about which we have been so silent can also be used to distinguish different types of atoms and can break symmetries. If you had a three band

Cu-O model of the cuprates you could, for example, make the oxygen atoms along the x and y bonds have distinct labels and hence different entries in #HAMILT)

But more fundamentally, QUEST knows about the symmetries of the lattice from the #SYMM block in the .geom file, which is the last one we shall discuss. In square.geom this looks like,

#SYMM

```
d 0.0 0.0 0.0 1.0 0.0 0.0
d 0.0 0.0 0.0 0.0 1.0 0.0
c4 0.0 0.0 0.0 0.0 0.0 1.0
```

QUEST supports 3 types of symmetry definitions. 'cn', where 'n' is an integer, tells QUEST the lattice is symmetric under rotations by $2\pi/n$. The six numbers following by 'cn' specify the three cartesian coordinates of a point belonging to the axis, following by the axis direction in the cartesian coordinates. In this case 'c4' is $\pi/2$ and indicates the x and y directions are equivalent. the point belonging to the axis is the origin '0.0 0.0 0.0' and the axis is the z direction '0.0 0.0 1.0'.

The symmetry 'd' is a mirror plane symmetry. It too is followed by six numbers. The first three are the cartesian coordinates of a point in the plane, and the final three are the cartesian coordinates of the normal to the plane.

Finally, 'I' is used for inversion symmetry. It is followed by three numbers, the cartesian coordinates of the inversion point.

In specifying #SYMM you must list all three components of teh vectors even if the lattice is two dimensional.

3.3 Create new lattice geometries

In QUEST, geometry dependent variables are placed together in a derived data type Struct. To create a new lattice geometry, one needs to fill out the data fields in Struct. There are several ways to achieve this; each has its advantages and drawbacks.

- 1. Write a program to fill out the data fields: most efficient way in execution, but need to write programs for different geometries.
- 2. Use primitive cell definition as input: ²⁰ Most compact representations of general geometry, but needs to understand the definitions of primitive cell.
- 3. Input data for each data field from files: Most easy way to create a new lattice geometry, but less flexibilities and less efficient.

In this section, we will illustrate the data fields in Struct and the file formats for method 3. Note not all the data fields are essential for simulations. Some of them are just for a particular measurements. When lacking one or some of the fields, QUEST will skip the corresponding measurements.

3.3.1 Struct

Derived data type Struct is defined as

 $^{^{20}}$ This feature is still underdevelopment.

```
type Struct
   integer :: nSite
                                   ! number of sites
   character(gname_len):: name
                                   ! name of the structure
   integer, pointer
                      :: dim(:)
                                 ! dim of the geometry
   integer :: n_t
                                   ! number of hopping types
   type(CCS):: T
                                   ! hopping matrix
   integer :: n_b
                                   ! number of neighbors types
  type(CCS):: N
                                   ! neighborhood matrix
   integer :: nClass
                                   ! number of distance classes.
   integer, pointer :: D(:,:)
                                   ! distance classes
   integer, pointer :: F(:)
                                   ! counts for each dist class.
   character(label_len), pointer :: &
                         label(:) ! label for distant class.
   integer :: nGroup
                                   ! number of diff singletons.
   integer, pointer :: map(:)
                                   ! site classification
  real(wp), pointer :: W(:,:)
                               ! wave functions
   integer :: nWave
   character(label_len), pointer :: &
                          wlabel(:)! label for wave functions.
  real(wp), pointer :: P(:)
                                ! phase assignment
  real(wp), pointer :: FT(:,:) ! Fourier Transform matrix
  logical::checklist(N_CHECKLIST) ! flags
end type Struct
```

Here are the detail comments for each field.

- Sites must be numbered from 1, with continuous numbering.
- String name is of length 80 characters, used in display.
- Vector dim is used to hold dimension parameters. For example, for a two dimensional square lattice, dim=(nx,ny), where nx and ny are the number of sites in x direction and in y direction. This field can be of arbitrary length. QUEST does not use this field directly.
- Hopping matrix T stores the indices of hopping parameter t for adjacent sites.²¹ It is normally a sparse matrix; and therefore is represented in the Compressed Column Storage (CCS) format.²² Field n_t is the number of different hopping parameters.
- Neighboring matrix N is similar to T, and is also represented in the CCS format. It is used in pair measurements, for which the link indices should be consistent with the wave function W. Field n_b is the number of different links.

²¹Two sites i, j are called adjacent if electrons can hop from site i to site j.

²²The detail of CCS format can be found in http://www.netlib.org.

- The distance classification is represented by D, F and label. Two pairs of sites are in the same class if they are translation/rotation invariant. The number of classes is specified by nClass. The class index is also started from 1. The number of pairs in each class is stored in F. String array label gives a label for each class, used in output. Matrix D records the classification for each pair of site. Number in D(i,j) denotes the class index for site i and j.
- The vector map classifies sites. Site i and site j are in the same class if they have the same physical parameters, like U and μ . The number nGroup denotes the number of site classes.
- Matrix W defines wave functions for the pair measurements, which are related to various spherical harmonic functions. The number of functions is specified by nWave. Each function has n_b elements, and is stored in a column of W matrix.
- The phase assignment vector P gives each site $\{+1, -1\}$ so that adjacent sites have opposite phases. This is used in spin correlation measurements, and in Green's function calculation when $\mu = 0$.
- Matrix FT is the Fourier transformation matrix for distance classes, which is used in time dependent measurements.
- Vector checklist is a set of flags that indicate which data fields are assigned. The flags include

```
STRUCT_INIT = 1
STRUCT_DIM = 2
STRUCT_ADJ = 3
STRUCT_CLASS = 4
STRUCT_WAVE = 5
STRUCT_NEIG = 6
STRUCT_PHASE = 7
STRUCT_FT = 8
```

3.3.2 Files for geometry definition

The file name of geometry definition is specified in the configuration file with parameter name gfile. For example,

```
gfile = strip.def
```

tells QUEST to use the file strip.def as geometry definition.

The format of the geometry definition is similar to the configuration file.

- 1. The symbol # is used to start a comment.
- 2. The data fields are set by the assignment

```
data_field_name = data_field_value
```

3. Some fields depend on the other. Depended fields should be declared first.

$$\begin{split} & T \rightarrow \text{nSite} \\ & B \rightarrow \text{nSite} \\ & D \rightarrow \text{nSite} \\ & FT \rightarrow \text{nClass} \\ & W \rightarrow \text{n_b} \text{ and nWave} \end{split}$$

- 4. Not every field in Struct need be defined in the file. For example, vector F and map will be derived from matrix D.
- 5. Array type assignment, including matrix or vector, is different from that in a configuration file. The right-hand-side of assignment should be the number of how many lines to read. For example

D = 64

means there are 64 immediate lines to read for matrix D. Those lines are called *content lines*. Note NO empty lines are allowed between content lines.

6. The format for a content line of a matrix is

i j value

where i is the row index, j is the column index, and value is the (i, j) element of the matrix.

7. Format for a content line of a vector is

value

Those values should be ordered sequentially, since indices are assumed to be implicitly embedded.

An example of how to create new geometry using method 3 can be found in the example program in EXAMPLE\gemo.

3.4 Add new measurements

New measurements need be made through programming in QUEST. Several subroutines can be used to create new measurements. The standard procedure to add a new measurement includes three steps.

- 1. Measuring.
- 2. Binning.
- 3. Postprocessing. (Statistics, Fourier transformation, output.)

3.4.1 Measuring

Several components may be needed to create a new measurements.

1. **Equal time Green's function**: Equal time Green's function is defined in the module dqmc_gfun, which will be initialized automatically when the subroutine DQMC_Hub_Init is called. Suppose Hub is typed Hubbard, the major data type of entire simulation. The Green's function matrices, spin-up and spin-down, can be obtained from

```
Hub%G_up%G
Hub%G_dn%G
```

And the signs of their determinants are recorded in

```
Hub%G_up%sgn
Hub%G_dn%sgn
```

- 2. Unequal time Green's function: Unequal time Green's function, denoted G_{ρ}^{τ} , $\rho \in \{\uparrow, \downarrow\}$, is defined in the module dqmc_gtau. Unlike equal time Green's functions, G^{τ} are not essential in the simulation. Therefore, user needs to initialize it by calling DQMC_Gtau_Init explicitly. The construction of G^{τ} can be made in two ways. The first way is to call DQMC_Gtau_Big, which returns entire G_{ρ}^{τ} . The second method is to invoke DQMC_MakeGtau, which returns block submatrices of G^{τ} . Since the signs of unequal time Green's functions are the same as the equal times, one can obtain the signs from Hub%G_up%sgn, Hub%G_dn%sgn.
- 3. Parameters of Hamiltonian: Parameters, such as t, μ and U, are stored in the data type Hubbard. The access is straightforward.
- 4. **Geometry related information**: Geometry information, such as hopping matrix, can be obtained from the data type struct, which is introduced in section 3.2.

3.4.2 Binning

In order to reduce correlation and bias, measurements in QUEST need be grouped into bins. Currently, equal divided binning strategy is used, which means the measurements are evenly divided by the total number of bins. Measurements in the same bin are averaged. The total number of bins are stored in the variable nbin.

3.4.3 Statistics

There are two special properties for the physical measurements produced by DQMC method.

- 1. The distribution is not normal.
- 2. Measurements are weighted with signs of the determinants of Green's functions, for which the average needs be normalized by the average of signs.

QUEST uses *Jackknife* resampling technique in error estimation. Two subroutines are provided to perform the statistics: DQMC_JackKnife and DQMC_SignJackKnife. The former is for error estimation of signs; the latter is used for other measurements. Those subroutines are defined in module dqmc_util.

3.4.4 Fourier transformation

For unequal time measurements, there are two possible Fourier transformations to be applied: transformation on the real space and transformation on the time domain. For the transformation on real space, since it is geometry dependent, a transformation matrix FT, defined in Struct, is required.²³ Once the matrix is available, the transformation is just a matrix-matrix multiplication. In the module dqmc_tdm, QUEST provides a subroutine DQMC_DCT_Space for the space transformation.

The Fourier transformation on the time domain is an integration. The numerical procedure is

- 1. Refine the time grid.
- 2. Interpolate the refined data points.
- 3. Integrate on the interpolated data with Fourier coefficients.

In step 1, QUEST evenly subdivides the time domain by the given parameter nitvl. In step 2, QUEST uses spline interpolation, which is supported by the subroutine DQMC_Spline. Step 3 requires an additional Fourier matrix, which can be generated from the subroutine DQMC_Make_FTM. The entire procedure is coded in the subroutine DQMC_FT_Time.

3.4.5 Printing

QUEST has two subroutines that prints out arrays of numbers. Subroutine DQMC_Print_RealArry prints out an array of real numbers; DQMC_Print_ComplexArray prints out an array of complex numbers. The title of the measurements and the labels of each array items are required for those two functions.

A Example programs

Eight example programs are included in the EXAMPLE directory. Here is an short introduction for each of them.

A.1 test

Program test demonstrates the simplest usage of the QUEST. Besides the timing commands, there is only one line in the program, which runs DQMC simulation on a two dimensional periodic rectangular lattice. In spite of its simplicity, this program can be configured dynamically for different lattice size, Hubbard parameters and execution iterations. Four sample configurations are accompanied within this program, as showing below.

Configuration	Lattice size	Time slice	Running time	
small.in	4×4	12	2	second
median.in	8×8	48	198	second
large.in	16×16	96	15,629	second
extra_large.in	32×32	192	unknown	

Their execution time also presented. This result is obtained by using checkerboard method, 1000 warmup steps and 5000 measurement steps, with Intel MKL BLAS/LAPACK library, on Intel Core 2 Duo 2.4G processors (but only run in one core).

²³see section 3.2 for more details.

A.2 verify

The verify program examines the correctness of the execution results of two special cases, single site (t=0) and no Coulomb interaction (U=0). Each test case runs through 9 configurations.

The correctness of those results are checked against the theoretical results. Statistically, 63.2 percent of the computed results are expected to have errors smaller than one standard error, and 86.5 percent of results should be within two standard errors. The verification program runs 1000 warm-up sweep and 5000 measurement sweep for t=0 cases, which gives error bars of 0.33% on the energy and less than 0.22% on the spin correlation SpinXX.

A.3 tdm

This example program demonstrates how unequal time measurements can be performed. This program also shows the flexibility of using the subroutines in QUEST. Four sample configurations, the same those in test, are available for this program. The time dependent measurements are made every 10 measurement sweeps. Their execution time are summarized in Table A.3.

Configuration	Lattice size	Time slice	Running	g time
small.in	4×4	12	3	second
median.in	8×8	48	266	second
large.in	16×16	96	19446.18	second
extra_large.in	32×32	192	unkno	own

A.4 Others

- parallel: This program uses MPI to parallelize the measurement steps. The compilation of this program requires mpif90. The base compiler of mpif90 should be the same as the one used in compiling library.
- wrap: This directory contains a module dqmc_wrapper and a program tw. The module wraps all computation components of QUEST into several simpler functions. The program tw, test wrapper, can perform equal time and unequal time measurements.
- sheet: This directory contains a module and programs for the multilayer lattice. Module dqmc_sheet defines the multilayer geometry. Program meas1 and meas2 provides new measurements for the multilayer structure.
- **geom**: The program inside this directory tests the general geometry method 3 mentioned in section 3.2.
- DCA: This is a project that interfacing QUEST with other programs. Basically, the entire project uses DQMC as a DCA solver. This demonstrates how to use QUEST as a library in other programs.