# Documentation for the General QMC code

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## 1 Theory part

#### 1.1 Definition of the physical Hamiltonian and its implementation

The physical Hamiltonians that we can simulate have the general form:

$$\mathcal{H} = \sum_{s=1}^{N_{fl}} \sum_{x,y} c_{xs}^{\dagger} M_{xy} c_{ys} - \sum_{k=1}^{M} U_k \left[ \sum_{s=1}^{N_{fl}} \sum_{x,y} \left( c_{xs}^{\dagger} T_{xy}^{(k)} c_{ys} - \alpha_k \right) \right]^2 . \tag{1}$$

The indices x, y are multi-indices that label sites and spin states:  $x = (i, \sigma)$ , where  $i = 1, \dots N_{sites}$  and  $\sigma = 1, \dots N_{sun}$ , so

$$\sum_{x,y} \equiv \sum_{i=1}^{N_{sites}} \sum_{\sigma=1}^{N_{sun}} . \tag{2}$$

Note, that we introduced two different labels for the number of spin states (flavours):  $N_{fl}$  and  $N_{sun}$ . The number of correlated sites which is a subset of all sites, is labelled by M ( $M \le N_{sites}$ ). Let us further define  $N_{dim} = N_{sun}N_{sites}$  such that the matrices  $\boldsymbol{M}$  and  $\boldsymbol{T}^{(k)}$  are of dimension  $N_{dim} \times N_{dim}$ .

I suggest to use a more intuitive notation and to label the hopping matrix by T and the interaction matrix by V:

$$\mathcal{H} = \sum_{s=1}^{N_{fl}} \sum_{x,y} c_{xs}^{\dagger} T_{xy} c_{ys} - \sum_{k=1}^{M} U_k \left[ \sum_{s=1}^{N_{fl}} \sum_{x,y} \left( c_{xs}^{\dagger} V_{xy}^{(k)} c_{ys} - \alpha_k \right) \right]^2 , \tag{3}$$

## 1.2 Structure of the matrices T and $V^{(k)}$ and their implementation

In general, the matrices  $\mathbf{T}$  and  $\mathbf{V}^{(k)}$  are sparse matrices. This property is used to minimize computational cost and storage requirements. In the following, we discuss the implementation of the matrix representation  $\mathbf{V}^{(k)}$  of the interaction operator. The same applies for the hopping matrix  $\mathbf{T}$ . We have

$$V_{xy}^{(k)} \neq 0$$
 only if  $x, y \in [z_1, \dots z_{N_{eff}}]$ . (4)

We define the projection matrices  $\mathbf{P}^{(k)}$  of dimension  $N_{eff}^{(k)} \times N_{dim}$ :

$$P_{i,z}^{(k)} = \delta_{z_i,z} \,, \tag{5}$$

where  $i \in [1, \dots, N_{eff}^{(k)}]$  and  $z \in [1, \dots, N_{dim}]$ . Evidently,  $\mathbf{P}^{(k)}$  picks out the non-vanishing entries of  $\mathbf{V}^{(k)}$ , which are contained in the  $(N_{eff}^{(k)} \times N_{eff}^{(k)})$  - dimensional matrix  $\mathbf{O}_V^{(k)}$ :

$$V^{(k)} = P^{(k)T} O_V^{(k)} P^{(k)},$$
 (6)

and

$$V_{xy}^{(k)} = P_{ix}^{(k)} \left[ O_V^{(k)} \right]_{ij} P_{jy}^{(k)} = \sum_{i,j}^{N_{eff}^{(k)}} \delta_{z_i,x} \left[ O_V^{(k)} \right]_{ij} \delta_{z_j,y} . \tag{7}$$

To set the interaction part, we therefore have to specify the matrix elements  $\left[O_V^{(k)}\right]_{ij}$ , the set  $[z_1, \cdots z_{N_{eff}}]$ , and the values  $U_k$  and  $\alpha_k$ .

# 2 Tutorial to set up the Hubbard model

The SU(2) symmetric Hubbard model is given by

$$\mathcal{H} = -t \sum_{\langle i,j \rangle, \sigma} \left( c_{i,\sigma}^{\dagger} c_{j,\sigma} + \text{H.c.} \right) + \frac{U}{2} \sum_{i} \left[ \sum_{\sigma} \left( c_{i\sigma}^{\dagger} c_{i\sigma} - 1/2 \right) \right]^{2} . \tag{8}$$

To bring Eq. (8) in the general form (3), we set

$$N_{fl} = 1$$

$$N_{sun} = 2$$

$$T_{xy} = -t\delta_{\langle i,j\rangle}\delta_{\sigma,\sigma'}$$

$$M = N_{sites}$$

$$U_k = -U/2$$

$$V_{xy}^{(k)} = \delta_{i,j}\delta_{i,k}\delta_{\sigma,\sigma'}$$

$$\alpha_k = 1/(N_{sites}N_{sun})^2 = 1/(2N_{sites})^2.$$
(9)

# 3 Using the code

Example simulation, tutorial: where to find and how to start

## 3.1 Parameter files

describe the input parameters, give sample values for the stabilization parameters

# 3.2 Analysis files

 $how\ the\ analysis\ of\ Monte\ Carlo\ data\ is\ done$ 

# 4 List of files

all files that constitute the code, with a brief description

 $4.1 \quad {\rm cgr1.f90} \ \& \ {\rm cgr2.f90}$ 

Stable computation of the physical single-particle equal time Green function  $G(\tau)$ .

## 4.2 control\_mod.f90

Includes a set of auxiliary routines, regarding the flow of the simulation. Examples are initialization of performance variables, precision tests and controlled termination of the code.

4.3 gperp.f90

## 4.4 Hamiltonian\_Hub.f90

Here, the physical simulation parameters (the model parameters) and the lattice parameters are read in. The lattice, the non-interacting and the interacting part of the Hubbard Hamiltonian are set according to the parameters and the chosen Hubbard-Stratonovich decomposition.

 $4.5 \quad \text{Hop\_mod.f90}$ 

## 4.6 inconfc.f90

The auxiliary-field QMC method is based on a Hubbard-Stratonovich decomposition of the interaction term. This decomposition introduces a space-time array of (discrete) configurations of auxiliary fields, i.e. Ising spins. In this routine, an existing configuration is read in, checks on its dimensionality are made and, in case no prior configuration exists, a random configuration of Ising spins is set up.

## 4.7 main.f90

Top-level part of the program. Here, the program flow which consists of initialization, sweeps through the space-time lattice, and finalizing the program, is coded.

# 4.8 nranf.f90

Auxiliary routine controlling the evaluation of random numbers.

## 4.9 Operator.f90

The algorithms is centered around evaluation of single-particle operators, represented as square matrices. In this routine, the abstract type Operator is defined, including information on the coupling strength, the sites that participate in the single-particle hopping process, and the type of Hubbard-Stratonovich transformation. This routine collects all program relevant operations that are applied to the type Operator, like initializations or multiplications.

#### 4.10 outconfc.f90

Description in plain text:

At the end of the simulation, the last configuration of Hubbard-Stratonovich variables, together with the last set of random numbers is written to the file confout. Prior to the start of a new simulation of the identical space-time dimesion, one can (manually) copy the file confout to the file confin and make the new run use the old configuration.

Doing this saves warmup time compared to a complete random (unphysical) configuration.

Input/output variables

in:
inout:
inout:
out:

Dependencies include: mpif.h modules: Hamiltonian

interfaces:

global variables: ntrot, nsigma

subroutines: MPI\_COMM\_RANK, MPI\_COMM\_SIZE, Get\_seed\_Len, Ranget

Things to check:

Rename subroutine to confout.f90 for consistency

#### 4.11 print\_bin\_mod.f90

Description in plain text: Here the way to write the measure bins to the respective output files is coded. A bin is an average over many individual measurements. The bin defines the unit of Monte Carlo time. Dependencies include: modules: interfaces: Print\_bin Contains subroutines: Print\_bin\_c, Print\_bin\_r, Print\_scal, Print\_bin\_tau Things to check: 4.11.1 Print\_bin\_C Description in plain text: Input/output variables in: Latt, type(Lattice) in: Phase\_bin\_tmp, complex in: File\_pr, character(Len=64) in: nobs, integer inout: Dat\_eq,(:,:,:), complex inout: Dat\_eq0(:), complex out: Dependencies include: mpif.h modules: Lattices\_v3 interfaces: global variables: TYPE(LATTICE), N, listk, b1\_p, b2\_p subroutines: MPI\_COMM\_RANK, MPI\_COMM\_SIZE, MPI\_REDUCE, Fourier\_R\_to\_k Things to check: STATUS(MPI\_STATUS\_SIZE), integer, needed? 4.11.2 Print\_bin\_R Description in plain text: Input/output variables in: Latt, type(Lattice) in: Phase\_bin\_tmp, complex in: File\_pr, character(Len=64) in: nobs, integer inout: Dat\_eq,(:,:,:), real inout: Dat\_eq0(:), real

Dependencies include: mpif.h

out:

modules: Lattices\_v3 interfaces: global variables: TYPE(LATTICE), N, listk, b1\_p, b2\_p subroutines: MPI\_COMM\_RANK, MPI\_COMM\_SIZE, MPI\_REDUCE, Fourier\_R\_to\_k Things to check: STATUS(MPI\_STATUS\_SIZE), integer, needed? 4.11.3 Print\_scal Description in plain text: Input/output variables in: File\_pr, character(Len=64) in: nobs, integer inout: Obs,(:), complex out: Dependencies include: mpif.h interfaces: global variables: subroutines: MPI\_COMM\_RANK, MPI\_COMM\_SIZE, MPI\_REDUCE Things to check: STATUS(MPI\_STATUS\_SIZE), integer, needed? change subroutine name to print\_bin\_scal for consistency 4.11.4 Print\_bin\_tau Description in plain text: Input/output variables in: Latt, type(Lattice) in: Phase\_bin, complex in: File\_pr, character(Len=64) in: nobs, integer in: dtau, real inout: Dat\_tau,(:,:,:), complex inout, optional: Dat0\_tau(:), complex out: Dependencies include: mpif.h modules: Lattices\_v3 interfaces: global variables: TYPE(LATTICE), N, listk, b1\_p, b2\_p subroutines: MPI\_COMM\_RANK, MPI\_COMM\_SIZE, MPI\_REDUCE, Fourier\_R\_to\_k Things to check:

STATUS(MPI\_STATUS\_SIZE), integer, needed?

```
Description in plain text:
module tau_m_mod, with several subroutines
Dependencies
include:
modules: Hamiltonian, Operator_mod, Precdef, Control, Hop_mod
Contains
subroutines: tau_m, propr, proprm1
Things to check:
change file name to tau_m_mod.f90 for consistency
4.12.1 TAU_M
Description in plain text:
Input/output variables
in: nstm, integer
in: nwrap, integer
in: ust(ndim,ndim; nstm,n_fl), complex
in: vst(ndim,ndim,nstm,n_fl), complex
in: dst(ndim,nstm,n_fl), complex
in: GR(ndim,ndim,n_fl), complex
in: phase, complex
in: stab_nt(0:nstm), integer
inout:
inout:
out:
Dependencies
include:
modules:
interfaces: wrapul, cgr2_1, cgr2_2, cgr2
global variables: ndim, n_fl, cone, ltrot
subroutines: obsert, initd, propr, proprm1, wrapur, cgr2_2, Control_Precision
Things to check:
cone?
4.12.2 propr
Description in plain text:
Input/output variables
in: nt
inout: Ain(ndim,ndim,n_fl)
inout:
out:
Dependencies
include:
```

4.12 tau\_m.f90

modules:

```
interfaces:
global variables: ndim, n_fl, op_v, nsigma, Phi, type
subroutines: Hop_mod_mmthr, Op_mmultR
Things to check:
4.12.3 proprm1
Description in plain text:
Input/output variables
in: nt
inout: Ain(ndim,ndim,n_fl)
inout:
out:
Dependencies
include:
modules:
interfaces:
global variables: ndim, n_fl, op_v, nsigma, Phi, type
subroutines: Hop_mod_mmthl, Op_mmultL
```

4.13 UDV\_WRAP.f90 Description in plain text: UDV\_Wrap\_mod is a module file, containing subroutines on the stabilization of matrix computations. Dependencies modules: MyMats, Files\_mod Contains subroutines: UDV\_Wrap\_Pivot, UDV\_Wrap Things to check: change file name to UDV\_Wrap\_mod.f90 for consistency 4.13.1 UDV\_Wrap\_Pivot Description in plain text: Input/output variables in: A(:,:), complex in: ncon, integer in: n1, integer in: n2, integer inout: U(:,:), complex inout: V(:,:), complex inout: D(:), complex out: Dependencies include: modules: interfaces: global variables: subroutines: UDV\_Wrap, MMULT, Compare Things to check: 4.13.2 UDV\_Wrap Description in plain text: Input/output variables in: A(:,:), complex in: ncon, integer inout: U(:,:), complex inout: V(:,:), complex inout: D(:), complex out: Dependencies include: mpif.h

modules:
interfaces:
global variables:

subroutines: MPI\_COMM\_SIZE, MPI\_COMM\_RANK, QR, SVD, MMULT

Things to check:

STATUS(MPI\_STATUS\_SIZE), integer: not used

## 4.14 upgrade.f90

Description in plain text:

log, logical (reserved name)

alpha: both a local and a global variable. CHECK!!

for each point in the space-time lattice, i.e. one Hubbard-Stratonovich Ising spin after the other. In this routine, an update (i.e. a spin flip) is accepted or rejected. The decision is made using the Metropolis method of importance sampling. Input/output variables in: N\_op, integer in: nt, integer in: OP\_dim, integer inout: GR(ndim,ndim,n\_fl), complex inout: Phase, complex out: Dependencies include: modules: Hamiltonian, Random\_wrap, Control, Precdef interfaces: global variables: ndim, n\_fl, op\_v, nflipl, Phi, n\_non\_zero, Gaml, P, nsigma, g, alpha, type, E subroutines: zgemm, control\_upgrade Things to check: nranf, integer, external (where is external fct. defined)

The update of the Hubbard-Stratonovich configuration is done sequentially

## 4.15 wrapgrdo.f90

Description in plain text:

Single-particle equal-time Green functions are the central object of the code.

The physical single-particle equal-time Green function  $G(\tau)$  is updated in wrapgrdo.f90 (down propagation, from  $\tau$ ).

The update is sequentially, over all (interacting) lattice sites or lattice bonds.

Input/output variables

in: ntau, integer

inout: gr (ndim,ndim,n\_fl), complex

inout: phase, complex

out:

Dependencies

include:

modules: Hamiltonian, MyMats, Hop\_mod

interfaces: upgrade

global variables: op\_v, phi, nsigma, ndim, n\_fl

subroutines: Hop\_mod\_mmthl, Hop\_mod\_mmthr\_m1, Op\_Wrapdo, Upgrade

## 4.16 wrapgrup.f90

Description in plain text:

Single-particle equal-time Green functions are the central object of the code.

The physical single-particle equal-time Green function  $G(\tau)$  is updated in wrapgrup.f90 (up propagation, from  $\tau$ ) to  $\tau$ .

The update is sequentially, over all (interacting) lattice sites or lattice bonds.

Input/output variables

in: ntau, integer

inout: gr (ndim,ndim,n\_fl), complex

inout: phase, complex

out:

Dependencies

include:

modules: Hamiltonian, Hop\_mod

interfaces: upgrade

global variables: op\_v, phi, nsigma, ndim, n\_fl

subroutines: Hop\_mod\_mmthr, Hop\_mod\_mmthl\_m1, Op\_Wrapup, Upgrade

## 4.17 wrapul.f90

Description in plain text: To stabilize the simulation at the time slice  $\hat \xi_2=i n_{stab}$ , the Green function has to be recomputed regularly, based on the stable matrices at an earlier stabilization point,  $\hat \xi_1=(i-1) n_{stab}$ . These stable matrices result from a singular-value-decomposition of the propagation matrix. They are computed in wrapul.f90 (down propagation).

Input/output variables
in: ntau1, integer
in: ntau, integer
inout: ulup (ndim,ndim,n\_fl), complex
inout: dlup (ndim,n\_fl), complex
inout: vlup (ndim,ndim,n\_fl), complex
out:

Dependencies include:

modules: Hamiltonian, Hop\_mod, UDV\_Wrap\_mod

interfaces:

global variables: ndim, n\_fl, Op\_V, Phi, nsigma,

subroutines: initd, Op\_mmultL, Hop\_mod\_mmthl, mmult, UDV\_Wrap

## 4.18 wrapur.f90

Description in plain text: To stabilize the simulation at the time slice  $\hat{2}=i n_{stab}$ , the Green function has to be recomputed regularly, based on the stable matrices at an earlier stabilization point,  $\hat{1}=(i-1) n_{stab}$ . These stable matrices result from a singular-value-decomposition of the propagation matrix. They are computed in wrapur.f90 (up propagation).

Input/output variables
in: ntau1, integer
in: ntau, integer
inout: ur (ndim,ndim,n\_fl), complex
inout: dr (ndim,n\_fl), complex
inout: vr (ndim,ndim,n\_fl), complex
out:

 ${\tt Dependencies}$ 

include:

modules: Hamiltonian, Hop\_mod, UDV\_Wrap\_mod

interfaces:

global variables: ndim, n\_fl, Op\_V, Phi, nsigma,

subroutines: initd, Op\_mmultR, Hop\_mod\_mmthr, mmult, UDV\_Wrap

```
Description in plain text:

Input/output variables
in:
inout:
inout:
out:

Dependencies
include:
modules:
interfaces:
global variables:
subroutines:
```

## 5 Module Hamiltonian

Detailed description of the module Hamiltonian since it will be modified by the users The module contains the following subroutines:

#### 5.1 ham\_set

Things to check:

It calls the subroutines

- ham\_latt
- ham\_hop
- ham\_v

It reads in the file

• parameters

It sets the variables ltrot,n\_fl,n\_sun. If compiled as a MPI-program, it broadcasts all variables that define the lattice, the model and the simulation process.

#### 5.2 ham latt

It sets the lattice, by calling the subroutine

• make\_lattice(l1\_p, l2\_p, a1\_p, a2\_p, latt)

#### 5.3 ham\_hop

Setup of the hopping amplitudes between the vertices of the graph (lattice sites and unit cell orbitals). It calls the subroutines

- op\_make(op\_t(nc,n),ndim
- $op_set(op_t(nc,n))$

#### 5.4 ham\_v

It calls the subroutines

- $op_make(op_v(i,nf),1)$
- $op_set(op_v(nc,nf))$

#### $5.5 \quad s0(n,nt)$

It is defined as s0(n, nt) = 1.d0. Why? It is superfluous.

#### 5.6 alloc\_obs(ltau)

Allocation of equal time and time-resolved quantities.

#### 5.7 init\_obs(ltau)

Initializes equal time and time-resolved quantities with zero.

#### 5.8 obser(gr,phase,ntau)

Includes the definition of all equal-time observables (scalars and correlation functions) that are built from the single-particle Green function based on Wick's theorem.

#### 5.9 pr\_obs(ltau)

Output (print) of the observables.

#### $5.10 \quad \text{obsert(nt,gt0,g0t,g00,gtt,phase)}$

Includes the definition of time-resolved observables that are built from the time-resolved single-particle Green function based on Wick's theorem.

## 6 Tutorial to set up a lattice

#### 7 Installation

#### 7.1 Dependencies

which software and libraries are needed and which version

- libraries: LAPACK, BLAS, EISPACK, NAG They are included in the package, but NAG is not public-domain (?)
- tools: cmake
- compiler: gfortran or ifort

#### 7.2 Build the GQMC program from source code

configuration, compile and Installation In the top level directory, where the README file resides, do:

```
mkdir build
cd build
cmake ..
make
```

# 8 Reference manual

# 9 License

Use of the GQMC code requires citation of the paper  $\dots$  The GQMC code is available for academic and non-commercial use under the terms of the license  $\dots$  For commercial licenses, please contact the GQMC development team.

# 10 ideas

FAQ, walkthroughs,