

# Documentation for the General QMC code

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## 1 Using the code

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

### 1.1 Parameter files

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

### 1.2 Analysis files

*how the analysis of Monte Carlo data is done*

## 2 List of files

*all files that constitute the code, with a brief description*

2.1 cgr1.f90 & cgr2.f90

2.2 control\_mod.f90

2.3 gperp.f90

2.4 Hamiltonian\_Hub.f90

2.5 Hop\_mod.f90

2.6 inconfc.f90

2.7 main.f90

2.8 nranf.f90

2.9 Operator.f90

2.10 outconfc.f90

2.11 print\_bin\_mod.f90

2.12 tau\_m.f90

2.13 truncation.f90

2.14 UDV\_WRAP.f90

- 2.15 upgrade.f90
- 2.16 wrapgrdo.f90
- 2.17 wrapgrup.f90
- 2.18 wrapul.f90
- 2.19 wrapur.f90

### 3 Module Hamiltonian

*Detailed description of the module Hamiltonian since it will be modified by the users*

The module contains the following subroutines:

#### 3.1 ham\_set

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.

#### 3.2 ham\_latt

It sets the lattice, by calling the subroutine

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

#### 3.3 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))

#### 3.4 ham\_v

It calls the subroutines

- op\_make(op\_v(i,nf),1)
- op\_set( op\_v(nc,nf) )

#### 3.5 s0(n,nt)

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

### 3.6 alloc\_obs(ltau)

Allocation of equal time and time-resolved quantities.

### 3.7 init\_obs(ltau)

Initializes equal time and time-resolved quantities with zero.

### 3.8 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.

### 3.9 pr\_obs(ltau)

Output (print) of the observables.

### 3.10 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.

## 4 Necessary background information

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

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

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

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

$$\sum_{\mathbf{x}, \mathbf{y}} \equiv \sum_{i=1, j=1}^{N_{sites}} \sum_{\sigma=1, \sigma'=1}^{N_{sun}}. \quad (2)$$

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  $M$  ( $M \leq N_{sites}$ ).

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

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

### 4.2 Which information does the type *operator* contain?

## 5 Tutorial to set up the Hubbard model

The  $SU(2)$  symmetric Hubbard model is given by

$$\mathcal{H} = -t \sum_{\langle i, j \rangle, \sigma} \left( c_{i, \sigma}^\dagger c_{j, \sigma} + \text{H.c.} \right) + \frac{U}{2} \sum_i \left[ \sum_\sigma (c_{i\sigma}^\dagger c_{i\sigma}) - 1 \right]^2. \quad (4)$$

To bring Eq. (4) in the general form (3), we define:

$$\begin{aligned}
N_{fl} &= 1 \\
N_{sun} &= 2 \\
T_{\mathbf{x}\mathbf{y}} &= -t\delta_{\langle i,j \rangle}\delta_{\sigma,\sigma'} \\
M &= N_{sites} \\
U_k &= -U/2 \\
V_{\mathbf{x}\mathbf{y}}^{(k)} &= \delta_{i,j}\delta_{i,k}\delta_{\sigma,\sigma'} \\
\alpha_k &= 1/(N_{sites}N_{sun})^2.
\end{aligned} \tag{5}$$

## 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 ... The GQMC code is available for academic and non-commercial use under the terms of the license ... For commercial licenses, please contact the GQMC development team.

## 10 ideas

FAQ, walkthroughs,