# User Guide

# $\mathcal{H}\Phi$ Version 2.0.4

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April, 3, 2017

# Contents

1	$\mathbf{W}\mathbf{h}$	at is $\mathcal{H}\Phi$ ?	1
	1.1	What is $\mathcal{H}\Phi$ ?	1
		1.1.1 License	2
		1.1.2 Copyright	2
		1.1.3 Contributors	2
	1.2	Operating environment	3
2	Hov	v to use $\mathcal{H}\Phi$	4
	2.1	Prerequisite	4
	2.2	Installation	4
		2.2.1 Using HPhiconfig.sh	5
		2.2.2 Using cmake	6
	2.3	Directory structure	7
	2.4	Basic usage	8
		2.4.1 Standard mode	8
		2.4.2 <i>Expert</i> mode	9
		2.4.3 Setting the process number for MPI/hybrid parallelization	10
		, , , -	10
3	Tut	orial	11
	3.1	Standard mode	11
			11
		9	18
	3.2		19
		•	19
			19
			21
		3.2.4 Setting output components	23
		3.2.5 Running	24
	3.3		25
	3.4		25
4	File	especification	26
•	4.1	Input files for Standard mode	
	1.1	<del>-</del>	27

ii Contents

	4.1.2	Parameters for the lattice	28
	4.1.3	Parameters for conserved quantities	32
	4.1.4	Parameters for the Hamiltonian	32
	4.1.5	Parameters for the numerical condition	36
	4.1.6	Parameters for the dynamical Green's function	38
4.2	Input f	files for Expert mode	40
	4.2.1	List file for the input files	40
	4.2.2	CalcMod file	42
	4.2.3	ModPara file	46
	4.2.4	LocSpin file	51
	4.2.5	Trans file	53
	4.2.6	InterAll file	55
	4.2.7	CoulombIntra file	58
	4.2.8	CoulombInter file	
	4.2.9	Hund file	62
	4.2.10	PairHop file	64
	4.2.11	Exchange file	66
	4.2.12	Ising file	68
	4.2.13	PairLift file	70
	4.2.14	OneBodyG file	72
	4.2.15	TwoBodyG file	74
	4.2.16	SingleExcitation file	76
	4.2.17	PairExcitation file	78
		SpectrumVec File	
4.3	Output	t files	81
	4.3.1	$CHECK\_Chemi.dat \dots \dots$	
	4.3.2	$CHECK\_InterAll.dat  .  .  .  .  .  .  .  .  .  $	82
	4.3.3	$CHECK\_CoulombIntra.dat \ . \ . \ . \ . \ . \ . \ . \ . \ . \ $	83
	4.3.4	$CHECK\_Hund.dat  .  .  .  .  .  .  .  .  .  $	84
	4.3.5	CHECK_INTER_U.dat	85
	4.3.6	$CHECK\_Memory.dat \ . \ . \ . \ . \ . \ . \ . \ . \ . \ $	86
	4.3.7	WarningOnTransfer.dat	87
	4.3.8	CalcTimer.dat	88
	4.3.9	TimeKeeper.dat	89
	4.3.10	sz_TimeKeeper.dat	89
	4.3.11	Time_CG_EigenVector.dat	90
	4.3.12	energy.dat	91
	4.3.13	Lanczos_Step.dat	93
	4.3.14	Time_TPQ_Step.dat	95
	4.3.15	Norm_rand.dat	96
	4.3.16	SS_rand.dat	98
	4.3.17	Flct_rand.dat	100
	4.3.18	Eigenvalue.dat	102
	4.3.19	phys.dat	103
	4.3.20	ham.dat	105

Contents

		4.3.21 cisajs.dat	107
		4.3.22 cisajscktalt.dat	109
		4.3.23 eigenvec.dat	
		4.3.24 tmpvec.dat	112
		4.3.25 DynamicalGreen.dat	113
		4.3.26 recalcvec.dat	114
		4.3.27 TMcomponents.dat	115
	4.4	Error messages	116
5	Alg	gorithm 1	117
	5.1	Lanczos method	117
		5.1.1 Details of Lanczos method	117
		5.1.2 Inverse iteration method	118
		5.1.3 Details of implementation	119
	5.2	Full Diagonalization method	120
		5.2.1 Overview	120
		5.2.2 Finite-temperature calculations	120
	5.3	Finite-temperature calculations by the TPQ method	120
		5.3.1 Details of implementation	121
	5.4	Bogoliubov representation	121
6	Ack	cnowledgement 1	122

# 1

# What is $\mathcal{H}\Phi$ ?

### 1.1 What is $\mathcal{H}\Phi$ ?

Comparison between experimental observation and theoretical analysis is a crucial step in condensed-matter physics research. The tTemperature dependence of specific heat and magnetic susceptibility, for example, has been studied to extract the nature of low energy excitations of and magnetic interactions between electrons, respectively, through comparison with theories such as Landau's Fermi liquid theory and the Curie-Weiss law.

For the flexible and quantitative comparison of theoretical and experimental data, the exact diagonalization approach [1] is one of the most reliable numerical tools that requires no approximation or inspiration of genius. For the last few decades, a numerical diagonalization package for quantum spin Hamiltonians, TIT-PACK, developed by Prof. Hidetoshi Nishimori of Tokyo Institute of Technology, has been widely used in the condensed-matter physics community. Nevertheless, limited computational resources have hindered the ability of non-expert users to apply the package to quantum systems with a large number of electrons or spins.

In contrast, the recent and rapid development of a parallel computing infrastructure has opened up new avenues for user-friendly larger scale diagonalizations up to 18-site Hubbard clusters or 36 S=1/2 quantum spins. In addition, recent advances in quantum statistical mechanics [2–5] allow the finite temperature properties of quantum many-body systems to be calculated at computational costs similar to those of the calculations of ground state properties, which also allows theoretical results for the temperature dependence of, for example, specific heat and magnetic susceptibility, to be compared with experimental results quantitatively [6]. To utilize the parallel computing infrastructure with narrow bandwidth and distributed-memory architectures, efficient, user-friendly, and highly parallelized diagonalization packages are highly desirable.

 $\mathcal{H}\Phi$ , a flexible diagonalization package for solving quantum lattice Hamiltonians, has been developed as a descendant of the pioneering package TITPACK. The Lanczos method for calculations of the ground state and a few excited states properties, as well as finite temperature calculations based on thermal pure quantum states [5], are implemented in the  $\mathcal{H}\Phi$  package, with an easy-to-use and flexible user interface. By using  $\mathcal{H}\Phi$ , you can analyze a wide range of quantum lattice Hamiltonians including simple Hubbard and Heisenberg models, multi-band extensions of the Hubbard model, exchange couplings that break the SU(2) symmetry of quantum spins, such as Dzyaloshinskii-Moriya and Kitaev interactions, and Kondo lattice models describing itinerant electrons coupled with quantum spins.  $\mathcal{H}\Phi$  calculates a

variety of physical quantities, such as internal energy at zero temperature or finite temperatures, temperature dependence of specific heat, and charge/spin structure factors. A broad spectrum of users including experimental scientists is cordially welcome.

#### 1.1.1 License

The distribution of the program package and the source codes for  $\mathcal{H}\Phi$  follow GNU General Public License version 3 (GPL v3).

### 1.1.2 Copyright

© 2015- The University of Tokyo. All rights reserved. This software was developed with the support of "Project for advancement of software usability in materials science" of The Institute for Solid State Physics, The University of Tokyo.

#### 1.1.3 Contributors

This software was developed by the following contributors.

- ver.2.0.2 (released on 2017/7/5)
- ver.2.0.1(released on 2017/6/20)
- ver.2.0 (released on 2017/4/11)
- ver.1.2 (released on 2016/11/14)
- ver.1.1 (released on 2016/5/13)
- ver.1.0 (released on 2016/4/5)

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# 1.2 Operating environment

 $\mathcal{H}\Phi$  was tested on the following platforms:

- The supercomputer system-B "sekirei" in ISSP
- Fujitsu Fx-10 and K computer
- Linux PC + Intel compiler
- Linux PC + gcc
- Mac + gcc.

# 2

# How to use $\mathcal{H}\Phi$

# 2.1 Prerequisite

 $\mathcal{H}\Phi$  requires the following packages:

- C/fortran compiler (Intel, Fujitsu, GNU, etc. )
- BLAS/LAPACK library (Intel MKL, Fujitsu, ATLAS, etc.)
- MPI library (if you do not use MPI, this is not requied).

## **Tips**

#### E.g. /Settings of Intel compiler

When you use the Intel compiler, you can easily use the scripts attached to the compiler. In the case of the bash in the 64-bit OS, write the following in your ~/.bashrc:

source /opt/intel/bin/compilervars.sh intel64

or

source /opt/intel/bin/iccvars.sh intel64
source /opt/intel/mkl/bin/mklvars.sh

Please read the manuals of your compiler/library for more information.

## 2.2 Installation

You can download  $\mathcal{H}\Phi$  at the following location. https://github.com/QLMS/HPhi/releases You can obtain the  $\mathcal{H}\Phi$  directory by typing

\$ tar xzvf HPhi-xxx.tar.gz

There are two types of procedure for installing  $\mathcal{H}\Phi$ .

2.2 Installation 5

### 2.2.1 Using HPhiconfig.sh

Please run HPhiconfig.sh script in the  $\mathcal{H}\Phi$  directory as follows (for ISSP system-B "sekirei"):

#### \$ bash HPhiconfig.sh sekirei

Then, the environmental configuration file make.sys is generated in the src/directory. The command-line argument of HPhiconfig.sh is as follows:

- sekirei : ISSP system-B "sekirei"
- fujitsu : ISSP system-C "maki"
- sr : HITACHI SR16000
- intel: Intel compiler
- intel-openmpi : Intel compiler + OpenMPI
- intel-mpich : Intel compiler + MPICH2
- intel-intelmpi : Intel compiler + IntelMPI
- gcc : GCC
- gcc-openmpi : GCC + OpenMPI
- gcc-mpich : GCC + MPICH2
- gcc-mac : GCC + Mac.

make.sys is as follows (for ISSP-system-B "sekirei"):

We explain the macros of this file as:

- CC : C compiler (icc, gcc, fccpx).
- F90 : fortran compiler (ifort, gfortran, frtpx)
- CFLAGS: C compile options. OpenMP utilization option (-openmp, -fopenmp, -qopenmp, etc.) must be specified. When you use MPI, please set -D MPI.
- FFLAGS: fortran compile options. Similar to CFLAGS.

• LIBS: Compilation options for LAPACK. -Dlapack can not be removed.

Then, you are ready to compile HPhi. Please type

#### \$ make HPhi

and obtain an executable HPhi in the src/directory; you should add this directory to the \$PATH.

If SSE2 is available in your system, please add -DHAVE\_SSE2 as an option of CMake.

# Tips

You can make a PATH to  $\mathcal{H}\Phi$  as follows:

\$ export PATH=\${PATH}: HPhi\_top\_directory/src/

If you retain this PATH, you should write above in ~/.bashrc (for bash as a login shell)

### 2.2.2 Using cmake

# Tips

Before using cmake for sekirei, you must type

source /home/issp/materiapps/tool/env.sh

while for maki, you must type

source /global/app/materiapps/tool/env.sh

We can compile  $\mathcal{H}\Phi$  as

```
cd $HOME/build/hphi
cmake -DCONFIG=gcc $PathTohphi
make
```

Here, we set a path to  $\mathcal{H}\Phi$  as \$PathTohphi and to a build directory as \$HOME/build/hphi. After compilation, src folder is constructed below a \$HOME/build/hphi folder and we obtain an executable HPhi in src/ directory. When no MPI library exists in the system, an executable HPhi is automatically compiled without an MPI library.

In the above example, we compile  $\mathcal{H}\Phi$  by using a gcc compiler. We can select a compiler by using the following options:

- sekirei : ISSP system-B "sekirei"
- fujitsu : Fujitsu compiler (ISSP system-C "maki")
- intel: Intel compiler + Linux PC
- gcc : GCC compiler + Linux PC.

An example of compiling  $\mathcal{H}\Phi$  by using the Intel compiler is shown as follows:

```
mkdir ./build
cd ./build
cmake -DCONFIG=intel ../
make
```

After compilation, src folder is created below the build folder and an execute  $\mathcal{H}\Phi$  in the src folder. Please note that we must delete the build folder and repeat the above operations when we change the compiler.

# 2.3 Directory structure

When HPhi-xxx.tar.gz is unzipped, the following directory structure is composed.

```
|--CMakeLists.txt
|--COPYING
|--config/
     |--fujitsu.cmake
     |--gcc.cmake
     |--intel.cmake
     ---sekirei.cmake
|--doc/
     |--en/
     |--jp/
     |--fourier/
          |--en/
          |--figs/
          ---ja/
     |--userguide_en.pdf
     ---userguide_jp.pdf
|--HPhiconfig.sh
|--samples/
|--src/
     |--*.c
     |--CMakeLists.txt
     |--include/*.h
     |--makefile_src
     ---StdFace/
|--test/
---test_tool/
```

# 2.4 Basic usage

 $\mathcal{H}\Phi$  has two modes: Standard mode and Expert mode. Here, the basic flows of calculations of the Standard and expert modes are shown.

#### 2.4.1 Standard mode

The procedure of calculation through the standard mode is as follows:

1. Create a directory for a calculation scenario

First, you create a working directory for the calculation.

#### 2. Create input files for Standard mode

In Standard mode, you can choose a model (the Heisenberg model, Hubbard model, etc.) and a lattice (the square lattice, triangular lattice, etc.) from those provided; you can specify some parameters (such as the first/second nearest neighbor hopping integrals and the on-site Coulomb integral) for them. Finally, you have to specify the numerical method (such as the Lanczos method) employed in this calculation. The input file format is described in Sec. 4.

#### 3. Run

Run an executable HPhi in the terminal by setting option "-s" (or "--standard") and the name of the input file written in the previous step.

- Serial/OpenMP parallel\$ Path/HPhi -s Input\_file\_name
- MPI parallel/ Hybrid parallel

\$ mpiexec -np number\_of\_processes Path/HPhi -s Input\_file\_name When you use a queuing system in workstations or super computers, sometimes the number of processes is specified as an argument for the job-submitting command. If you need more information, please refer to your system manuals. The number of processes depends on the target system of the models. The details of setting the number of processes are shown in Sec. 2.4.3.

#### 4. Watch calculation logs

Log files are outputted in the "output" folder, which is automatically created in the directory for a calculation scenario. The details of the output files are shown in Sec. 4.3.

#### 5. Results

If the calculation is completed normally, the result files are outputted in the "output" folder. The details of the output files are shown in Sec. 4.3.

# Tips

#### The number of threads for OpenMP

If you specify the number of OpenMP threads for  $\mathcal{H}\Phi$ , you should set it as follows (in the case of 16 threads) before running:

export OMP\_NUM\_THREADS=16

### **2.4.2** *Expert* mode

The calculation procedure for Expert mode is as follows.

1. Create a directory for a calculation scenario First, you create a directory and give it the name of a calculation scenario (you can attach an arbitrary name to a directory).

#### 2. Create input files for Expert mode

For Expert mode, you should create input files for constructing Hamiltonian operators, calculation conditions, and a list file for the filenames of the input files (see the file formats shown in Sec. 4.2).

**Note:** A list file can be easily created by using Standard mode.

#### 3. Run

Run  $\mathcal{H}\Phi$  in the terminal by setting option "-e" (or "--expert") and the file name of a list file.

- Serial/OpenMP
  - \$ Path/HPhi -e Input\_List\_file\_name
- MPI/Hybrid

\$ mpiexec -np number\_of\_processes Path/HPhi -e Input\_List\_file\_name A number of processes depend on a target of system for models. The details of setting a number of processes are shown in 2.4.3.

### 4. While running

Log files are outputted in the "output" folder which is automatically created in the directory for a calculation scenario. The details of the output files are shown in Sec. 4.3.

#### 5. Results

If the calculation is finished normally, the result files are outputted in the "output" folder. The details of the output files are shown in Sec. 4.3.

# 2.4.3 Setting the process number for MPI/hybrid parallelization

For using MPI/hybrid parallelization, the process number must be set as follows.

#### 1. Standard mode

- Hubbard/Kondo model When model in the input file for Standard mode is set as "Fermion Hubbard", "Kondo Lattice", or "Fermion HubbardGC", the process number must be equal to  $4^n$ .
- Spin model
  When model in the input file for Standard mode is set as "Spin" or
  "SpinGC", the process number must be equal to (2S+1)<sup>n</sup>, where 2S is set
  in the input file (the default value is 1).

#### 2. Expert mode

Hubbard/Kondo model
 When the model is selected as the Fermion Hubbard model or Kondo model by setting CalcModel in a CalcMod file, the process number must be equal to 4<sup>n</sup>. See Sec. 4.2.2 for details of the CalcModel file.

• Spin model

When the model is selected as the spin model by setting CalcModel in a CalcMod file, the process number is fixed by a LocSpin file. The process number must be equal to the number calculated by multiplying the state number of the localized spin (2S+1) in descending order by the site number. See Sec. 4.2.4 for details of the LocSpin file.

For example, when a **LocSpin** file is given as follows, the process number must be equal to 2 = 1 + 1,  $6 = 2 \times (2 + 1)$ ,  $24 = 6 \times (3 + 1)$ .

=====================================	
NlocalSpin 3	
=======================================	
======i_0IteElc_2S =====	
=======================================	
0 3	
1 2	
2 1	

# 2.4.4 Printing version ID

By using the -v option as follows, you can check which version of  $\mathcal{H}\Phi$  you are using.

\$ PATH/HPhi -v

# **Tutorial**

## 3.1 Standard mode

## 3.1.1 Heisenberg model

This tutorial should be performed in

samples/CG/Heisenberg/

The input file is provided as follows:

samples/CG/Heisenberg/stan.in

In this case, we treat the two-dimensional antiferromagnetic Heisenberg model that has a nearest neighbor spin coupling.

$$\hat{H} = J \sum_{i,j=1}^{4} (\hat{\mathbf{S}}_{ij} \cdot \hat{\mathbf{S}}_{i+1j} + \hat{\mathbf{S}}_{ij} \cdot \hat{\mathbf{S}}_{ij+1},)$$
(3.1)

where we use the periodic boundary condition  $(S_{15} = S_{51} = S_{11})$ .

The input file is as follows:

```
model = "Spin"
method = "CG"
lattice = "square"
W = 4
L = 4
J = 1.0
2Sz = 0
```

In this tutorial, J and the number of sites are set to 1 (arbitral unit) and 16, respectively.

#### Log output

Log messages are outputted to the standard output. Log files for the calculation procedure are created in the "output" directory which is automatically created. In this example, the following files are outputted.

```
CHECK_InterAll.dat Time_CG_EigenVector.dat zvo_Lanczos_Step.dat CHECK_Memory.dat WarningOnTransfer.dat zvo_sz_TimeKeeper.dat check_Sdim.dat zvo_TimeKeeper.dat
```

The details of the outputted files are shown in Secs. 4.3.1-4.3.11. We execute

#### \$ パス/HPhi -s stan.in

and obtain the following standard outputs (the compilation mode is MPI parallel/hybrid parallel):

```
Welcome to the
      ,ammmmmmmmmmmb,,
    ,00' dm
                     mb ===m
  ,00' d000000000000000000 Pm,
                                  00
                                               00
                                                          @@
 d0 d000 000 000000 0000b ~0a
                                  00
                                               00
                                                      00000000
     0000 ^^^ 0000 m m 000
                                               00
                                                          00
     0000_000_0000mm mm000
                              @|
                                  @@mmmmmmmm@@
                                                   00
                                                          @@
P@
      9000000000000000000P
                              @~
                                  00000000000000000
                                                   @@
                                                          @@
                                                                @@
         ~~9@@@@@PPP~
 00
                             @P
                                  00
                                                          00
                                                              000
                                               00
                                                    000
                          ,@@~
  ~@@b
            0000000
                                  00
                                               00
                                                      000000000
                      ,m@~'
    ^{\sim}000m,,0000000000
                                  00
                                                          @@
                                               00
        ~~9@@@@@@@@
           9@P~~~9@P
                                  Version 2.0.3
```

#### ##### Parallelization Info. #####

OpenMP threads : 1 MPI PEs : 1

#### ###### Standard Intarface Mode STARTS ######

#### Open Standard-Mode Inputfile stan.in

```
      KEYWORD : model
      | VALUE : Spin

      KEYWORD : method
      | VALUE : CG

      KEYWORD : lattice
      | VALUE : square

      KEYWORD : w
      | VALUE : 4

      KEYWORD : j
      | VALUE : 4

      KEYWORD : j
      | VALUE : 1.0

      KEYWORD : 2sz
      | VALUE : 0
```

#### ###### Parameter Summary ######

#### @ Lattice Size & Shape

```
a = 1.00000 ##### DEFAULT VALUE IS USED #####
Wlength = 1.00000 ##### DEFAULT VALUE IS USED #####
Llength = 1.00000 ##### DEFAULT VALUE IS USED #####
Wx = 1.00000 ##### DEFAULT VALUE IS USED #####
```

```
Wy = 0.00000
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
              Lx = 0.00000
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
              Ly = 1.00000
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
           phase0 = 0.00000
                                ###### DEFAULT VALUE IS USED
                                                               ######
           phase1 = 0.00000
                                ###### DEFAULT VALUE IS USED
                                                               ######
  @ Super-Lattice setting
                L = 4
                W = 4
           Height = 1
                                ######
                                        DEFAULT VALUE IS USED ######
        Number of Cell = 16
  @ Hamiltonian
               h = 0.00000
                                ###### DEFAULT VALUE IS USED
                                                               ######
            Gamma = 0.00000
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
              2S = 1
                                ###### DEFAULT VALUE IS USED
                                                               ######
                D = 0.00000
                                ###### DEFAULT VALUE IS USED
                                                               ######
              J0x = 1.00000
              J0y = 1.00000
              J0z = 1.00000
              J1x = 1.00000
              J1y = 1.00000
              J1z = 1.00000
  @ Numerical conditions
       LargeValue = 4.50000
                                ##### DEFAULT VALUE IS USED #####
##### Print Expert input files #####
    locspn.def is written.
    coulombinter.def is written.
    hund.def is written.
    exchange.def is written.
                                        DEFAULT VALUE IS USED
    CDataFileHead = zvo
                                ######
                                                               ######
     Lanczos_max = 2000
                                        DEFAULT VALUE IS USED
                                ######
                                                               ######
       initial_iv = -1
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
             exct = 1
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
       LanczosEps = 14
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
                                        DEFAULT VALUE IS USED
   LanczosTarget = 2
                                ######
                                                               ######
           NumAve = 5
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
   ExpecInterval = 20
                                        DEFAULT VALUE IS USED
                                ######
                                                               ######
           NOmega = 200
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
         OmegaMax = 72.00000
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
         OmegaMin = -72.00000
                                ###### DEFAULT VALUE IS USED
                                                               ######
```

###### DEFAULT VALUE IS USED

######

OmegaIm = 0.04000

```
2Sz = 0
    modpara.def is written.
  @ Spectrum
       SpectrumQW = 0.00000
                                ######
                                        DEFAULT VALUE IS USED
                                                               ######
       SpectrumQL = 0.00000
                                ###### DEFAULT VALUE IS USED
                                                               ######
       SpectrumQH = 0.00000
                                ###### DEFAULT VALUE IS USED
                                                               ######
     SpectrumType = szsz
                                ###### DEFAULT VALUE IS USED
                                                               ######
        pair.def is written.
  @ CalcMod
                                ###### DEFAULT VALUE IS USED
          Restart = none
                                                               ######
   InitialVecType = c
                                ###### DEFAULT VALUE IS USED
                                                               ######
       EigenVecIO = none
                                ###### DEFAULT VALUE IS USED
                                                               ######
                                ###### DEFAULT VALUE IS USED
         CalcSpec = none
                                                               ######
     calcmod.def is written.
                                ##### DEFAULT VALUE IS USED #####
      ioutputmode = 1
    greenone.def is written.
    greentwo.def is written.
   namelist.def is written.
       Input files are generated. #####
######
 Read File 'namelist.def'.
 Read File 'calcmod.def' for CalcMod.
 Read File 'modpara.def' for ModPara.
 Read File 'locspn.def' for LocSpin.
 Read File 'coulombinter.def' for CoulombInter.
 Read File 'hund.def' for Hund.
 Read File 'exchange.def' for Exchange.
  Read File 'greenone.def' for OneBodyG.
 Read File 'greentwo.def' for TwoBodyG.
  Read File 'pair.def' for PairExcitation.
##### Definition files are correct. #####
  Read File 'locspn.def'.
 Read File 'coulombinter.def'.
 Read File 'hund.def'.
 Read File 'exchange.def'.
 Read File 'greenone.def'.
 Read File 'greentwo.def'.
 Read File 'pair.def'.
```

###### Indices and Parameters of Definition files(\*.def) are complete. ######

MAX DIMENSION idim\_max=12870
APPROXIMATE REQUIRED MEMORY max\_mem=0.001647 GB

#### ##### MPI site separation summary #####

INTRA process site

Bit
2
2
2
2
2
2
2
2
2
2
2
2
2
2
2
2

INTER process site
Site Bit

Process element info

Process Dimension Nup Ndown Nelec Total2Sz State
0 12870 8 8 8 0

Total dimension: 12870

#### ##### LARGE ALLOCATE FINISH ! #####

Start: Calculate HilbertNum for fixed Sz.
End : Calculate HilbertNum for fixed Sz.

Start: Calculate diagaonal components of Hamiltonian. End : Calculate diagaonal components of Hamiltonian.

#### ##### Eigenvalue with LOBPCG ######

initial\_mode=1 (random): iv = -1 i\_max=12870 k\_exct =1

```
Step
           Residual-2-norm
                                Threshold
                                                Energy
              2.44343e+00
                               1.00000e-07
                                                     -5.27456e-01
        1
        2
              2.76604e+00
                               1.87217e-07
                                                     -1.87217e+00
        3
              2.61923e+00
                               4.19088e-07
                                                     -4.19088e+00
              2.57106e+00
                               5.97098e-07
                                                     -5.97098e+00
(snip)
       40
              7.39431e-06
                               1.12285e-06
                                                     -1.12285e+01
       41
              4.15948e-06
                               1.12285e-06
                                                     -1.12285e+01
       42
              2.04898e-06
                               1.12285e-06
                                                     -1.12285e+01
              9.92048e-07
                               1.12285e-06
                                                     -1.12285e+01
       43
######
        End
             : Calculate Lanczos EigenValue.
                                                ######
######
        End
             : Calculate Lanczos EigenVec.
                                              ######
      O Energy=-11.228483 N= 16.000000 Sz= 0.000000 Doublon= 0.000000
i =
```

In the beginning of this run, files describing the details of the considered Hamiltonian (locspin.def, trans.def, exchange.def, coulombintra.def, hund.def, namelist.def, calcmod.def, modpara.def) and files specifying the elements of the correlation functions that will be calculated (greenone.def, greentwo.def) are generated.

#### Outputs for calculation results

#### Locally Optimal Block Conjugate Gradient (LOBCG) method

When a calculation by the LOBCG method is finished normally, eigenenergies, one-body Green's functions, and two-body Green's functions are calculated and outputted to the files, respectively. In this sample, the following files are outputted.

```
zvo_energy.dat
zvo_cisajscktalt_eigen_xx.dat zvo_phys_Nup4_Ndown4.dat
```

where xx is the number of the eigenstate counting from 0.

#### Lanczos method

When a calculation by the Lanczos method is completed normally, eigenenergies, one-body Green's functions, and two-body Green's functions are calculated and outputted to the files, respectively. In this sample, the following files are outputted.

```
zvo_energy.dat zvo_cisajs.dat
zvo_cisajscktalt.dat
```

For Standard mode, all pairs of  $\langle n_{i\sigma} \rangle$  are calculated as one-body Green's functions and those of  $\langle n_{i\sigma}n_{j\sigma'} \rangle$  are calculated as two-body Green's functions on the basis of the definition files, greenone.def and greentwo.def.

When the accuracy of the Lanczos vectors is sufficient, one-body and two-body Green's functions are calculated by the eigenvectors obtained by the Lanczos method. When the accuracy of the Lanczos vectors is *not* sufficient, a message "Accuracy of Lanczos vector is not enough" is outputted to the standard output and the one-body and two-body Green's functions are calculated by the eigenvectors obtained by CG method. The details of output files are shown in Secs. 4.3.12, 4.3.21, 4.3.22.

#### TPQ method

When method="TPQ" is selected in an input file, a calculation by the TPQ method is started. After the calculation is completed normally, the following files are outputted, where %% is the number of runs and && is the number of steps for the TPQ method.

```
Norm_rand%%.dat SS_rand%%.dat zvo_cisajs_set%%step&&.dat zvo_cisajscktalt_set%%step&&.dat
```

In Norm\_rand%%.dat, basic information such as the inverse of temperature and the norm of the wave function before normalization is outputted with a TPQ step for each number of runs. In SS\_rand%%.dat, physical quantities such as the inverse of temperature, energy, and expected value of the square of the Hamiltonian are outputted with a TPQ step for each number of runs. In zvo\_cisajs\_set%%step&&.dat and zvo\_cisajscktalt\_set%%step&&.dat, one-body and two-body Green's functions are outputted for each number of a TPQ steps and runs. The details of these files are shown in Secs. 4.3.15, 4.3.16, 4.3.21, 4.3.22.

#### Full diagonalization method

When method = "fulldiag" is selected in an input file, a calculation by the full diagonalization method is started. After the calculation is completed normally, the following files are outputted, where xx is the number of the eigenstate counting from 0.

```
Eigenvalue.dat zvo_cisajs_eigen_xx.dat zvo_cisajscktalt_eigen_xx.dat zvo_phys_Nup4_Ndown4.dat
```

In Eigenvalue.dat, an eigennumber and an eigenvalue are outputted for each line. In zvo\_cisajs\_eigen\_xx.dat and zvo\_cisajscktalt\_eigen\_xx.dat, one-body

Green's functions and two-body Green's functions are outputted for each eigennumber. In zvo\_phys\_Nup4\_Ndown4.dat, physical quantities, such as the expected values of energy and the doublon are outputted. The details of these files are shown in Secs. 4.3.18 - 4.3.22.

# 3.1.2 Other tutorials

There are many tutorials in samples/Standard/. For more details, please see README.md at each directory.

# 3.2 Expert mode

For Expert mode, the following input files are needed.

- 1. A file list for input files
- 2. Files for basic parameters
- 3. Files for constructing Hamiltonian
- 4. Files for setting output components.

The process after calculation is the same as in Standard mode. In this section, we demonstrate Expert mode in the directory where the tutorial at the previous section was performed.

### 3.2.1 File list for input files

In namelist.def, the types of input files and filenames are defined as shown below. By writing the keyword and filenames at each line, the types of files are distinguished. The details of namelist.def are shown in Sec. 4.2.1.

```
modpara.def
       ModPara
       LocSpin
                locspn.def
  CoulombInter
                coulombinter.def
          Hund
               hund.def
                exchange.def
      Exchange
      OneBodyG
               greenone.def
      TwoBodyG
                greentwo.def
       CalcMod
               calcmod.def
PairExcitation
               pair.def
   SpectrumVec
               zvo_eigenvec_0
```

### 3.2.2 Files for basic parameters

In this subsection, we show how to set a calculation mode, the parameters for the calculation, and the positions of the localized spins.

#### Setting a calculation mode

The calculation mode is set in a CalcMod file (in this sample file, calcmod.def). The contents of the files are as follows.

We select a calculation method in CalcType and a target model in CalcModel. In this sample, we set the Lanczos method as a calculation method and the target model as the spin system (canonical ensemble). The details of a CalcMod file are shown in Sec. 4.2.2.

### Setting parameters for calculation

The parameters for the calculation are set in a ModPara file (in this sample, modpara.def). The contents of this file are as follows.

```
Model_Parameters
_____
HPhi_Cal_Parameters
_____
CDataFileHead zvo
CParaFileHead zqp
Nsite
              16
2Sz
              2000
Lanczos_max
initial_iv
              -1
exct
              1
LanczosEps
              14
LanczosTarget
LargeValue
              4.50000000000000e+00
NumAve
ExpecInterval 20
NOmega
              200
OmegaMax
              7.200000000000000e+01
                                      4.000000000000000e-02
OmegaMin
              -7.200000000000000e+01
                                      4.00000000000000e-02
OmegaOrg
              0.0 0.0
```

In this file, we set the parameters for the calculation, such as the site number, the total number of conduction electrons, the total  $S_z$  and the number of Lanczos steps. The details of the ModPara file are shown in Sec. 4.2.3.

#### Setting positions of localized spins

The positions and S of the localized spins are defined by a LocSpin file (in this sample, locspn.def). The contents of the files are as follows.

======		 ======	=		
NlocalSp	in 16				
=======	====== i_0LocSp ======		= =		
0	1				
1	1				
2	1				
3	1				
	1				
5	1				
					,

When CalcModel in a CalcMod file is set as the spin system, all the sites are automatically treated as localized spins. The details of a LocSpin file are shown in Sec. 4.2.4.

# 3.2.3 Files for constructing Hamiltonian

After setting the basic parameters, we create input files for constructing the Hamiltonian. Since the calculations are performed by using the representation of the fermion operators in  $\mathcal{H}\Phi$ , we must rewrite the spin operator. For example, in the case of S=1/2, we rewrite the equation by using the relation

$$S_z^{(i)} = (c_{i\uparrow}^{\dagger} c_{i\uparrow} - c_{i\downarrow}^{\dagger} c_{i\downarrow})/2, \tag{3.2}$$

$$S_{+}^{(i)} = c_{i\uparrow}^{\dagger} c_{i\downarrow}, \tag{3.3}$$

$$S_{-}^{(i)} = c_{i,\downarrow}^{\dagger} c_{i\uparrow}. \tag{3.4}$$

#### Setting transfer integrals

In a Trans file (in this sample, zTrans.def), we set the transfer part of the Hamiltonian,

$$H + = -\sum_{ij\sigma_1\sigma_2} t_{ij\sigma_1\sigma_2} c_{i\sigma_1}^{\dagger} c_{j\sigma_2}. \tag{3.5}$$

The contents of the files are as follows.

We can use this term when an electric magnetic field is added in the spin system. For example, when a magnetic field is added at a site 1 such as  $-0.5S_z^{(1)}$  for S=1/2, this term can be rewritten as  $-0.5/2(c_{1\uparrow}^{\dagger}c_{1\uparrow}-c_{1\downarrow}^{\dagger}c_{1\downarrow})$ . Thus, the input file becomes as follows.

The details for a Trans file are shown in Sec. 4.2.5.

#### Setting general two-body interactions

In an InterAll file (in this sample, zInterall.def), we set the general two-body interaction part of the Hamiltonian,

$$H + = \sum_{i,j,k,l} \sum_{\sigma_1,\sigma_2,\sigma_3,\sigma_4} I_{ijkl\sigma_1\sigma_2\sigma_3\sigma_4} c_{i\sigma_1}^{\dagger} c_{j\sigma_2} c_{k\sigma_3}^{\dagger} c_{l\sigma_4}. \tag{3.6}$$

The contents of the files are as follows.

=====		=====	===						
NInter#	111	96							
======		=====	===						
======	=zInte	rAll===	===						
======	=====	=====	===						
0	0	0	0	1	0	1	0	0.500000	0.000000
0	0	0	0	1	1	1	1	-0.500000	0.000000
0	1	0	1	1	0	1	0	-0.500000	0.000000
0	1	0	1	1	1	1	1	0.500000	0.000000
0	0	0	1	1	1	1	0	1.000000	0.000000
0	1	0	0	1	0	1	1	1.000000	0.000000

Here, we explain the interaction between site i and site j in the case of S = 1/2, for simplicity. Using fermion operators, the interaction terms for the spin op-

erators can be rewritten as

$$\begin{split} H_{i,i+1} &= J(S_x^{(i)} S_x^{(i+1)} + S_y^{(i)} S_y^{(i+1)} + S_z^{(i)} S_z^{(i+1)}) \\ &= J\left(\frac{1}{2} S_+^{(i)} S_-^{(i+1)} + \frac{1}{2} S_-^{(i)} S_+^{(i+1)} + S_z^{(i)} S_z^{(i+1)}\right) \\ &= J\left[\frac{1}{2} c_{i\uparrow}^{\dagger} c_{i\downarrow} c_{i+1\downarrow}^{\dagger} c_{i+1\uparrow} + \frac{1}{2} c_{i\downarrow}^{\dagger} c_{i\uparrow} c_{i+1\uparrow}^{\dagger} c_{i+1\downarrow} + \frac{1}{4} (c_{i\uparrow}^{\dagger} c_{i\uparrow} - c_{i\downarrow}^{\dagger} c_{i\downarrow}) (c_{i+1\uparrow}^{\dagger} c_{i+1\uparrow} - c_{i+1\downarrow}^{\dagger} c_{i+1\downarrow})\right]. \end{split}$$

Thus, the interaction  $S_z^{(i)} S_z^{(i+1)}$  for J=2 can be written as

i	0	i	0	i+1	0	i+1	0	0.500000	0.000000
i	0	i	0	i+1	1	i+1	1	-0.500000	0.000000
i	1	i	1	i+1	0	i+1	0	-0.500000	0.000000
i	1	i	1	i+1	1	i+1	1	0.500000	0.000000

in the format of an InterAll file. The other terms can be written as follows.

/	/										1
1	i	0	i	1	i+1	1	i+1	0	1.000000	0.000000	
ļ	i	1	i	0	i+1	0	i+1	1	1.000000	0.000000	
١	(										1

There are other file formats for constructing the Hamiltonian. The details of the input formats of two-body interactions are shown in Secs. 4.2.6-4.2.13.

### 3.2.4 Setting output components

In OneBodyG and TwoBodyG files, the indices of one-body and two-body Green's functions are defined, respectively.

#### Setting indices of one-body Green's functions

In a OneBodyG file (in this sample, greenone.def), the indices of  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} \rangle$  are defined. The contents of files are as follows.

NCi	sAjs 		32		
===	=====	Green	func	tions	======
	0	0	0	0	
	0	0 1 0 1 0	0	1	
	1	0	1	0	
	1	1	1	1	
	2	0	2	0	

The details of the input formats of a OneBodyG file are shown in Sec. 4.2.14.

#### Setting indices of two-body Green's functions

In the TwoBodyG file (in this sample, greentwo.def), the indices of  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} c_{k\sigma_3}^{\dagger} c_{l\sigma_4} \rangle$ 

1.C . 1	$\sigma_{1}$ .		. C / 1	C1.			C 11
are defined.	- i ne	contents	or this	ше	are	as	TOHOWS.

NCisAjsC	===== ktAltD(	======= C :	-==== 1024	=====	======	==
=======================================	====== Green 	function	====== ns for S	q AND	Nq ====	== == 
0	0	0 (	) 0	0	0	0
						1
0	0	0 0	) 1	0	1	0
0	0	0 (	) 1	1	1	1
0	0	0 (	) 2	0	2	0

The details of the input formats of the TwoBodyG file are shown in Sec. 4.2.15.

# 3.2.5 Running

After creating all the input files above, we are ready to run a program. For Expert mode, we must set an option "-e" and a file name list (in this sample, namelist.def) as arguments to run  $\mathcal{H}\Phi$ .

#### \$ Path/HPhi -e namelist.def

The process after the calculation is the same as that of Standard mode.

# 3.3 Creating input files for Expert mode

This mode is for creating input files for *Expert* mode. A set of input files created using this mode gives a model provided in *Standard* mode. The usage is shown as follows.

- 1. Create an input file for *Standard* mode.
- 2. Setting an option "-sdry" and an input file (in this example, StdFace.def), run  $\mathcal{H}\Phi$ .
  - \$ Path/HPhi -sdry StdFace.def

In this case, you should not use MPI parallelization (mpirun, mpiexec, etc.).

3. The following files are created as the input files for *Expert* mode in the current working directory.

```
calcmod.def greentwo.def namelist.def zTrans.def greenone.def modpara.def zInterAll.def zlocspn.def
```

## 3.4 Fourier transformation of correlation functions

This package has a utility which performs the Fourier transformation of the correlation function and plots that function. The manual of this utility is located in

```
doc/fourier/ja/_build/html/index.html
doc/fourier/ja/_build/latex/fourier.pdf
doc/fourier/en/_build/html/index.html
doc/fourier/en/_build/latex/fourier.pdf
```

# File specification

# 4.1 Input files for Standard mode

An example of an input file for the standard mode is as follows:

```
W = 2
L = 4
model = "spin"
method = "Lanczos"

lattice = "triangular lattice"
//mu = 1.0
// t = -1.0
// t' = -0.5
// U = 8.0
//V = 4.0
//V'=2.0
J = -1.0
J'=-0.5
// nelec = 8
2Sz = 0
```

#### Basic rules for input files

- In each line, there is a set of a keyword (before an "=") and a parameter (after an "="); they are separated by "=".
- You can describe keywords in a random order.
- Empty lines and lines beginning with a "//" (comment outs) are skipped.
- Upper- and lowercase are not distinguished. Double quotes and blanks are ignored.
- There are three types of parameters.
  - 1. Parameters that must be specified (if not,  $\mathcal{H}\Phi$  will stop with error messages),
  - 2. Parameters that it is not necessary to specified (if not specified, default values are used),
  - 3. Parameters that must not be specified (if specified,  $\mathcal{H}\Phi$  will stop with error

messages).

An example of type 3 is the transfer t parameter for the Heisenberg spin system. If you choose "model=spin", you should not specify "t".

We explain each keyword as follows.

### 4.1.1 Parameters for the type of calculation

#### • model

Type: String (hoose from "Fermion Hubbard", "Spin", "Kondo Lattice", "Fermion HubbardGC", "SpinGC", "Kondo LatticeGC", "SpinGCCMA")\*1

**Description:** The target model is specified with this parameter; the expressions above denote the canonical ensemble of the Fermion in the Hubbard model

$$H = -\mu \sum_{i\sigma} c_{i\sigma}^{\dagger} c_{i\sigma} - \sum_{i \neq j\sigma} t_{ij} c_{i\sigma}^{\dagger} c_{j\sigma} + \sum_{i} U n_{i\uparrow} n_{i\downarrow} + \sum_{i \neq j} V_{ij} n_{i} n_{j}, \tag{4.1}$$

the canonical ensemble in the Spin model( $\{\sigma_1, \sigma_2\} = x, y, z$ )

$$H = -h \sum_{i} S_{iz} - \Gamma \sum_{i} S_{ix} + D \sum_{i} S_{iz} S_{iz} + \sum_{ij,\sigma_{1} \neq \sigma_{2}} J_{ij\sigma_{1}\sigma_{2}} S_{i\sigma_{1}} S_{j\sigma_{2}},$$

$$+ \sum_{ij,\sigma_{1}} J_{ij\sigma_{1}} S_{i\sigma_{1}} S_{j\sigma_{1}} + \sum_{ij,\sigma_{1} \neq \sigma_{2}} J_{ij\sigma_{1}\sigma_{2}} S_{i\sigma_{1}} S_{j\sigma_{2}},$$
(4.2)

the canonical ensemble in the Kondo lattice model

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

$$(4.3)$$

the grand canonical ensemble of the Fermion in the Hubbard model [Eqn. (4.1)], the grand canonical ensemble in the Spin model [Eqn. (4.2)], and the grand canonical ensemble in the Kondo lattice model [Eqn. (4.3)], respectively.

When model="SpinGCCMA", by using a more efficient algorithm\*2,  $\mathcal{H}\Phi$  calculates a system that is the same as "SpinGC". However, supported models and MPI processes are highly limited. See "Lattice" section.

#### • method

Type: String (choose from "Lanczos", "TPQ", "Full Diag", "CG")

**Description:** The calculation type is specified with this parameter; the above expressions above denote the single eigenstate calculation by using the Lanczos method, at the finite-temperature by using the thermally pure quantum state,

<sup>\*1</sup>GC=Grand Canonical

<sup>\*2</sup>Y. Yamaji et. al., manuscript in preparation.

the full diagonalization method, and the multiple eigenstates calculation by using the LOBCG method [7,8], respectively.

The scheme employed for the spectrum calculation is also specified with this parameter. If "CG" is chosen, the shifted bi-conjugate gradient method [9] together with the seed-switch technique [10] is employed with the help of the  $K\omega$  library [11].

#### • lattice

Type: String (choose from "Chain Lattice", "Square Lattice", "Triangular Lattice", "Honeycomb Lattice", "Ladder", "Kagome")

**Description:** The lattice shape is specified with this parameter; the expressions above denote the one-dimensional chain lattice (Fig. 4.1(a)), the two-dimensional square lattice (Fig. 4.1(b)), the two-dimensional triangular lattice (Fig. 4.1(c)), the two-dimensional anisotropic honeycomb lattice (Fig. 4.2), the ladder lattice (Fig. 4.4), and the Kagome Lattice (Fig. 4.3) respectively.

In method="SpinGCCMA", only "Chain Lattice", "Honeycomb Lattice", "Ladder", and "Kagome" are supported. The limits of L, W, and the number of MPI processes ( $N_{\rm proc}$ ) are as follows:

- "Chain Lattice"

L=8n (where n is an integer number under the condition  $n\geq 1$ ),  $N_{\text{proc}}\leq 2(L=8),\,N_{\text{proc}}\leq 2^{L/2-2}(L>8).$ 

- "Honeycomb Lattice"

$$W = 3, L \ge 2, N_{\text{proc}} \le 2(L = 2), N_{\text{proc}} \le 64(L > 2).$$

- "Ladder"

W=2, L=2n (where n is an integer number under the condition  $n\geq 4$ ),  $N_{\rm proc}\leq 2^{L-4}$ .

- "Kagome"

$$W = 3, L \ge 2, N_{\text{proc}} \le 1(L = 2), N_{\text{proc}} \le 512(L > 2).$$

#### 4.1.2 Parameters for the lattice

Chain [Fig. 4.1(a)]

• L

Type: Integer

**Description**: The length of the chain is specified with this parameter.

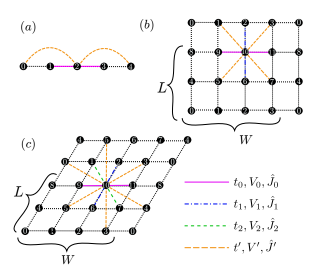


Figure 4.1: Schematic illustration of (a) one-dimensional chain lattice, (b) two-dimensional square lattice, and (c) two-dimensional triangular lattice. They have t, V, and J as the nearest neighbor hopping, an offsite Coulomb integral, and a spin-coupling constant, respectively (magenta solid lines); they also have t', V', and J' as the next nearest neighbor hopping, offsite Coulomb integral, and spin-coupling constant, respectively (green dashed line).

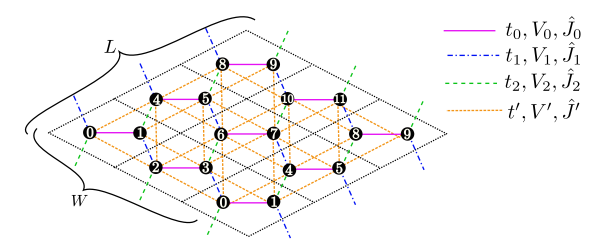


Figure 4.2: Schematic illustration of the anisotropic honeycomb lattice. The nearest neighbor hopping integral, spin coupling, and offsite Coulomb integral depend on the bond direction. Those between the second nearest neighbor sites are not supported.

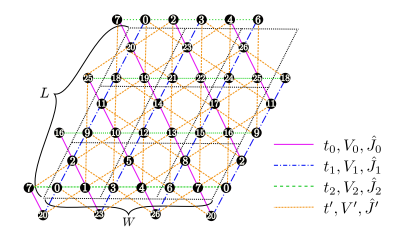


Figure 4.3: Schematic illustration of the Kagome lattice.

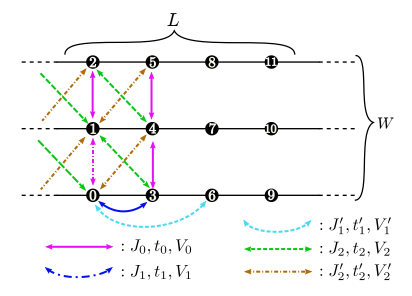


Figure 4.4: Schematic illustration of the ladder lattice.

## Ladder (Fig. 4.4)

• L

Type: Integer

**Description**: The length of the ladder is specified with this parameter.

W

Type: Integer

**Description**: The number of the ladder is specified with this parameter.

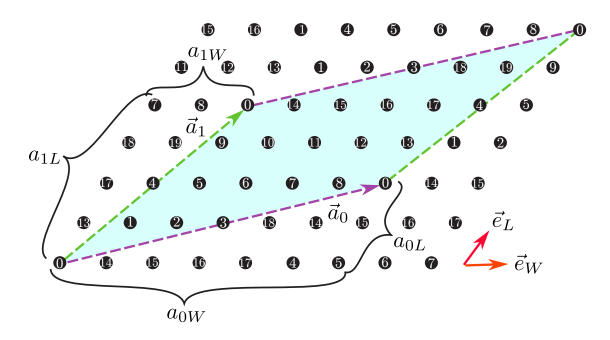


Figure 4.5: Shape of the numerical cell when  $\mathbf{a}_0 = (6, 2)$ ,  $\mathbf{a}_1 = (2, 4)$  in the triangular lattice. The region surrounded by  $\mathbf{a}_0$  (magenta dashed arrow) and  $\mathbf{a}_1$  (green dashed arrow) becomes the cell to be calculated (20 sites).

Tetragonal lattice [Fig. 4.1(b)], triangular lattice [Fig. 4.1(c)], honeycomb lattice (Fig. 4.2), Kagome lattice (Fig. 4.3)

In these lattices, we can specify the shape of the numerical cell by using the following two methods.

• W, L

Type: Integer

**Description:** The alignment of the original unit cells (dashed black lines in Figs. 4.1 - 4.3) is specified with this parameter.

• a0W, a0L, a1W, a1L

Type: Integer

**Description:** We can specify two vectors  $(\boldsymbol{a}_0, \boldsymbol{a}_1)$  that surround the numerical cell (Fig. 4.5). These vectors should be specified in the fractional coordinate.

If we use both these methods,  $\mathcal{H}\Phi$  stops. When model=SpinGCCMA, we can use only the former.

We can check the shape of the numerical cell by using a file lattice.gp which is written in Standard mode. This file can be read by gnuplot as follows:

\$ gnuplot lattice.gp

## 4.1.3 Parameters for conserved quantities

• nelec

Type: Positive integer

**Description:** The number of valence electrons is specified with this parameter. When model = "Fermion HubbardGC", "Spin", or "SpinGC", it should not be specified.

• 2Sz

Type: Integer

**Description:** The z component of the twofold total spin is specified with this parameter. When model = "Fermion HubbardGC" or "SpinGC", it should not be specified.

## 4.1.4 Parameters for the Hamiltonian

A default value is 0 unless a specific value is written in the description. Table 4.1 shows the parameters for each models. In the case of a complex type, a file format is "a real part, an imaginary part" while in the case of a real type, only "a real part"

#### Local terms

• mu

Type: Real

**Description :** It is available only for the Hubbard and Kondo lattice model. The chemical potential  $\mu$  (including the site potential) is specified with this parameter.

#### • U

Type: Real

**Description :** It is available only for the Hubbard and Kondo lattice model. The onsite Coulomb integral U is specified with this parameter.

• Jx, Jy, Jz, Jxy, Jyx, Jxz, Jzx, Jyz, Jzy

Type: Real

**Description:** It is available only for the Kondo lattice model. The spin-coupling constant between the valence and the local electrons is specified with this parameter. If the exchange coupling J is specified in the input file, instead of Jx, Jy, Jz, the diagonal exchange couplings, Jx, Jy, Jz, are set as Jx = Jy = Jz = J. When both the set of exchange couplings (Jx, Jy, Jz) and the exchange coupling J are specified in the input file,  $\mathcal{H}\Phi$  will stop.

### • h, Gamma, D

Type: Real

**Description:** (Spin model) The longitudinal magnetic field, transverse magnetic field, and the single-site anisotropy parameter are specified with these parameters. The single-site anisotropy parameter is not available for model=SpinGCCMA.

The non-local terms described below should be specified differently according to the lattice structure: For lattice=Ladder, the non-local terms are specified differently from those for the lattice=Chain Lattice, Square Lattice, Triangular Lattice, Honeycomb Lattice, Kagome. Below, the available parameters for each lattice are shown in Table 4.1.

Interactions	1D chain	2D square	2D triangular	Honeycomb	Kagome	Ladder
J, t, V (simplified)	0	0	0	0	0	-
J', t', V'	0	0	0	0	0	-
J0, t0, V0	0	0	0	0	0	0
J1, t1, V1	-	0	0	0	0	0
J2, t2, V2	-	-	0	0	0	0
J1', t1', V1'	-	-	-	-	-	0
J2',t2', V2'	_	_	_	_	_	0

Table 4.1: Interactions for each model defined in an input file. We can define spin couplings as a matrix format.

## Non-local terms for Ladder (Fig. 4.4)]

• t0, t1, t1', t2, t2'

Type: Complex

**Description:** (Hubbard and Kondo lattice model) Hopping integrals in the ladder lattice (see Fig. 4.4) are specified with this parameter.

• V0, V1, V1', V2, V2'

Type: Real

**Description:** (Hubbard and Kondo lattice model) Offsite Coulomb integrals on the ladder lattice (Fig. 4.2) are specified with these parameters.

- J0x, J0y, J0z, J0xy, J0yx, J0xz, J0zx, J0yz, J0zy
- J1x, J1y, J1z, J1xy, J1yx, J1xz, J1zx, J1yz, J1zy
- J1'x, J1'y, J1'z, J1'xy, J1'yx, J1'xz, J1'zx, J1'yz, J1'zy
- J2x, J2y, J2z, J2xy, J2yx, J2xz, J2zx, J2yz, J2zy
- J2'x, J2'y, J2'z, J2'xy, J2'yx, J2'xz, J2'zx, J2'yz, J2'zy.

Type: Real

**Description:** (Spin model) Spin-coupling constants in the ladder lattice (see Fig. 4.4) are specified with these parameters. If the simplified parameter J0 is specified in the input file instead of the diagonal couplings, J0x, J0y, J0z, these diagonal couplings are set as J0x = J0y = J0z = J0. If both J0 and the set of the couplings (J0x, J0y, J0z) are specified,  $\mathcal{H}\Phi$  will stop. The above rules are also valid for the simplified parameters, J1, J1', J2, and J2'.

## Non-local terms [other than Ladder (Figs. 4.1, 4.2, 4.3)]

• t0, t1, t2

Type: Complex

**Description:** (Hubbard and Kondo lattice model) The nearest neighbor hoppings for each direction (see Figs. 4.1-4.3) are specified with these parameters. If there is no bond dependence of the hoppings, the simplified parameter t is available to specify t0, t1, and t2 as t0 = t1 = t2 = t. If both t and the set of the hoppings (t0, t1, t2) are specified,  $\mathcal{H}\Phi$  will stop.

• V0, V1, V2

Type: Real

- JOx, JOy, JOz, JOxy, JOyx, JOxz, JOzx, JOyz, JOzy
- J1x, J1y, J1z, J1xy, J1yx, J1xz, J1zx, J1yz, J1zy

• J2x, J2y, J2z, J2xy, J2yx, J2xz, J2zx, J2yz, J2zy

Type: Real

**Description:** (Spin model) The nearest neighbor exchange couplings for each direction are specified with these parameters. If the simplified parameter J0 is specified, instead of J0x, J0y, J0z, the exchange couplings, J0x, J0y, J0z, are set as J0x = J0y = J0z = J0. If both J0 and the set of the exchange couplings (J0x, J0y, J0z) are specified,  $\mathcal{H}\Phi$  will stop. The above rules are valid for J1 and J2.

If there is no bond dependence of the exchange couplings, the simplified parameters, Jx, Jy, Jz, Jxy, Jyx, Jxz, Jzx, Jyz, Jzy, are available to specify the exchange couplings for every bond as J0x = J1x = J2x = Jx. If any simplified parameter (Jx-Jzy) is specified in addition to its counterparts (J0x-J2zy),  $\mathcal{H}\Phi$  will stop. Below, examples of parameter sets for nearest neighbor exchange couplings are shown.

- If there are no bond-dependent, and no anisotropic and offdiagonal exchange couplings (such as  $J_{xy}$ ), please specify J in the input file.
- If there are no bond-dependent and offdiagonal exchange couplings but there are anisotropic couplings, please specify the non-zero couplings in the diagonal parameters, Jx, Jy, Jz.
- If there are no bond-dependent exchange couplings but there are anisotropic and offdiagonal exchange couplings, please specify the non-zero couplings in the nine parameters, Jx, Jy, Jz, Jxy, Jyz, Jxz, Jyx, Jzy, Jzx.
- If there are no anisotropic and offdiagonal exchange couplings, but there
  are bond-dependent couplings, please specify the non-zero couplings in
  the three parameters, J0, J1, J2.
- If there are no anisotropic exchange couplings, but are bond-dependent and offdiagonal couplings, please specify the non-zero couplings in the nine parameters, J0x, J0y, J0z, J1x, J1y, J1z, J2x, J2y, J2z.
- If there are bond-dependent, anisotropic, and offdiagonal exchange couplings, please specify the non-zero couplings in the twenty-seven parameters from J0x to J2zy.

### • t'

Type: Complex

**Description:** (Hubbard and Kondo lattice model) The nearest neighbor hoppings for each direction (see Figs. 4.1-4.3) are specified with these parameter.

#### V'

Type: Real

**Description :** (Hubbard and Kondo lattice model) The nearest neighbor-offsite Coulomb integrals V for each direction (see Figs. 4.1-4.3) are specified with this parameter.

• J'x, J'y, J'z, J'xy, J'yx, J'xz, J'zx, J'yz, J'zy

Type: Real

**Description:** (Spin model) The second nearest neighbor exchange couplings are specified. However, for lattice = Honeycomb Lattice and lattice = Kagome with model=SpinGCCMA, the second nearest neighbor exchange couplings are not available in the Standard mode. If the simplified parameter J' is specified, instead of J'x, J'y, J'z, the exchange couplings are set as J'x = J'y = J'z = J'. If both J' and the set of the couplings (J'x, J'y, J'z) are specified,  $\mathcal{H}\Phi$  will stop.

## • phase0, phase1

Type: Double (0.0 as defaults)

**Description:** We can specify the phase for the hopping through the cell boundary with these parameter (unit: degree). These fuctor for the  $a_0$  direction and the  $a_1$  direction can be specified independently. For the one-dimensional system, only phase0 can be used. For example, a fopping from i-th site to j-th site through the cell boundary with the positive direction becomes as

$$\exp(i \times \text{phase0} \times \pi/180) \times t \hat{c}_{j\sigma}^{\dagger} \hat{c}_{i\sigma} + \exp(-i \times \text{phase0} \times \pi/180) \times t^* \hat{c}_{i\sigma}^{\dagger} \hat{c}_{j\sigma}$$
(4.4)

## 4.1.5 Parameters for the numerical condition

• 2S

**Type:** Positive integer (1 as a default)

**Description :** The 2S at each site in the localized spin system is specified. (E.g. 1 for the 1/2 system)

• Restart

Type: String (choose from "None", "Restart\_out", "Restart\_in", "Restart". "None" as a default)

**Description:** The condition of the restart is specified. "None" for omitting file IOs for the restart, "Restart\_out" for starting calculation from scratch and generating a restart-file after the calculation finishes, "Restart\_in" for starting calculation with the restart-file generated in the previous run, "Restart" for "Restart\_out" + "Restart\_in".

• Lanczos\_max

**Type:** Positive integer (default value: 2000)

**Description :** The upper limit of the Lanczos step is specified with this parameter.

## • initial\_iv

Type: Integer (default value: -1)

**Description**: An initial vector is specified with this parameter.

- Lanczos method
  - \* For the canonical ensemble and initial\_iv  $\geq 0$ The non-zero components of an initial vector are specified with this parameter.
  - \* For the grand canonical ensemble or initial\_iv < 0
    The seed of the random generator is given by this parameter and the random vector is used as the initial vector.

### - TPQ method

The seed of the random generator is given by this parameter and the random vector is used as the initial vector.

See Sec. 5 for details of setting an initial vector.

#### • exct

Type: Positive integer (default value: 1)

**Description:** The number of eigenvectors obtained from the ground energy by the Lanczos method are specified.

When exct=2, the eigenvector of the first-excited state is obtained. When method="CG", the number of states to be calculated is specified.

**Note**: The condition nvec >= exct must be satisfied.

### • LanczosEps

**Type:** Positive integer (default value: 14)

**Description :** The convergence criterion for the Lanczos method is specified with this parameter. If the difference between the old and the new target eigenvalue falls below  $10^{-\text{LanczosEps}}$ , the Lanczos step will finish. For method="CG", we assume the calculation is converged when the 2-norm of the residual vector becomes smaller than  $10^{-\text{LanczosEps}/2}$ .

## • LancczosTarget

**Type:** Positive integer (default value: 2)

**Description :** The target eigenenergy for the convergence criterion is specified. If it is set to 1, the target eigenenergy becomes the ground state.

## • LargeValue

**Type:** Double (the default value is provided below)

**Description :** (Only for TPQ) l as  $l - \hat{H}/N_s$  is used in the TPQ calculation. Usually, the largest eigenvalue of the Hamiltonian is used as l. Thus, the

default value of l is taken as the summation of the absolute values of each coefficient in the Hamiltonian divided by the number of sites.

### • NumAve

Type: Positive integer (default value: 5)

**Description :** (Only for TPQ) The number of independent runs for the TPQ method is specified with this parameter.

## • ExpecInterval

Type: Positive integer (default value: 20)

**Description:** (Only for TPQ) The interval of calculating correlation functions in the TPQ iteration is specified.

**Note:** A small interval increases the time cost of calculations.

### • OutputMode

Type: Choose from "none", "correlation", and "full" (correlation as default)

**Description :** Indices of correlation functions are specified with this keyword. "none" indicates correlation functions will not be calculated. When outputmode="correlation", the correlation function supported by the utility fourier is computed. For more details, see the document in doc/fourier/. If "full" is selected,  $\langle c_{i\sigma}^{\dagger}c_{j\sigma'}\rangle$  is computed at all  $i,j,\sigma,\sigma'$ , and  $\langle c_{i_1\sigma_1}^{\dagger}c_{i_2\sigma_2}c_{i_3\sigma_3}^{\dagger}c_{i_4\sigma_4}\rangle$  is computed at all  $i_1,i_2,i_3,i_4,\sigma_1,\sigma_2,\sigma_3,\sigma_4$ .

In a spin system, the indices are specified as those of the Bogoliubov representation (see Sec. 5.4).

### • InitialVecType

Type: Character (choose from "C", "R". "C" as a default)

**Description :** The type of the initial eigenvector is specified. C for the complex number, and R for the real number.

## • EigenVecIO

Type: String (choose from "None", "Out", "In". "None" as a default)

**Description :** The I/O of the eigenvector is specified. "None" for omitting the IO of the eigenvector, "Out" for writing the eigenvector to a file, "In" for reading the eigenvector from a file and using it in the subsequent calculation (such as the Green's function).

# 4.1.6 Parameters for the dynamical Green's function

## • CalcSpec

Type: String(choose from "None", "Normal", "NoIteration", "Restart\_out", "Restart\_in", "Restart". "None" as default.)

Description: The condition for the calculation of the dynamical Green's function is specified. "None" for omitting the calculation of the dynamical Green's function. "Normal" for calculating that function from scratch, "NoIteration" for calculating that function with the same iteration in the previous run (In this case, the Hamiltonian-vector product is not performed. Although the numerical cost is very small, the convergence is not guaranteed), "Restart\_out" for calculating that function from scratch and writing the restart-file at the end, "Restart\_in" for starting the calculation with the previously written restart-file, "Restart" for "Restart\_out" + "Restart\_in".

The scheme for the spectrum calculation is specified by using the parameter method. If method="CG" is chosen, the shifted bi-conjugate gradient method [9] together with the seed-switch technique [10] is employed with the help of the  $K\omega$  library [11].

## • SpectrumType

Type: String (choose from "SzSz", "S+S-", "Density", "up", "down". "SzSz" as default.)

**Description :** The type of the dynamical Green's function to be computed is specified. "SzSz" for  $\langle \hat{S}_{zq} \hat{S}_{zq} \rangle$ , "S+S-" for  $\langle \hat{S}_q^+ \hat{S}_q^- \rangle$ , "Density" for  $\langle \hat{n}_q \hat{n}_q \rangle$ , "up" for  $\langle \hat{c}_{q\uparrow}^{\dagger} \hat{c}_{q\uparrow} \rangle$ , "down" for  $\langle \hat{c}_{q\downarrow}^{\dagger} \hat{c}_{q\downarrow} \rangle$ .

## • SpectrumQW, SpectrumQL

Type: Double (default value: 0.0)

**Description:** The wave number (Fractional coordinate) of the dynamical Green's function is specified. The reciplocal lattice vector is computed from the direct lattice vector shown in Figs. 4.1, 4.2, 4.4, 4.3.

#### • OmegaMin

Type: Double (-LargeValue times the number of sites as default.) Description: The lower limit of the real part of the frequency.

#### OmegaMax

Type: Double (LargeValue times the number of sites as default.) Description: The upper limit of the real part of the frequency.

## • OmegaIm

Type: Double (0.01\*LargeValue as a default.) Description: The imaginary part of the frequency.

#### • NOmega

**Type:** Positive integer (200 as a default.) **Description:** The number of frequencies.

#### Input files for Expert mode 4.2

In this section, the details of the input files for the expert mode are explained. The input files are categorized according to the following four parts.

- (1) List: This file is a list of the input file names with the keywords. Each keyword is fixed, but the file names can be determined freely.
- (2) Basic parameters: The following input files give the basic parameters. The types of input files are determined by the keywords.

**CalcMod**: Set the parameters for the calculation modes.

**ModPara**: Set the parameters for the basic parameters, such as site number, electron number, and Lanczos step.

**LocSpin**: Set the location of the local spin (used only in the Kondo model).

(3) Hamiltonian: The Hamiltonian for  $\mathcal{H}\Phi$  is denoted by the format of the interactions for the electron system. The types of interaction are determined by the following keywords.

**Trans**: The one-body part,  $c_{i\sigma_1}^{\dagger}c_{j\sigma_2}$ 

**InterAll**: The general two-body interactions,  $c_{i\sigma_1}^{\dagger}c_{j\sigma_2}c_{k\sigma_3}^{\dagger}c_{l\sigma_4}$ .

We can set the interactions that are frequently used by the following keywords.

**CoulombIntra**: On-site Coulomb interactions,  $n_{i\uparrow}n_{i\downarrow}$   $(n_{i\sigma} = c^{\dagger}_{i\sigma}c_{i\sigma})$ 

**CoulombInter:** Off-site Coulomb interactions,  $n_i n_j$   $(n_i = n_{i\uparrow} + n_{i\downarrow})$ 

**Hund**: Hund couplings,  $n_{i\uparrow}n_{j\uparrow} + n_{i\downarrow}n_{j\downarrow}$ 

**PairHop**: Pair hopping couplings,  $c_{i\uparrow}^{\dagger}c_{j\uparrow}c_{i\downarrow}^{\dagger}c_{j\downarrow}$ 

**Exchange**: Exchange couplings,  $c_{i\uparrow}^{\dagger}c_{j\uparrow}c_{j\downarrow}^{\dagger}c_{i\downarrow}$ **Ising**: Ising interactions,  $S_i^z S_j^z$ 

**PairLift**: PairLift couplings,  $c_{i\uparrow}^{\dagger} c_{i\downarrow} c_{j\uparrow}^{\dagger} c_{j\downarrow}$ .

(4) Output: The target for the output is determined.

OneBodyG: One-body Green's functions,  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} \rangle$ TwoBodyG: Two-body Green's functions,  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} c_{k\sigma_3}^{\dagger} c_{l\sigma_4} \rangle$ .

#### 4.2.1 List file for the input files

This file determines the input filenames, which are correlated with the keywords. The file format is as follows.

```
CalcMod calcmod.def
ModPara modpara.def
LocSpin zlocspn.def
Trans ztransfer.def
InterAll zinterall.def
OneBodyG zcisajs.def
TwoBodyG zcisajscktaltdc.def
```

#### File format

```
[string01] [string02]
```

#### **Parameters**

• [string01]

Type: String

**Description**: Select a word from keywords.

• [string02]

Type: String

**Description**: An input filename that is correlated with the keywords.

## Use rules

- After setting keywords at [string 01], the half-width state is needed for writing a filename. You can set the filename freely.
- Keywords for input files are shown in Table 4.2.
- Essential keywords are "CalcMod", "ModPara", and "LocSpin".
- Keywords can be set in random order.
- If the keywords or filenames are incorrect, the program is terminated.
- When the head of a line is "#", the line is skipped.

Keywords	Details of corresponding files
CalcMod	Parameters for modes of calculation
ModPara	Parameters for calculation
LocSpin	Configurations of the local spins for Hamiltonian
Trans	Transfer and chemical potential for Hamiltonian
InterAll	Two-body interactions for Hamiltonian
CoulombIntra	CoulombIntra interactions
CoulombInter	CoulombInter interactions
Hund	Hund couplings
PairHop	Pair hopping couplings
Exchange	Exchange couplings
Ising	Ising interactions
PairLift	Pair lift couplings.
OneBodyG	Output components for one-body Green's functions $\langle c_{i\sigma}^{\dagger} c_{j\sigma} \rangle$
TwoBodyG	Output components for two-body Green's functions $\langle c_{i\sigma}^{\dagger} c_{j\sigma} c_{k\tau}^{\dagger} c_{l\tau} \rangle$
SingleExcitation	Operators for generating a single excited state
PairExcitation	Operators for generating a pair excited state
SpectrumVec	An input vector to calculate a restart vector

Table 4.2: List of the definition files

## 4.2.2 CalcMod file

This file determines the parameters for the calculation method, model, and output mode. The file format is as follows.

```
CalcType 0
CalcModel 2
CalcEigenVec 0
```

## File format

[string01] [int01]

## **Parameters**

• [string01]

Type: String

**Description :** Select a word from keywords.

• [int01]

Type: Int

**Description**: A parameter that is correlated with a keyword.

#### Use rules

- After setting the keywords at [string 01], a half-width blank is needed for setting a parameter.
- Keywords can be set in random order.
- If the keywords or filenames are incorrect, the program is terminated.
- The keywords "CalcType" and "CalcModel" are essential.
- When a head of line is "#", the line is skipped.

## Keywords and parameters

The parameters correlated with the keywords are as follows.

• CalcType

Type: Int

**Description:** Select the method for calculation from the following list:

- 0: Lanczos method
- 1: Analysis of the physical properties by using TPQ
- 2: Full diagonalization method.
- 3: LOBCG for the ground state.
- CalcModel

Type: Int

**Description**: Select the model from the following list:

- 0: Fermion Hubbard model (canonical ensemble: conservation of particles or conservation of particles and the component of  $S_z$ )
- 1: Spin model (canonical ensemble: conservation of the component of  $S_z$ )
- 2: Kondo lattice model (canonical ensemble: conservation of particles, the component of  $S_z$ )
- 3: Fermion Hubbard model (grand canonical ensemble)
- 4: Spin model (grand canonical ensemble)
- 5: Kondo lattice model (grand canonical ensemble).

For the fermion Hubbard model, you can select the model under the conservation of the particles by setting NCond in the ModPara file. When you want to select the model under the conservation of particles and the component of  $S_z$ , set both NCond and 2Sz in the ModPara file.

• CalcEigenVec

**Type:** Int (default value: 0)

**Description :** Select the method to calculate the eigenvectors:

0: Lanczos+CG methods (when the convergence of eigenvectors is not sufficient for using the Lanczos method, the CG method is applied to calculate eigenvectors).

1: Lanczos method.

## • InitialVecType

**Type:** Int (default value: 0)

**Description :** Select the type of an initial vector:

0: Complex type1: Real type.

## • OutputEigenVec

Type: Int (default value: 0)

**Description**: Select the mode of outputting an eigenvector:

0: Not output an eigenvector1: Output an eigenvector.

## • InputEigenVec

**Type:** Int (default value: 0)

**Description**: Select the mode of inputting an eigenvector:

0: Not input an eigenvector1: Input an eigenvector.

#### • ReStart

**Type:** Int (default value: 0)

**Description:** Select the mode of inputting a restart vector:

- 0: Not restart calculation
- 1: Output a restart vector
- 2: Input a restart vector and output a new restart vector
- 3: Input a restart vector.

## • CalcSpec

**Type:** Int (default value: 0)

**Description**: Select the mode of calculating dynamical Green's functions:

0: Not calculate dynamical Green's functions

1: (not restart) Input an initial vector and files for generating single excited or pair excited states

- 2: Input components of triangular diagonal matrix
- 3: Output both components of triangular diagonal matrix and a restart vector
- 4: Input both components of triangular diagonal matrix and a restart vector
- 5: Input and output both components of triangular diagonal matrix and a restart vector.

## • OutputHam

**Type:** Int (default value: 0)

 $\bf Description:$  Full Diag) Select the mode of outputting Hamiltonian:

0: not output Hamiltonian.

1: output Hamiltonian.

## • InputHam

**Type:** Int (default value: 0)

**Description:** (Full Diag)Select the mode of inputting Hamiltonian:

0: not input Hamiltonian.

1: input Hamiltonian.

## 4.2.3 ModPara file

This file determines the parameters for calculation. The file format is as follows.

Model_Paramete	rs 0
VMC_Cal_Parame	 ters
CDataFileHead	zvo
CParaFileHead	zqp
N + -	1.0
Nsite	16
Ncond	16
2Sz	0
Lanczos_max	1000
initial_iv	12
exct	1
LanczosEps	14
LanczosTarget	2
LargeValue	12
NumAve	5
ExpecInterval	20

## File format

- Lines 1-4: Header
- Line 6: [string01] [string02]
- Lines 7-8: Header
- Lines 9- : [string01] [int01].

## **Parameters**

• [string01]

Type: String

**Description:** Select a word from keywords.

• [string02]

**Type:** String (a blank parameter is not allowed)

**Description**: Set a header for output files.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description**: A parameter that is correlated with a keyword.

#### Use rules

- From Line 9: After setting keywords at [string 01], a half-width blank is needed for setting a parameter
- All the parameters are needed and the order for the parameters is fixed

## Keywords and parameters

• CDataFileHead

**Type:** String (a blank parameter is not allowed)

**Description:** A header for output files. For example, the output filename for one-body Green's function becomes "xxx\_Lanczos\_cisajs.dat" (xxx are the characters set by CDataFileHead).

• Nsite

Type: Int (positive integer)

**Description**: The number of sites.

• Ncond

Type: Int (positive integer)

**Description :** The number of conduction electrons (not used in grand canonical ensemble).

• 2Sz

**Type:** Int (positive integer)

**Description:** The total value of  $2S_z$  (not used in grand canonical ensemble). For conservation of  $S_z$  in the case of CalcModel = 0 (fermion Hubbard model) or 2 (Kondo lattice model), set Ncond.

• Lanczos\_max

Type: Int (positive integer)

**Description:** The maximum number of Lanczos steps in the calculation. When the convergence within the specified accuracy is satisfied, the calculation is completed before a step reaches Lanczos\_max. For TPQ calculation, the total number of TPQ steps is specified with this parameter. In the case of restart calculation, Lanczos\_max must be larger than that of the previous calculation.

### • initial\_iv

Type: Int

**Description**: An initial vector is specified with this parameter.

#### - Lanczos method

- \* For canonical ensemble and initial\_iv  $\geq 0$ The non-zero components of an initial vector are specified with this parameter.
- \* For grand canonical ensemble or initial\_iv < 0
  The seed of the random generator is given by this parameter and the random vector is used as the initial vector.

## - TPQ method

The seed of the random generator is given by this parameter and the random vector is used as the initial vector.

See Sec. 5 for details of setting an initial vector.

#### • exct

Type: Int (positive integer)

**Description :** An integer for setting the number of eigenvectors obtained from the ground energy by the Lanczos method.

For method="CG", the number of eigenvectors is specified.

## • LanczosEps

**Type:** Int (positive integer)

**Description :** An integer for judging the convergence of the Lanczos method. The convergence is determined by whether the condition is satisfied that the relative error between an eigenvalue and an eigenvalue at the Lanczos step of the one step before is less than  $10^{-\text{LanczosEps}}$ . For method="CG", the calculation finishes when the 2-norm of the residual vector becomes smaller than  $10^{-\text{LanczosEps}/2}$ .

## • LanczosTarget

**Type:** Int (positive integer)

**Description:** An integer giving the target of the eigenvalue for judging the convergence of the Lanczos method. For example, the target becomes a ground state when LanczosTarget is equal to one, and a first excited state when LanczosTarget is equal to two.

#### • CalcHS

**Type:** Int (positive integer)

**Description :** If CalcHS=1, an efficient algorithm for generating the restricted Hilbert space with the specified quantum number is used (Details of algorithm is shown in http://qlms.github.io/HPhi/develop/tips.pdf[in Japanese]). Default value is 1 and the efficient algorithm is used.

## • LargeValue

Type: Double

**Description :** (Use only for the TPQ method) An integer giving l of  $l - \hat{H}/N_s$  used in the TPQ method.

#### • NumAve

Type: Int

**Description:** (Use only for the TPQ method) An integer giving the number of independent runs for the TPQ method.

### • ExpecInterval

Type: Int

**Description:** (Use only for the TPQ method) An integer giving the interval steps of calculating the correlation functions in the TPQ method.

**Note:** A small interval increases the time cost of calculations.

## • OmegaOrg

Type: Complex

**Description:** (Use only for calculating dynamical Green's functions) The center value of the frequency. Specify the real and imaginary parts in that order separated by a space, and if there is no imaginary part, the real part of the frequency is only given.

## • OmegaIm

Type: Double

**Description:** (Use only for calculating dynamical Green's functions) The imaginary part of the frequency. When OmegaOrg is defined in a modpara file, OmegaIm is added to the imaginary value of OmegaOrg.

## • OmegaMin

Type: Complex

**Description:** (Use only for calculating dynamical Green's functions) The lower limit of the frequency from OmegaOrg. Specify the real and imaginary parts in that order separated by a space, and if there is no imaginary part, the real part of the frequency is only given.

## • OmegaMax

Type: Complex

**Description:** (Use only for calculating dynamical Green's functions) The upper limit of the frequency from OmegaOrg. Specify the real and imaginary parts in that order separated by a space, and if there is no imaginary part, a real part of the frequency is only given.

## • NOmega

Type: Int

**Description :** (Use only for calculating dynamical Green's functions) The integer for defining the step size of the frequency  $\Delta \omega = (\text{OmegaMax-OmegaMin})/N_{\omega}$ . The frequency is given by  $z_n = \text{OmegaOrg} + \text{OmegaMax} + \Delta \omega \times n$ .

# 4.2.4 LocSpin file

This file determines sites with localized spins. The file format is as follows.

NlocalS	====== pin ======	6 
======	=i_0Loc	Spn_1IteElc ======
0	1	
1	0	
2	1	
3	0	
4	1	
5	0	
6	1	
7	0	
8	1	
9	0	
10	1	
11	0	

### File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03].

#### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description**: A keyword for the total number of localized spins. You can freely give a name to the keyword.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving the total number of localized spins.

• [int02]

Type: Int (a blank parameter is not allowed)

**Description :** An integer giving a site index  $(0 \le [int02] \le Nsite)$ .

• [int03]

Type: Int (a blank parameter is not allowed)

**Description :** An integer for selecting an electron state whether the electron state is a localized spin or an itinerant electron state:

0: Itinerant electron state

n > 0: Localized spin state with 2S = n.

## Use rules

- Headers cannot be omitted.
- A program is terminated when [int01] is different from the total number of localized spins indicated by [int03].
- A program is terminated, when [int02] is different from the total number of sites.
- A program is terminated under the condition [int02] < 0 or Nsite <= [int02].

## 4.2.5 Trans file

This file determines the values of the transfer integrals  $t_{ij\sigma_1\sigma_2}$ ,

$$H + = -\sum_{ij\sigma_1\sigma_2} t_{ij\sigma_1\sigma_2} c_{i\sigma_1}^{\dagger} c_{j\sigma_2}. \tag{4.5}$$

An example of the file format is as follows.

======						
NTransfe	er	24				
=======	=====					
======	i_j_s_	_tijs==				
		-====		4 000000	0 00000	
0	0	2	0	1.000000	0.000000	
2	0	0	0	1.000000	0.000000	
0	1	2	1	1.000000	0.000000	
2	1	0	1	1.000000	0.000000	
2	0	4	0	1.000000	0.000000	
4	0	2	0	1.000000	0.000000	
2	1	4	1	1.000000	0.000000	
4	1	2	1	1.000000	0.000000	
4	0	6	0	1.000000	0.000000	
6	0	4	0	1.000000	0.000000	
4	1	6	1	1.000000	0.000000	
6	1	4	1	1.000000	0.000000	
6	0	8	0	1.000000	0.000000	
8	0	6	0	1.000000	0.000000	

## File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03] [int04] [int05] [double01] [double02].

#### **Parameters**

• [string01]

Type: String (a blank parameter is not allowed)

**Description**: A keyword for the total number of transfer integrals. You can freely give a name to the keyword.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description:** An integer giving the total number of transfer integrals.

• [int02], [int04]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving a site index  $(0 \le [int02], [int04] < Nsite)$ .

• [int03], [int05]

Type: Int (a blank parameter is not allowed)

**Description:** An integer giving a spin index:

0: Up-spin1: Down-spin.

• [double01]

Type: Double (a blank parameter is not allowed)

**Description**: A value for a real part of  $t_{ij\sigma_1\sigma_2}$ .

• [double02]

Type: Double (a blank parameter is not allowed)

**Description**: A value for an imaginary part of  $t_{ij\sigma_1\sigma_2}$ .

### Use rules

- Headers cannot be omitted.
- Since the Hamiltonian must be Hermitian, the relation  $t_{ij\sigma_1\sigma_2} = t^{\dagger}_{ji\sigma_2\sigma_1}$  must be satisfied. A program is terminated when this relation is broken.
- A program is terminated when the components of the on-site interactions are double counted.
- A program is terminated when [int01] is different from the total number of transfer integrals defined in this file.
- A program is terminated when [int02]-[int05] are outside the range of the defined values.

## 4.2.6 InterAll file

This file determines the values of generalized two-body interactions integrals  $I_{ijkl\sigma_1\sigma_2\sigma_3\sigma_4}$ ,

$$H + = \sum_{i,j,k,l} \sum_{\sigma_1,\sigma_2,\sigma_3,\sigma_4} I_{ijkl\sigma_1\sigma_2\sigma_3\sigma_4} c_{i\sigma_1}^{\dagger} c_{j\sigma_2} c_{k\sigma_3}^{\dagger} c_{l\sigma_4}. \tag{4.6}$$

For spin, the conditions i=j and k=l must be satisfied. An example of the file format is as follows.

===:	=====	=====	=====	====					
NIN'	terAl 		36 						
=====zInterAll====									
0	0	0	1	1	1	1	0	0.50	0.0
	1	0	0	1	0	1	1	0.50	0.0
	0	0	0	1	0	1	0	0.25	0.0
)	0	0	0	1	1	1	1	-0.25	0.0
)	1	0	1	1	0	1	0	-0.25	0.0
)	1	0	1	1	1	1	1	0.25	0.0
	0	2	1	3	1	3	0	0.50	0.0
	1	2	0	3	0	3	1	0.50	0.0
	0	2	0	3	0	3	0	0.25	0.0
	0	2	0	3	1	3	1	-0.25	0.0
	1	2	1	3	0	3	0	-0.25	0.0
	1	2	1	3	1	3	1	0.25	0.0
	0	4	1	5	1	5	0	0.50	0.0
:	1	4	0	5	0	5	1	0.50	0.0
	0	4	0	5	0	5	0	0.25	0.0
	0	4	0	5	1	5	1	-0.25	0.0
	1	4	1	5	0	5	0	-0.25	0.0
Ŀ	1	4	1	5	1	5	1	0.25	0.0

## File format

- Line 1: Header
- Line 2: [string01] [int01]
- Lines 3-5: Header
- $\bullet$  Lines 6-: [int02] [int03] [int04] [int05] [int06] [int07] [int08] [int09] [double01] [double02].

#### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description:** A keyword for the total number of generalized two-body interactions. You can freely give a name to the keyword.

• [int01]

**Type:** Int (a blank parameter is not allowed)

**Description :** An integer giving the total number of generalized two-body interactions.

• [int02], [int04], [int06], [int08]

Type: Int (a blank parameter is not allowed)

**Description :** An integer giving a site index  $(0 \le [int02], [int04], [int06], [int08] \le Nsite).$ 

• [int03], [int05], [int07], [int09]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a spin index:

0: Up-spin,

1: Down-spin.

• [double01]

**Type:** Double (a blank parameter is not allowed)

**Description**: A value for a real part of  $I_{ijkl\sigma_1\sigma_2\sigma_3\sigma_4}$ .

• [double02]

**Type:** Double (a blank parameter is not allowed)

**Description**: A value for an imaginary part of  $I_{ijkl\sigma_1\sigma_2\sigma_3\sigma_4}$ .

#### Use rules

- Headers cannot be omitted.
- Since the Hamiltonian must be Hermitian, the relation  $I_{ijkl\sigma_1\sigma_2\sigma_3\sigma_4} = I^{\dagger}_{lkji\sigma_4\sigma_3\sigma_2\sigma_1}$  must be satisfied. A program is terminated when this relation is broken. It is noted that the term of the Hermitian conjugate for  $I_{ijkl\sigma_1\sigma_2\sigma_3\sigma_4}c^{\dagger}_{i\sigma_1}c_{j\sigma_2}c^{\dagger}_{k\sigma_3}c_{l\sigma_4}$  should be inputted as  $I_{lkji\sigma_4\sigma_3\sigma_2\sigma_1}$   $c^{\dagger}_{l\sigma_4}c_{k\sigma_3}c^{\dagger}_{j\sigma_2}c_{i\sigma_1}$ .
- A program is terminated when the conditions i = j and k = l are not satisfied for the spin model.
- A program is terminated when the components of the on-site interactions are double counted.

- A program is terminated when [int01] is different from the total number of generalized two-body interactions defined in this file.
- A program is terminated when [int02]-[int09] are outside the range of the defined values.

## 4.2.7 CoulombIntra file

This file determines the values of the on-site interactions  $U_i$  (for S=1/2 system),

$$H + = \sum_{i} U_{i} n_{i\uparrow} n_{i\downarrow}. \tag{4.7}$$

An example of the file format is as follows.

- 0 4.000000
- 1 4.000000
- 2 4.000000
- 3 4.000000
- 4 4.000000
- 5 4.000000

#### File format

- Line 1: Header
- Line 2: [string01] [int01]
- Lines 3-5: Header
- Lines 6-: [int02] [double01].

## **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description :** A keyword for the total number of on-site interactions. You can freely give a name to the keyword.

• [int01]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving the total number of on-site interactions.

• [int02]

**Type:** Int (a blank parameter is not allowed)

**Description :** An integer giving a site index  $(0 \le [int02] \le Nsite)$ .

• [double01]

Type: Double (a blank parameter is not allowed)

**Description**: A value for  $U_i$ .

## Use rules

• Headers cannot be omitted.

- A program is terminated when the components of on-site interactions are double counted.
- A program is terminated when [int01] is different from the total number of on-site interactions defined in this file.
- A program is terminated when [int02] is outside the range of the defined values.

## 4.2.8 CoulombInter file

This file determines the values of off-site interactions  $V_{ij}$  (for S=1/2 system),

$$H + = \sum_{i,j} V_{ij} n_i n_j. \tag{4.8}$$

An example of the file format is as follows.

========	
NCoulombInt	ser 6
========	=========
======Cou	lombInter =====
========	========
0 1	1.0000
1 2	1.0000
2 3	1.0000
3 4	1.0000
4 5	1.0000
5 0	1.0000

#### File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03] [double01].

### **Parameters**

• [string01]

Type: String (a blank parameter is not allowed)

**Description :** A keyword for the total number of off-site interactions. You can freely give a name to the keyword.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving the total number of off-site interactions.

• [int02], [int03]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a site index  $(0 \le [int02], [int03] \le Nsite)$ .

• [double01]

Type: Double (a blank parameter is not allowed)

**Description**: A value for  $V_{ij}$ .

## Use rules

• Headers cannot be omitted.

- A program is terminated when the components of off-site interactions are double counted.
- A program is terminated when [int01] is different from the total number of off-site interactions defined in this file.
- A program is terminated when either [int02] or [int03] is outside the range of the defined values.

## **4.2.9** Hund file

This file determines the values of Hund couplings  $J_{ij}^{\text{Hund}}$  (for S=1/2 system),

$$H + = -\sum_{i,j} J_{ij}^{\text{Hund}} (n_{i\uparrow} n_{j\uparrow} + n_{i\downarrow} n_{j\downarrow}). \tag{4.9}$$

An example of the file format is as follows.

======		
NHund 6	3	
======		
======	==Hund =====	
======		
0	1 -0.250000	
1	2 -0.250000	
2	3 -0.250000	
3	4 -0.250000	
4	5 -0.250000	
5	0 -0.250000	
5	0 -0.250000	

### File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03] [double01].

### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description :** A keyword for the total number of Hund couplings. You can freely give a name to the keyword.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving the total number of Hund couplings.

• [int02], [int03]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a site index  $(0 \le [int02], [int03] \le Nsite)$ .

• [double01]

Type: Double (a blank parameter is not allowed)

**Description :** A value for  $J_{ij}^{\text{Hund}}$ .

## Use rules

• Headers cannot be omitted.

- A program is terminated when the components of the Hund couplings are double counted.
- A program is terminated when [int01] is different from the total number of Hund couplings defined in this file.
- A program is terminated when either [int02] or [int03] is outside the range of the defined values.

## 4.2.10 PairHop file

This file determines the values of PairHop couplings  $J_{ij}^{\text{Pair}}$  (for S=1/2 system),

$$H + = \sum_{i,j} J_{ij}^{\text{Pair}} (c_{i\uparrow}^{\dagger} c_{j\uparrow} c_{i\downarrow}^{\dagger} c_{j\downarrow} + h.c.). \tag{4.10}$$

An example of the file format is as follows.

======	====	=======
NPairho	p 6	
======	====	=======
======	=Pai	rhop =====
======	:====	=======
0	1	0.50000
1	2	0.50000
2	3	0.50000
3	4	0.50000
4	5	0.50000
5	0	0.50000

### File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03] [double01].

### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description :** A keyword for the total number of PairHop couplings. You can freely give a name to the keyword.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving the total number of PairHop couplings.

• [int02], [int03]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a site index  $(0 \le [int02], [int03] \le Nsite)$ .

• [double01]

Type: Double (a blank parameter is not allowed)

**Description :** A value for  $J_{ij}^{\text{Pair}}$ .

## Use rules

• Headers cannot be omitted.

- A program is terminated when [int01] is different from the total number of the PairHop couplings defined in this file.
- A program is terminated when either [int02] or [int03] is outside the range of the defined values.

# 4.2.11 Exchange file

This file determines the values of Exchange couplings  $J_{ij}^{\text{Ex}}$  (for S=1/2 system). For the fermion electronic system, the exchange terms are given as

$$H + = \sum_{i,j} J_{ij}^{\text{Ex}} (c_{i\uparrow}^{\dagger} c_{j\uparrow} c_{j\downarrow}^{\dagger} c_{i\downarrow} + c_{i\downarrow}^{\dagger} c_{j\downarrow} c_{j\uparrow}^{\dagger} c_{i\uparrow}), \tag{4.11}$$

while for the spin system, they are given as

$$H + = \sum_{i,j} J_{ij}^{\text{Ex}} (S_i^+ S_j^- + S_i^- S_j^+). \tag{4.12}$$

We note that  $(S_i^+ S_j^- + S_i^- S_j^+)$  in the spin system is written by the operators for electrons as  $-(c_{i\uparrow}^{\dagger} c_{j\uparrow} c_{j\downarrow}^{\dagger} c_{i\downarrow} + c_{i\downarrow}^{\dagger} c_{j\downarrow} c_{j\uparrow}^{\dagger} c_{i\uparrow})$ . An example of the file format is as follows.

======	====	========		
NExchan	ge 6			
	====	========		
======	=Exc	hange =====		
======	====	========		
0	1	0.50000		
1	2	0.50000		
2	3	0.50000		
3	4	0.50000		
4	5	0.50000		
5	0	0.50000		

#### File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03] [double01].

## **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description:** A keyword for the total number of Exchange couplings. You can freely give a name to the keyword.

• [int01]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving the total number of Exchange couplings.

• [int02], [int03]

Type: Int (a blank parameter is not allowed)

**Description :** An integer giving a site index  $(0 \le [int02], [int03] \le Nsite)$ .

• [double01]

Type: Double (a blank parameter is not allowed)

**Description :** A value for  $J_{ij}^{\text{Ex}}$ .

## Use rules

• Headers cannot be omitted.

- A program is terminated when the components of the exchange couplings are double counted.
- A program is terminated when [int01] is different from the total number of exchange couplings defined in this file.
- A program is terminated when either [int02] or [int03] is outside the range of the defined values.

## 4.2.12 Ising file

This file determines the values of Ising interactions  $J_{ij}^z$  (for S=1/2 system). For the fermion electronic system, the Ising terms are given as

$$H + = \sum_{i,j} J_{ij}^{z} (n_{i\uparrow} - n_{i\downarrow})(n_{j\uparrow} - n_{j\downarrow}). \tag{4.13}$$

For the spin system, they are given as

$$H + = \sum_{i,j} J_{ij}^z S_i^z S_j^z. \tag{4.14}$$

An example of the file format is as follows.

======	====			
NIsing	6			
======	====	========		
======	=Isi	ng =====		
======	====	========		
0	1	0.50000		
1	2	0.50000		
2	3	0.50000		
3	4	0.50000		
4	5	0.50000		
5	0	0.50000		

## File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

 $\bullet$  Lines 6-: [int02] [int03] [double01].

#### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description :** A keyword for the total number of Ising interactions. You can freely give a name to the keyword.

• [int01]

**Type:** Int (a blank parameter is not allowed)

**Description:** An integer giving the total number of Ising interactions.

• [int02], [int03]

Type: Int (a blank parameter is not allowed)

**Description :** An integer giving a site index  $(0 \le [int02], [int03] \le Nsite)$ .

• [double01]

Type: Double (a blank parameter is not allowed)

**Description**: A value for  $J_{ij}^{z}$ .

## Use rules

• Headers cannot be omitted.

- A program is terminated when the components of the Ising interactions are double counted.
- A program is terminated when [int01] is different from the total number of Ising interactions defined in this file.
- A program is terminated when either [int02] or [int03] is outside the range of the defined values.

## 4.2.13 PairLift file

This file determines the values of PairLift couplings  $J_{ij}^{\text{PairLift}}$  (for S=1/2 system),

$$H + = \sum_{i,j} J_{ij}^{\text{PairLift}} (c_{i\uparrow}^{\dagger} c_{i\downarrow} c_{j\uparrow}^{\dagger} c_{j\downarrow} + c_{i\downarrow}^{\dagger} c_{i\uparrow} c_{j\downarrow}^{\dagger} c_{j\uparrow}). \tag{4.15}$$

An example of the file format is as follows.

======	====	=======	
NPairLi	ft 6		
======		=======	
======	=NPa	irLift =====	
======	====	=======	
0	1	0.50000	
1	2	0.50000	
2	3	0.50000	
3	4	0.50000	
4	5	0.50000	
5	0	0.50000	
\			

#### File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03] [double01] .

#### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description:** A keyword for the total number of PairLift couplings. You can freely give a name to the keyword.

• [int01]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving the total number of PairLift couplings.

• [int02], [int03]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a site index  $(0 \le [int02], [int03] \le Nsite)$ .

• [double01]

Type: Double (a blank parameter is not allowed)

**Description**: A value for  $J_{ij}^{\text{PairLift}}$ .

## Use rules

• Headers cannot be omitted.

- A program is terminated when the components of the PairLift couplings are double counted.
- A program is terminated when [int01] is different from the total number of PairLift couplings defined in this file.
- A program is terminated when either [int02] or [int03] is outside the range of the defined values.

# 4.2.14 OneBodyG file

This file determines the target components of the one-body Green's function  $\langle c_{i\sigma_1}^{\dagger}c_{j\sigma_2}\rangle$ . An example of the file format is as follows.

NCisAjs		24	<b></b>	
======				
======	Gree	n funct	cions =	====
				=====
0	0	0	0	
0	1	0	1	
1	0	1	0	
1	1	1	1	
2	0	2	0	
2	1	2	1	
3	0	3	0	
3	1	3	1	
4	0	4	0	
4	1	4	1	
5	0	5	0	
5	1	5	1	
6	0	6	0	
6	1	6	1	
7	0	7	0	
7	1	7	1	
8	0	8	0	
8	1	8	1	
9	0	9	0	
9	1	9	1	
10	0	10	0	
10	1	10	1	
11	0	11	0	
11	1	11	1	

## File format

• Line 1: Header

• Line 2: [string01] [int01]

• Lines 3-5: Header

• Lines 6-: [int02] [int03] [int04] [int05].

#### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description:** A keyword for the total number of one-body Green's functions. You can freely give a name to the keyword.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description:** An integer giving the total number of one-body Green's functions.

• [int02], [int04]

Type: Int (a blank parameter is not allowed)

**Description :** An integer giving a site index  $(0 \le [int02], [int04] < Nsite)$ .

• [int03], [int05]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a spin index:

For Hubbard or Kondo system, the index can be selected from

0: Up-spin,

1: Down-spin.

For Spin system, the index can be selected from

 $0, 1, \dots, 2S + 1$  (corresponding to  $-S - 0.5, -S + 0.5, \dots S + 0.5$ ).

#### Use rules

- Headers cannot be omitted.
- A program is terminated when the components of the one-body Green's functions are double counted.
- A program is terminated when [int01] is different from the total number of one-body Green's functions defined in this file.
- A program is terminated when [int02]-[int05] are outside the range of the defined values.

# 4.2.15 TwoBodyG file

This file determines the target components of the two-body Green's function  $\langle c_{i\sigma_1}^{\dagger}c_{j\sigma_2}c_{k\sigma_3}^{\dagger}c_{l\sigma_4}\rangle$ . For the spin system, the conditions i=j and k=l must be satisfied. An example of the file format is as follows.

CisAjsC 	ktAltD(	C	5	76 			
						Nq =====	
0	0	0	0	0	0	0	(
0	0	0	0	0	1	0	1
0	0	0	0	1	0	1	(
0	0	0	0	1	1	1	1
0	0	0	0	2	0	2	C
0	0	0	0	2	1	2	1
0	0	0	0	3	0	3	(
0	0	0	0	3	1	3	1
0	0	0	0	4	0	4	C
0	0	0	0	4	1	4	1
0	0	0	0	5	0	5	0
0	0	0	0	5	1	5	1
0	0	0	0	6	0	6	C
0	0	0	0	6	1	6	1
0	0	0	0	7	0	7	0
0	0	0	0	7	1	7	1
0	0	0	0	8	0	8	0
0	0	0	0	8	1	8	1
0	0	0	0	9	0	9	C
0	0	0	0	9	1	9	1
0	0	0	0	10	0	10	0
0	0	0	0	10	1	10	1
0	0	0	0	11	0	11	0
0	0	0	0	11	1	11	1
0	1	0	1	0	0	0	0

## File format

- Line 1: Header
- Line 2: [string01] [int01]
- Lines 3-5: Header
- $\bullet$  Lines 6-: [int02] [int03] [int04] [int05] [int06] [int07] [int08] [int09].

#### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description:** A keyword for the total number of two-body Green's functions. You can freely give a name to the keyword.

• [int01]

Type: Int (a blank parameter is not allowed)

**Description:** An integer giving the total number of two-body Green's functions.

• [int02], [int04], [int06], [int08]

Type: Int (a blank parameter is not allowed)

**Description :** An integer giving a site index  $(0 \le [int02], [int04], [int06], [int08] \le Nsite).$ 

• [int03], [int05], [int07], [int09]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a spin index:

For Hubbard or Kondo system, the index can be selected from

0: Up-spin,

1: Down-spin.

For Spin system, the index can be selected from

 $0, 1, \dots, 2S + 1$  (corresponding to  $-S - 0.5, -S + 0.5, \dots S + 0.5$ ).

## Use rules

- Headers cannot be omitted.
- A program is terminated when the components of the two-body Green's functions are double counted.
- A program is terminated when the conditions i = j and k = l are not satisfied for the spin model.
- A program is terminated when [int01] is different from the total number of two-body Green's functions defined in this file.
- A program is terminated, when [int02]-[int09] are outside the range of the defined values.

## 4.2.16 SingleExcitation file

The operators to generate the single excited state  $c_{i\sigma_1}(c_{i\sigma_1}^{\dagger})$  are defined. An example of the file format is as follows.

======	=====			=====
NSingle		24		
=======	Sing	le Exc	citatior	===== 1 ======
0	0	0	1.0	0.0
0	1	0	1.0	0.0
1	0	0	1.0	0.0
(cont	inue.	)		
11	0	0	1.0	0.0
11	1	0	1.0	0.0

## File format

- Line 1: Header
- Line 2: [string01] [int01]
- Lines 3-5: Header
- Lines 6-: [int02] [int03] [int04] [double01] [double02].

## **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description :** A keyword for the total number of single excitation operators. You can freely give a name to the keyword.

• [int01]

**Type:** Int (a blank parameter is not allowed)

**Description:** An integer giving the total number of total number of single excitation operators.

• [int02]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving a site index  $(0 \le [int02] \le Nsite)$ .

• [int03]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving a spin index:

For Hubbard or Kondo system, the index can be selected from

0: Up-spin,

1: Down-spin.

For Spin system, the index can be selected from

 $0, 1, \dots, 2S + 1$  (corresponding to  $-S - 0.5, -S + 0.5, \dots S + 0.5$ ).

• [int04]

**Type:** Int (a blank parameter is not allowed)

**Description:** An integer giving a type of single excitation operators:

0:  $c_{i\sigma_1}$ 1:  $c_{i\sigma_1}^{\dagger}$ .

• [double01], [double02]

**Type:** Double (a blank parameter is not allowed)

**Description**: [double01] gives the real part of  $c_{i\sigma_1}(c_{i\sigma_1}^{\dagger})$ , while [double02] gives the imaginary part of  $c_{i\sigma_1}(c_{i\sigma_1}^{\dagger})$ .

#### Use rules

- Headers cannot be omitted.
- A program is terminated when the components of the single excitation operators are double counted.
- A program is terminated when the conditions i = j and k = l are not satisfied for spin model.
- A program is terminated when [int01] is different from the total number of two-body Green's functions defined in this file.
- A program is terminated, when [int02]-[int04] are outside the range of the defined values.

## 4.2.17 PairExcitation file

The operators to generate the pair excited state  $c_{i\sigma_1}c_{j\sigma_2}^{\dagger}(c_{i\sigma_1}^{\dagger}c_{j\sigma_2})$  are defined. The type of pair excitation operators  $(c_{i\sigma_1}c_{j\sigma_2}^{\dagger})$  or  $c_{i\sigma_1}^{\dagger}c_{j\sigma_2}$  must be same in the input file. In the  $S_z$  conserved system,  $\sigma_1$  must be equal to  $\sigma_2$ . An example of the file format is as follows.

======	=====	=====		=====	=		
NPair		24					
=======	Pair	Excita	ation	===== =====	= = -		
0	0	0	0	0	1.0	0.0	
0	1	0	1	0	1.0	0.0	
1	0	1	0	0	1.0	0.0	
(cont	inue.	)					
11	0	11	0	0	1.0	0.0	
11	1	11	1	0	1.0	0.0	

#### File format

- Line 1: Header
- Line 2: [string01] [int01]
- Lines 3-5: Header
- Lines 6-: [int02] [int03] [int04] [int05] [int06] [double01] [double02].

#### **Parameters**

• [string01]

**Type:** String (a blank parameter is not allowed)

**Description:** A keyword for the total number of the pair excitation operators. You can freely give a name to the keyword.

• [int01]

**Type:** Int (a blank parameter is not allowed)

**Description:** An integer giving the total number of pair excitation operators.

• [int02], [int04]

**Type:** Int (a blank parameter is not allowed)

**Description**: An integer giving a site index  $(0 \le [int02], [int04] < Nsite)$ .

• [int03], [int05]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving a spin index:

For Hubbard or Kondo system, the index can be selected from

0: Up-spin,

1: Down-spin.

For Spin system, the index can be selected from

 $0, 1, \dots, 2S + 1$  (corresponding to  $-S - 0.5, -S + 0.5, \dots S + 0.5$ ).

• [int06]

Type: Int (a blank parameter is not allowed)

**Description**: An integer giving a type of pair excitation operators:

 $0: c_{i\sigma_1} c_{j\sigma_2}^{\dagger}$ 

1:  $c_{i\sigma_1}^{\dagger}c_{j\sigma_2}$ .

 $\bullet$  [double01], [double02]

**Type:** Double (a blank parameter is not allowed)

**Description:** [double01] gives the real part of  $c_{i\sigma_1}c_{j\sigma_2}^{\dagger}(c_{i\sigma_1}^{\dagger}c_{j\sigma_2})$ , while [double02] gives the imaginary part of  $c_{i\sigma_1}c_{j\sigma_2}^{\dagger}(c_{i\sigma_1}^{\dagger}c_{j\sigma_2})$ .

#### Use rules

- Headers cannot be omitted.
- A program is terminated when the components of the pair excitation operators are double counted.
- A program is terminated when the conditions i = j and k = l are not satisfied for the spin model.
- A program is terminated when [int01] is different from the total number of two-body Green's functions defined in this file.
- A program is terminated when [int02]-[int06] are outside the range of the defined values.

## 4.2.18 SpectrumVec File

The header of the input file for the initial vector to calculate the dynamical Green's functions is defined. The file name and the file format (binary type) are as follows.

## File name

• ##\_rank\_\$\$.dat

## is the name of the head indicated by the key word SpectrumVec in the CalcMod file, and \$\$ is the number of the rank.

#### File format

• Line 1: [int01]

• Line 2: [int02]

• Lines 3 -: [double01] [double02].

#### **Parameters**

• [int01]

Type: Int

**Description:** The total number of the targets of the Hilbert spaces.

• [int02]

Type: Int

**Description:** The number of Lanczos or TPQ steps.

• [double01], [double02]

Type: double

**Description:** The coefficient value of the input vector. [double01] is a real part and [double02] is an imaginary part.

# 4.3 Output files

In this section, the details of the output files of the expert mode are explained.

## 4.3.1 CHECK\_Chemi.dat

This file is outputted to check the input of chemical potential  $\mu_{i\sigma}$ ,

$$H + = \sum_{i,\sigma} \mu_{i\sigma} c_{i\sigma}^{\dagger} c_{i\sigma}. \tag{4.16}$$

An example of the file format is as follows.

```
i=0 spin=0 isite1=1 tmp_V=0.000000
i=1 spin=0 isite1=2 tmp_V=0.000000
i=2 spin=0 isite1=3 tmp_V=0.000000
i=3 spin=0 isite1=4 tmp_V=0.000000
i=4 spin=0 isite1=5 tmp_V=0.000000
i=5 spin=0 isite1=6 tmp_V=0.000000
...
```

## File format

• i=[int01] spin=[int02] isite1=[int03] tmp\_V=[double01]

## **Parameters**

• [int01]

Type: Int

**Description:** The counted number of inputting terms.

• [int02]

Type: Int

**Description**: An integer for showing the spin index of  $\mu_{i\sigma}$ :

0: Up-spin,1: Down-spin.

• [int03]

Type: Int

**Description**: An integer for showing the site index of  $\mu_{i\sigma}$ .

• [double01]

Type: Double

**Description**: A value for  $\mu_{i\sigma}$ .

## 4.3.2 CHECK\_InterAll.dat

This file is outputted to check the input of the diagonal components of general two-body interactions,

$$H + = \sum_{i,j,\sigma} I_{iijj\sigma_1\sigma_1\sigma_2\sigma_2} c_{i\sigma_1}^{\dagger} c_{i\sigma_1} c_{i\sigma_2}^{\dagger} c_{i\sigma_2}. \tag{4.17}$$

An example of the file format is as follows.

```
i=0 isite1=1 A_spin=0 isite2=2 B_spin=0 tmp_V=0.500000
i=1 isite1=1 A_spin=0 isite2=2 B_spin=1 tmp_V=-0.500000
i=2 isite1=1 A_spin=1 isite2=2 B_spin=0 tmp_V=-0.500000
i=3 isite1=1 A_spin=1 isite2=2 B_spin=1 tmp_V=0.500000
i=4 isite1=2 A_spin=0 isite2=3 B_spin=0 tmp_V=0.500000
i=5 isite1=2 A_spin=0 isite2=3 B_spin=1 tmp_V=-0.500000
...
```

#### File format

• i=[int01] isite1=[int02] A\_spin=[int03] isite2=[int04] B\_spin=[int05]  $tmp_V=[double01]$ 

#### **Parameters**

• [int01]

Type: Int

**Description**: The counted number of inputting terms.

• [int02], [int04]

Type: Int

**Description :** An integer for showing the site index of  $I_{iijj\sigma_1\sigma_1\sigma_2\sigma_2}$ . [int02] and [int04] correspond to i and j, respectively.

• [int03], [int05]

Type: Int

**Description**: An integer for showing the spin index of  $I_{iijj\sigma_1\sigma_1\sigma_2\sigma_2}$ :

0: Up-spin

1: Down-spin.

[int03] and [int05] correspond to  $\sigma_1$  and  $\sigma_2$ , respectively.

• [double01]

Type: Double

**Description**: A value for  $I_{iijj\sigma_1\sigma_1\sigma_2\sigma_2}$ .

## 4.3.3 CHECK\_CoulombIntra.dat

This file is outputted to check the input of the on-site interactions  $U_i$ ,

$$H + = \sum_{i} U_{i} n_{i\uparrow} n_{j\downarrow}. \tag{4.18}$$

An example of the file format is as follows.

## File format

• i=[int01] isite1=[int02]  $tmp_V=[double01]$ 

#### **Parameters**

• [int01]

Type: Int

**Description :** The counted number of inputting terms.

• [int02]

Type: Int

**Description**: An integer for showing the site index of  $U_i$ .

• [double01]

Type: Double

**Description**: A value for  $U_i$ .

## 4.3.4 CHECK\_Hund.dat

This file is outputted to check the input of the Hund couplings  $J_{ij}^{\text{Hund}}$ ,

$$H + = -\sum_{i,j} J_{ij}^{\text{Hund}} (n_{i\uparrow} n_{j\uparrow} + n_{i\downarrow} n_{j\downarrow}). \tag{4.19}$$

An example of the file format is as follows.

```
i=0 isite1=1 isite2=2 tmp_V=0.250000
i=1 isite1=2 isite2=3 tmp_V=0.250000
i=2 isite1=3 isite2=4 tmp_V=0.250000
i=3 isite1=4 isite2=5 tmp_V=0.250000
i=4 isite1=5 isite2=6 tmp_V=0.250000
i=5 isite1=6 isite2=1 tmp_V=0.250000
```

## File format

• i=[int01] isite1=[int02] isite2=[int03]  $tmp_V=[double01]$ 

## **Parameters**

• [int01]

Type: Int

**Description:** The counted number of inputting terms.

• [int02], [int03]

Type: Int

**Description :** An integer for showing the site index of  $J_{ij}^{\text{Hund}}$ . [int02] and [int03] correspond to i and j, respectively.

• [double01]

Type: Double

**Description :** A value for  $J_{ij}^{\text{Hund}}$ .

## 4.3.5 CHECK\_INTER\_U.dat

This file is outputted to check the input of the diagonal components of the off-site interactions  $V_{ij}$ ,

$$H + = \sum_{i} V_{ij} n_i n_j \tag{4.20}$$

An example of the file format is as follows.

```
i=0 isite1=1 isite2=2 tmp_V=-0.125000
i=1 isite1=2 isite2=3 tmp_V=-0.125000
i=2 isite1=3 isite2=4 tmp_V=-0.125000
i=3 isite1=4 isite2=5 tmp_V=-0.125000
i=4 isite1=5 isite2=6 tmp_V=-0.125000
i=5 isite1=6 isite2=1 tmp_V=-0.125000
```

#### File format

• i=[int01] isite1=[int02] isite2=[int03]  $tmp_V=[double01]$ 

#### **Parameters**

• [int01]

Type: Int

**Description:** The counted number of inputting terms.

• [int02], [int03]

Type: Int

**Description :** An integer giving the site index of  $V_{ij}$ . [int02] and [int03] correspond to i and j, respectively.

• [double01]

Type: Double

**Description**: A value for  $V_{ii}$ .

## 4.3.6 CHECK\_Memory.dat

This file shows the size of the memory used in the calculation. An example of the file format is as follows.

```
MAX DIMENSION idim_max=400
REQUIRED MEMORY max_mem=0.000019 GB
```

## File format

- MAX DIMENSION idim\_max=[int01]
- REQUIRED MEMORY max\_mem =[double01] GB

#### **Parameters**

• [int01]

Type: Int

**Description:** An integer to show the total numbers of the Hilbert space under a calculation.

• [double01]

Type: Double

**Description :** The size of memory to store the Hilbert space in a calculation (GB unit).

## 4.3.7 WarningOnTransfer.dat

This file shows the double counted components of transfer integrals. An example of the file format is as follows.

```
double conuntings in transfers: i=0 j=2 spni 0 spnj 0 double conuntings in transfers: i=2 j=0 spni 0 spnj 0 double conuntings in transfers: i=0 j=2 spni 1 spnj 1 double conuntings in transfers: i=2 j=0 spni 1 spnj 1
```

#### File format

• Double countings in transfers: i=[int01] j=[int02] spni [int03] spnj [int04]

#### **Parameters**

• [int01], [int02]

Type: Int

**Description :** The integer of the site number where the transfer integrals are double counted.

• [int03], [int04]

Type: Int

**Description**: The integer of the spin index of a transfer integral:

0: Up-spin1: Down-spin.

# 4.3.8 CalcTimer.dat

The name of the calculation process, the process number, and the calculation process time are outputted in order at each line in the CalcTime.dat file. An example of the file format for the TPQ method is as follows.

All	[0000]	12.94052
SZ	[1000]	0.01795
diagonalcalc	[2000]	0.00693
CalcByTPQ	[3000]	12.90670
FirstMultiply	[3100]	0.08416
rand in FirstMultiply	[3101]	0.00172
mltply in FirstMultiply	[3102]	0.07707
expec_energy_flct	[3200]	9.06255
<pre>calc flctuation in expec_energy_flct</pre>	[3201]	1.67779
<pre>mltply in expec_energy_flct</pre>	[3202]	7.31207
expec_onebody	[3300]	0.11640
expec_twobody	[3400]	3.28796
Multiply	[3500]	0.14840
FileIO	[3600]	0.20493
	:	
All mltply	[0001]	7.38883
diagonal	[0100]	0.04153
Hubbard	[0300]	7.34636
trans in Hubbard	[0310]	7.34595
double	[0311]	0.00000
single	[0312]	0.00000
inner	[0313]	7.34299
interall in Hubbard	[0320]	0.00008
interPE	[0321]	0.00000
inner	[0322]	0.00000
pairhopp in Hubbard	[0330]	0.00006
interPE	[0331]	0.00000
inner	[0332]	0.00000
exchange in Hubbard	[0340]	0.00004
interPE	[0341]	0.00000
inner	[0342]	0.00000
	:	

## 4.3.9 TimeKeeper.dat

This file is outputted to show the calculation process information. An example of the file format for the Lanczos method is as follows.

```
diagonal calculation finishes: Wed Sep 16 22:58:49 2015
Lanczos Eigen Value start: Wed Sep 16 22:58:49 2015
1 th Lanczos step: Wed Sep 16 22:58:49 2015
...

122 th Lanczos step: Wed Sep 16 22:58:49 2015
Lanczos Eigenvalue finishes: Wed Sep 16 22:58:49 2015
Lanczos Eigenvector finishes: Wed Sep 16 22:58:49 2015
Lanczos expec energy finishes: Wed Sep 16 22:58:49 2015
CG Eigenvector finishes: Wed Sep 16 22:58:49 2015
CG expec energy finishes: Wed Sep 16 22:58:50 2015
CG expec_cisajs finishes: Wed Sep 16 22:58:50 2015
CG expec_cisajacktalt begins: Wed Sep 16 22:58:50 2015
```

#### File name

• ##\_TimeKeeper.dat

## indicates a header defined by [string02] in a ModPara file.

## 4.3.10 sz\_TimeKeeper.dat

This file is outputted to show the process information to obtain the Hilbert space needed for the calculation. An example of the file format is as follows.

```
initial sz : Wed Sep 16 22:58:49 2015
num_threads==4
omp parallel sz finishes: Wed Sep 16 22:58:49 2015
mid omp parallel sz : Wed Sep 16 22:58:49 2015
omp parallel sz finishes: Wed Sep 16 22:58:49 2015
```

#### File name

• ##\_sz\_TimeKeeper.dat

## indicates a header defined by [string02] in a ModPara file.

## 4.3.11 Time\_CG\_EigenVector.dat

(For the Lanczos method) The process for calculating the eigenvector by the CG method is outputted. An example of the file format is as follows.

```
allocate succeed !!!
b[4341]=1.000000 bnorm== 1.000000
i_itr=0 itr=5 0.0411202543 0.0000100000
i_itr=0 itr=155 0.0000000058 0.0000100000
CG OK:
         t_itr=155
i_itr=0 itr=155 time=0.000000
fabs(fabs(xb)-1.0)=0.9955114473313577
b[4341]=0.004489 bnorm== 1.000000
i_itr=1 itr=5 13.0033983157 0.0000100000
. . .
CG OK:
         t_itr=275
i_itr=1 itr=120 time=0.000000
fabs(fabs(xb)-1.0)=0.0000000000001295
number of iterations in inv1:i_itr=1 itr=120
t_itr=275 0.000000
```

#### File name

• ##\_Time\_CG\_EigenVector.dat

## indicates a header defined by [string02] in a ModPara file.

## 4.3.12 energy.dat

(For the Lanczos method) The values of the energy, doublon, and  $\langle S_z \rangle$  calculated by using the eigenvector obtained by the Lanczos or CG method are outputted. An example of the file format is as follows.

#### For method="Lanczos"

```
Energy -7.1043675920
Doublon 0.4164356536
Sz 0.0000000000
```

## For method="CG"

```
State 0
    Energy -7.1043675920
    Doublon 0.4164356536
    Sz 0.0000000000

State 1
:
```

#### File name

• ##\_energy.dat

## indicates a header defined by [string02] in a ModPara file.

## File format

- Line 1: Energy [double01]
- Line 2: Doublon [double02]
- Line 3: Sz [double02].

## **Parameters**

• [double01]

Type: Double

**Description :** The value of the energy calculated by the eigenvetor obtained by the Lanczos or CG method.

• [double02]

Type: Double

**Description :** The value of the doublon calculated by the eigenvetor obtained by the Lanczos or CG method,  $\frac{1}{N_s} \sum_i \langle n_{i\uparrow} n_{i\downarrow} \rangle$  ( $N_s$  is the total number of sites).

# • [double03]

 $\mathbf{Type}: Double$ 

**Description :** The value of  $S_z$  calculated by the eigenvetor obtained by the Lanczos or CG method.

## 4.3.13 Lanczos\_Step.dat

(For the Lanczos method) This file is outputted to show the process information for calculating the eigenvector by Lanczos method. An example of the file format for the Lanczos method is shown as follows.

#### For method="Lanczos"

#### For method="CG"

(	Step	Residual-2-norm	Threshold	Energy			
	1	6.79819e+00	8.19743e-07	7.86586e+00	8.19743e+00	8.02804e+00	
	2	7.47402e+00	3.69905e-07	3.35827e+00	3.63546e+00	3.69905e+00	
	3	5.30943e+00	2.44472e-07	-2.44472e+00	-2.23296e+00	-1.95487e+00	
	4	4.52737e+00	5.10297e-07	-5.10297e+00	-4.92390e+00	-4.58682e+00	
	5	3.66168e+00	7.14105e-07	-7.14105e+00	-6.91226e+00	-6.44532e+00	
	6	3.12717e+00	8.27201e-07	-8.27201e+00	-7.93262e+00	-7.44680e+00	
	152	1.05602e-06	1.04544e-06	-1.04544e+01	-9.89605e+00	-9.89605e+00	
	153	1.07401e-06	1.04544e-06	-1.04544e+01	-9.89605e+00	-9.89605e+00	
	154	9.45018e-07	1.04544e-06	-1.04544e+01	-9.89605e+00	-9.89605e+00	
/	_						

## File name

• ##\_Lanczos\_Step.dat

## indicates a header defined by [string02] in a ModPara file.

#### File format

- For method="Lanczos" stp= [int01] [double01] [double02] [double03] [double04] [double-a] [double-b]
- For method="CG"

  [int01] [double-c] [double-d] [double01] [double02] ...

#### **Parameters**

• [int01]

Type: Int

**Description:** The iteration number of the Lanczos and LOBCG method.

• [double01], [double02], [double03], [double04] ...

Type: Double

**Description:** Eigenvalues computed with the Lanczos or LOBCG method (ascending order). Four and exct eigenvalues are printed for the Lanczos and LOBCG method, respectively (in the above case, exct=3). While the degenerate eigenstates are printed as a single state in the Lanczos method, they are printed separately in LOBCG method. In the above case, we can find there is a degeneracy in the first excited state.

• [double-a]

Type: Double

**Description:** (Only for the Lanczos method) The eigenvalue used for the convergence check. It was specified by LanczosTarget (in the above case, LanczosTarget=3).

• [double-b]

Type: Double

**Description:** (Only for the Lanczos method) The maximum eigenvalue divided by the number of sites. It is the lower limit of LargeValue in TPQ method.

• [double-c]

Type: Double

**Description:** (Only for LOBCG method) The maximum of the 2-norm of each residual vector. It is used for the convergence check.

• [double-d]

Type: Double

**Description:** (Only for LOBCG method) The convergence threshold. This is obtained by the value specified with LanczosEps times the absolute value of the energy of the ground state.

## 4.3.14 Time\_TPQ\_Step.dat

(For the TPQ method) This file is outputted to show the time for starting the calculation of the TPQ method at each seed and step. In the restart calculation, the values are added to the previous file. An example is as follows.

```
set 0 step 1:TPQ begins: Wed Jul 13 07:59:20 2016
set 0 step 2:TPQ begins: Wed Jul 13 07:59:20 2016
set 0 step 3:TPQ begins: Wed Jul 13 07:59:20 2016
...
set 4 step 1997:TPQ begins: Wed Jul 13 07:59:32 2016
set 4 step 1998:TPQ begins: Wed Jul 13 07:59:32 2016
set 4 step 1999:TPQ begins: Wed Jul 13 07:59:32 2016
```

#### File name

•  $##_TPQ_Step.dat$ 

## indicates a header defined by [string02] in a ModPara file.

#### File format

• Set [int01] stp [int02]: TPQ begins: [string01]

#### **Parameters**

• [int01]

Type: Int

**Description**: The seed number in the calculation of the TPQ method.

• [int02]

Type: Int

**Description**: The step number in the calculation of the TPQ method.

• [string01]

Type: String

**Description :** The time for starting the calculation of the TPQ method at each seed and step.

## 4.3.15 Norm\_rand.dat

(For the TPQ method) This file is outputted to show the calculation process information for the TPQ method. In the restart calculation, the values are added to the previous file. An example of the file format is as follows.

```
# inv_temp, global_norm, global_1st_norm, step_i
0.017471 19.046586 11.288975 1
0.034863 19.089752 11.288975 2
...
31.999572 20.802362 11.288975 1997
32.015596 20.802362 11.288975 1998
32.031620 20.802362 11.288975 1999
```

#### File name

• Norm\_rand??.dat

?? indicates the number of runs in the calculation of the TPQ method.

#### File format

• Line 1: Header

• Lines 2-: [double01] [double02] [double03] [int01].

#### **Parameters**

• [double01]

Type: Double

**Description :** Inverse temperature  $1/k_{\rm B}T$ .

• [double02]

Type: Double

**Description :** The norm of a wave function before normalization given by  $\langle \tilde{\psi}_k | \tilde{\psi}_k \rangle$ , where  $|\tilde{\psi}_k \rangle \equiv (l - \hat{H}/N_s) |\psi_{k-1}\rangle$ .

• [double03]

Type: Double

**Description**: The norm of an initial wave function before normalization given by  $\langle \tilde{\psi}_0 | \tilde{\psi}_0 \rangle$ , where  $|\tilde{\psi}_0 \rangle$  is an initial random vector.

• [int01]

Type: Int

**Description :** The number of operations of  $(l - \hat{H}/N_s)$  for an initial wave function, where l is LargeValue defined in a ModPara file and  $N_s$  is the total number of sites.

## 4.3.16 SS\_rand.dat

(For the TPQ method) This file is outputted to show the calculation results for the TPQ method. In the restart calculation, the values are added to the previous file. An example of the file format is as follows.

```
# inv_tmp, energy, phys_var, phys_doublon, phys_num, step_i
0.017471  5.526334  45.390269  1.464589  6.000000  1
0.034863  5.266718  42.655559  1.434679  6.000000  2
...
31.999572  -4.814170  23.176231  0.590568  6.000000  1997
32.015596  -4.814170  23.176231  0.590568  6.000000  1998
32.031620  -4.814170  23.176231  0.590568  6.000000  1999
```

#### File name

• SS\_rand??.dat

?? indicates the number of runs in the calculation of the TPQ method.

#### File format

- Line 1: Header
- Lines 2-: [double01] [double02] [double03] [double04] [double05] [int01].

#### **Parameters**

• [double01]

Type: Double

**Description**: Inverse temperature  $1/k_{\rm B}T$ .

• [double02]

Type: Double

**Description :** The expected value of the energy  $\langle H \rangle$ .

• [double03]

Type: Double

**Description**: The expected value of the square of the Hamiltonian  $\langle H^2 \rangle$ .

• [double03]

Type: Double

**Description :** The expected value of the doublon,  $\frac{1}{N_s} \sum_i \langle n_{i\uparrow} n_{i\downarrow} \rangle$  ( $N_s$  is the total number of sites).

• [double05]

Type: Double

**Description :** The total number of particles  $\langle \hat{n} \rangle$ .

• [int01]

Type: Int

**Description :** The number of operations of  $(l - \hat{H}/N_s)$  for an initial wave function, where l is LargeValue defined in a ModPara file and  $N_s$  is the total number of sites.

## 4.3.17 Flct\_rand.dat

(For the TPQ method) This file is outputted to show the calculation results of the fluctuation of the particle number, doublon, and  $S_z$  for the TPQ method. In the restart calculation, the values are added to the previous file. An example of the file format is as follows.

```
# inv_temp, N, N^2, D, D^2, Sz, Sz^2, step_i
0.0826564 12.00 144.00 0.00 0.00 0.0009345626081113 0.2500 1
0.1639935 12.00 144.00 0.00 0.00 0.0023147006319775 0.2500 2
0.2440168 12.00 144.00 0.00 0.00 0.0037424057659867 0.2500 3
...
135.97669 12.00 144.00 0.00 0.00 -0.000000000167368 0.2500 1998
136.04474 12.00 144.00 0.00 0.00 -0.000000000165344 0.2500 1999
```

#### File name

• Flct\_rand??.dat

?? indicates the number of runs in the calculation of the TPQ method.

### File format

- Line 1: Header
- $\bullet$  Lines 2-: [double01] [double02] [double03] [double04] [double05] [double06] [double07] [int01].

#### **Parameters**

• [double01]

Type: Double

**Description**: Inverse temperature  $1/k_{\rm B}T$ .

• [double02]

Type: Double

**Description**: A total particle number  $\sum_{i} \langle \hat{n}_i \rangle$ .

• [double03]

Type: Double

**Description :** The expected value of the square of the particle number  $\sum_{i} \langle \hat{n}_{i}^{2} \rangle$ .

• [double04]

Type: Double

**Description :** The expected value of doublon  $\frac{1}{N_s} \sum_i \langle n_{i\uparrow} n_{i\downarrow} \rangle$  ( $N_s$  is the total number of sites).

• [double05]

Type: Double

**Description :** The expected value of the square of doublon  $\frac{1}{N_s} \sum_i \langle (n_{i\uparrow} n_{i\downarrow})^2 \rangle$   $(N_s \text{ is the total number of sites}).$ 

• [double06]

Type: Double

**Description :** The expected value of  $S_z \frac{1}{N_s} \sum_i \langle \hat{S}_i^z \rangle$  ( $N_s$  is the total number of sites).

• [double07]

Type: Double

**Description**: The expected value of the square of  $S_z \frac{1}{N_s} \sum_i \langle (\hat{S}_i^z)^2 \rangle$  ( $N_s$  is the total number of sites).

• [int01]

Type: Int

**Description:** The number of operations of  $(l - \hat{H}/N_s)$  for an initial wave function, where l is LargeValue defined in a ModPara file and  $N_s$  is the total number of sites.

## 4.3.18 Eigenvalue.dat

(For the FullDiag method) This file is outputted to show the energies calculated by the FullDiag method. An example of the file format is as follows.

```
0 -4.8141698096

1 -3.7968502453

2 -3.2462822372

...

397 13.9898305290

398 14.4896221034

399 14.8525199079
```

## File format

• [int01] [double01]

## **Parameters**

• [int01]

Type: Int

**Description:** The index of eigenvalues. The index 0 is for the energy of the ground state and the indexes are labeled in descending order for energies.

• [double01]

Type: Double

**Description :** The expected value of energy  $\langle H \rangle$ .

## 4.3.19 phys.dat

(For the FullDiag method) This file is outputted to show the physical values calculated by the FullDiag method. The data are outputted in descending order for energies. An example of the file format is as follows.

> 4.814170 3.796850	<n> 0.000000 0.000000</n>	<sz> 0.000000 0.000000</sz>	<s2> -0.000000 1.333333</s2>	<d> 0.590568 0.423804</d>
 .489622 .852520	0.000000	0.000000	0.000000	2.550240 2.329157

#### File name

- Canonical ensemble: ##\_phys\_Nup\_\$\$Ndown%%.dat
- Grand canonical ensemble: ##\_phys.dat.

##, \$\$, and %% indicate [string02], Nup, and Ndown defined in a ModPara file, respectively.

#### File format

- Line 1: Header
- Lines 2-: [double01] [double02] [double03] [double04] [double05].

#### **Parameters**

• [double01]

Type: Double

**Description :** The energy  $\langle H \rangle$ .

• [double02]

Type: Double

**Description :** The total number of particles  $\langle \hat{n} \rangle$ .

• [double03]

Type: Double

**Description :** The expected value of  $S_z$ ,  $\langle S_z \rangle$ .

• [double04]

Type: Double

**Description**: The expected value of  $S^2$ ,  $\langle S^2 \rangle$ .

## • [double05]

 $\mathbf{Type}: Double$ 

**Description :** The expected value of doublon,  $\frac{1}{N_s} \sum_i \langle n_{i\uparrow} n_{i\downarrow} \rangle$  ( $N_s$  is the total number of sites).

#### 4.3.20 ham.dat

(For the FullDiag method) When OutputHam=1 in the CalcMod file, the Hamiltonian calculated by  $\mathcal{H}\Phi$  is outputted by the MatrixMarket format. The recalculation by using this file can be down when InputHam=1 in the CalcMod file. An example of the file format is as follows.

```
%%%%MatrixMarket matrix coordinate complex hermitian
28 28 56
1 1 1.000000 0.000000
2 1 0.500000 0.000000
3 2 0.500000 0.000000
4 3 0.500000 0.000000
5 4 0.500000 0.000000
6 5 0.500000 0.000000
7 6 0.500000 0.000000
7 7 1.000000 0.000000
...
```

#### File name

• ##\_ham.dat

## indicates [string02] in a ModPara file.

#### File format

• Line 1: Header

• Line 2: [int01] [int02] [int03]

• Lines3-: [int04] [int05] [double01] [double02]

#### **Parameters**

• [int01]

Type: Int

**Description:** Number of rows of Hamiltonian.

• [int02]

Type: Int

**Description:** Number of columns of Hamiltonian.

• [int03]

Type: Int

**Description:** Number of nonzero elements of Hamiltonian.

 $\bullet$  [double01], [double02]

Type: Double

**Description :** The value of Hamiltonian; [double01] and [double02] represent real and imaginary part of the Hamiltonian, respectively.

## 4.3.21 cisajs.dat

This file is the outputted files for one-body Green's function  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} \rangle$ . The target components are set in the input file with the keyword "OneBodyG". An example of the file format is as follows.

```
0
     0
               0 0.4452776740 0.0000000000
0
     1
          0
               1 0.4452776740 0.0000000000
1
               0 0.500000000 0.0000000000
          1
1
     1
          1
               1 0.500000000 0.0000000000
2
          2
               0 0.4452776740 0.0000000000
     0
2
          2
               1 0.4452776740 0.0000000000
     1
3
          3
               0 0.500000000 0.0000000000
     0
               1 0.5000000000 0.0000000000
3
```

#### File name

• Lanczos method: ##\_cisajs.dat

• TPQ method: ##\_cisajs\_set??step%%.dat

• Full diagonalization method, LOBCG method: ##\_cisajs\_eigen&&.dat.

##, ??, %%, and && indicate [string02] in a ModPara file, the number of runs in calculation in the TPQ method, the number of steps in the TPQ method, and the index of the eigenvalues, respectively.

#### File format

• [int01] [int02] [int03] [int04] [double01] [double02]

#### **Parameters**

• [int01], [int03]

Type: Int

**Description:** The integer of the site number. [int01] and [int03] show the i and j site numbers, respectively.

• [int02], [int04]

Type: Int

**Description :** The integer of the spin index:

0: Up-spin1: Down-spin.

[int02] and [int04] show  $\sigma_1$  and  $\sigma_2$ , respectively.

 $\bullet$  [double01], [double02]

 $\mathbf{Type}: \mathbf{Double}$ 

**Description :** The value of  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} \rangle$ . [double01] and [double02] show the real and imaginary part of  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} \rangle$ , re-

spectively.

## 4.3.22 cisajscktalt.dat

This file is the outputted files for the two-body Green's function  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} c_{k\sigma_3}^{\dagger} c_{l\sigma_4} \rangle$ . The target components are set in the input file with the keyword "TwoBodyG". An example of the file format is as follows.

	0	0	0	0	0	0	0	0 0.4452776740 0.0000000000
	0	0	0	0	0	1	0	1 0.1843355815 0.0000000000
	0	0	0	0	1	0	1	0 0.1812412105 0.0000000000
	0	0	0	0	1	1	1	1 0.2640364635 0.0000000000
	0	0	0	0	2	0	2	0 0.0279690007 0.0000000000
	0	0	0	0	2	1	2	1 0.2009271524 0.0000000000
	0	0	0	0	3	0	3	0 0.2512810778 0.0000000000
	0	0	0	0	3	1	3	1 0.1939965962 0.0000000000
/								

#### File name

- Lanczos method: ##\_cisajscktalt.dat
- TPQ method: ##\_cisajscktalt\_set??step%%.dat
- Full diagonalization method, LOBCG method: ##\_cisajscktalt\_eigen&&.dat

##, ??, %%, and && indicate [string02] in a ModPara file, the number of runs in calculation in the TPQ method, the number of steps in the TPQ method, and the index of the eigenvalues, respectively.

#### File format

 $\bullet$  [int01] [int02] [int03] [int04] [int05] [int06] [int07] [int08] [double01] [double02].

#### **Parameters**

• [int01], [int03], [int05], [int07]

Type: Int

**Description:** The integer of the site number. [int01], [int03], [int05], and [int07] show the i, j, k, and l site numbers, respectively.

• [int02], [int04], [int06], [int08]

Type: Int

**Description:** The integer of the spin index:

0: Up-spin1: Down-spin.

[int02], [int04], [int06], and [int08] show  $\sigma_1$ ,  $\sigma_2$ ,  $\sigma_3$ , and  $\sigma_4$ , respectively.

 $\bullet$  [double01], [double02]

 $\mathbf{Type}: Double$ 

**Description :** The value of  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} c_{k\sigma_3}^{\dagger} c_{l\sigma_4} \rangle$ . [double01] and [double02] show the real and imaginary part of  $\langle c_{i\sigma_1}^{\dagger} c_{j\sigma_2} c_{k\sigma_3}^{\dagger} c_{l\sigma_4} \rangle$ ,

respectively.

## 4.3.23 eigenvec.dat

When OutputEigenVec=1 in a CalcMod file, the eigenvectors calculated by the Lanczos method are outputted. When InputEigenVec=1 in a CalcMod file, eigenvectors are inputted by this outputted file. The file format is of the binary type.

#### File name

• ##\_eigenvec\_&&\_rank\_\$\$.dat

## indicates [string02] in a ModPara file, && is the number of eigenvalues, and \$\$ is a number of rank.

#### File format

This file is written through the following source code (a little different fron the actual  $\mathcal{H}\Phi$  source).

```
fp = fopen("zvo_eigenvec_0_rank_0.dat", "wb");
fwrite(&number_of_interations, sizeof(int), 1,fp);
fwrite(&local_size, sizeof(unsigned long int),1,fp);
fwrite(&eigen_vector[0], sizeof(complex double),local_size+1, fp);
fclose(fp);
```

where number\_of\_interations is the number of iterations, local\_size is the size of eigenvector (if MPI is used, it differs from the dimension of the Hilbert space), eigen\_vector is the (complex) eigenvector.

**Note:** The fist component of eigen\_vector (eigen\_vector[0]) is not used for calculation.

#### 4.3.24 tmpvec.dat

When ReStart=1, 2 in a CalcMod file, vectors after the calculation stops at an indicated step are outputted. The file format is of the binary type. An example of the file format is as follows.

#### File name

- Lanczos method: ##\_tmpvec\_rank\_\$\$.dat
- TPQ and LOBPCG method: ##\_tmpvec\_set\_&&\_rank\_\$\$.dat .

## indicates [string02] in a ModPara file, and \$\$ is the number of rank. && is the sampling number for the TPQ calculation.

#### File format

This file is written through the following source code (a little different fron the actual  $\mathcal{H}\Phi$  source).

Lanczos

```
fp = fopen("zvo_tmpvec_rank_0.dat", "wb");
fwrite(&number_of_interations, sizeof(int), 1,fp);
fwrite(&local_size, sizeof(unsigned long int),1,fp);
fwrite(&last_vector[0], sizeof(complex double),local_size+1, fp);
fwrite(&second_last_vector[0], sizeof(complex double),local_size+1, fp);
fclose(fp);
```

#### TPQ and LOBPCG

```
fp = fopen("zvo_tmpvec_set_0_rank_0.dat", "wb");
fwrite(&number_of_interations, sizeof(int), 1,fp);
fwrite(&local_size, sizeof(unsigned long int),1,fp);
fwrite(&last_vector[0], sizeof(complex double),local_size+1, fp);
fclose(fp);
```

where number\_of\_interations is the number of iterations, local\_size is the size of eigenvector (if MPI is used, it differs from the dimension of the Hilbert space), last\_vector is the vector at the last iteration and second\_last\_vector is the vector at the second last iteration.

**Note:** The fist component of last\_vector and second\_last\_vector (last\_vector[0] and second\_last\_vector[0]) are not used for calculation.

#### 4.3.25 DynamicalGreen.dat

This file is the outputted file for calculating the dynamical Green's function. An example of the file format is as follows.

#### File name

• ##\_DynamicalGreen.dat

## indicates [string02] in a ModPara file.

#### File format

• Lines 1-: [double01] [double02] [double03] [double04]

#### **Parameters**

• [double01], [double02]

Type: Double

**Description:** The value of the frequency z. [double01] and [double02] are a real and an imaginary part of z, respectively.

• [double03], [double04]

Type: Double

**Description:** The value of dynamical Green's functions G(z). [double03] and [double04] are a real and an imaginary part of G(z), respectively.

#### 4.3.26 recalcvec.dat

This file is the outputted file for two vectors to recalculate the dynamical Green's function by the Lanczos method. The file format is of the binary type. An example of the file format is as follows.

#### File name

• ##\_recalcvec\_rank\_\$\$.dat

## indicates [string02] in a ModPara file and \$\$ is the number of rank.

#### File format

- Line 1: [int01]
- Line 2: [int02]
- Lines 3 3+[int02]: [double01] [double02]
- Lines  $4+[int02] 4+2\times[int02]$ : [double03] [double04].

#### **Parameters**

• [int01]

Type: Int

**Description :** The step for calculating dynamical Green's functions by the Lanczos method  $N_d$ .

• [int02]

Type: Long Int

**Description**: The total number of targets of the Hilbert spaces.

• [double01], [double02]

Type: Double

**Description:** The value of the vector  $v_{k+1}$  for recalculating dynamical Green's functions by the Lanczos method.

[double01] and [double02] are a real part and an imaginary part of  $v_{k+1}$ , respectively. The fist component is not used for calculation.

• [double03], [double04]

Type: Double

**Description :** The value of the vector  $v_k$  for recalculating dynamical Green's functions by the Lanczos method.

[double03] and [double04] are a real part and an imaginary part of  $v_k$ , respectively. The fist component is not used for calculation.

#### 4.3.27 TMcomponents.dat

This file is the outputted files for the components of the tridiagonal matrix and the norm of the excited state to recalculate the dynamical Green's function by the Lanczos method. The file format is of the binary type. An example of the file format is as follows.

#### File name

• ##\_TMcomponents.dat

## indicates [string02] in a ModPara file and \$\$ is the number of rank.

#### File format

• Line 1: [int01]

• Line 2: [double01]

• Lines 3 -: [double02] [double03].

#### **Parameters**

• [int01]

Type: Int

**Description :** The step for calculating dynamical Green's functions by the Lanczos method  $N_d$ .

• [double01]

Type: Double

**Description**: The value of the norm of the excited state.

• [double02], [double03]

Type: Double

**Description:** The value of the components of the tridiagonal matrix to recalculate dynamical Green's functions by the Lanczos method  $\alpha_i$ ,  $\beta_i$  ( $i = 1, \dots, N_d$ ). [double02] is  $\alpha_i$  and [double03] is  $\beta_i$ .

## 4.4 Error messages

• ERROR! Unsupported Keyword!

The program stops because unsupported keyword is specified.

- "ERROR! Keyword is duplicated!

  The program stops because a parameter is specified twice.
- ERROR ! Unsupported Solver : solver
- ullet ERROR ! Unsupported Model : model
- $\bullet$  Sorry, this system is unsupported in the STANDARD MODE... Please use the EXPART MODE, or write a NEW FUNCTION and post it us.

The program stops because unsupported parameter for method, model, or lattice is specified.

- ERROR ! abs(2 \* Sz) > nsite in Hubbard model !
- ERROR ! Nelec > 2 \* nsite in Hubbard model !
- ERROR ! (nelec + 2 \* Sz) % 2 != 0 in Hubbard model !
- ERROR ! nelec <= nsite && 2 \* |Sz| > nelec in Hubbard model !
- ERROR ! nelec > nsite && 2 \* |Sz| > 2 \* nsite nelec in Hubbard model !
- ERROR ! abs(2 \* Sz) > nsite in Spin model !
- ERROR ! (nsite + 2 \* Sz) % 2 != 0 in Spin model !
- ERROR ! abs(2 \* Sz) > nsite in Hubbard model !
- ERROR ! Nelec\_cond / 2 + Nelec\_loc > nsite in Kondo model !
- ERROR ! (nelec\_cond + nelec\_loc + 2 \* Sz) % 2 != 0 in Kondo model !
- ERROR ! nelec\_cond <= nsite / 2 && 2 \* |Sz| > nelec\_cond + nelec\_loc ...
- ERROR ! nelec\_cond > nsite / 2 && abs(Sz2) > nsite / 2 \* 3 nelec...

In the calculation of the canonical ensemble, there are some irrelevant combinations of the number of electrons, the number of sites, and the total spin moment (the number of electrons is larger twice than the number of sites); If these situations are detected, the program will stop.

• Check ! keyword is SPECIFIED but will NOT be USED.

Please COMMENT-OUT this line

or check this input is REALLY APPROPRIATE for your purpose !

Because an unnecessary parameter is specified, the program suggests checking the input file. If that parameter is actually unnecessary, please delete or comment out this line.

- ERROR ! *keyword* is NOT specified !

  The program stops because a prerequisite keyword is not specified.
- keyword = value ##### DEFAULT VALUE IS USED #####

  This is not an error message. The program states that the default value is used because this keyword is not specified.

# Algorithm

## 5.1 Lanczos method

#### 5.1.1 Details of Lanczos method

Some parts of this section are based on the manual of TITPACK [12] and the textbook published by M. Sugihara and K. Murota [13](these references are written in Japanese).

In the Lanczos method, by successively operating the Hamiltonian to the initial vector, we obtain the accurate eigenvalues around the maximum and minimum eigenvalues and associated eigenvectors. Because we can perform the Lanczos method by using only two vectors, the dimensions of which are the dimensions of the total Hilbert space\*1, the Lanczos method is frequently used for the diagonalization of the large matrices. As explained in detail below, one additional vector is necessary for obtaining the eigenvector.

The principle of the Lanczos method is based on the power method. In the power method, by successively operating the Hamiltonian  $\hat{\mathcal{H}}$  to the arbitrary vector  $\mathbf{x}_0$ , we generate  $\hat{\mathcal{H}}^n \mathbf{x}_0$ . The obtained space  $\mathcal{K}_{n+1}(\hat{\mathcal{H}}, \mathbf{x}_0) = \{\mathbf{x}_0, \hat{\mathcal{H}}^1 \mathbf{x}_0, \dots, \hat{\mathcal{H}}^n \mathbf{x}_0\}$  is called the Krylov subspace. The initial vector is represented by the superposition of the eigenvectors  $\mathbf{e}_i$  (the corresponding eigenvalues are  $E_i$ ) of  $\hat{\mathcal{H}}$  as

$$\boldsymbol{x}_0 = \sum_i a_i \boldsymbol{e}_i. \tag{5.1}$$

Here,  $E_0$  denotes the maximum absolute values of the eigenvalues. We note that all the eigenvalues are real numbers because the Hamiltonian is Hermitian. By operating  $\hat{\mathcal{H}}^n$  to the initial vector, we obtain the relation as

$$\hat{\mathcal{H}}^n \boldsymbol{x}_0 = E_0^n \left[ a_0 \boldsymbol{e}_0 + \sum_{i \neq 0} \left( \frac{E_i}{E_0} \right)^n a_i \boldsymbol{e}_i \right]. \tag{5.2}$$

This relation indicates that the eigenvector of  $E_0$  becomes dominant for sufficiently large n. In the Lanczos method, we obtain the eigenvalues and eigenvectors by performing the appropriate transformation for the obtained Krylov subspace.

In the Lanczos method, we successively generate the normalized orthogonal basis  $v_0, \ldots, v_{n-1}$  from the Krylov subspace  $\mathcal{K}_n(\hat{\mathcal{H}}, x_0)$ . We define an initial vector and

 $<sup>^{*1}</sup>$ In  $\mathcal{H}\Phi$ , to reduce the numerical cost, we use some additional vectors; a vector for accumulating the real-space diagonal elements of the Hamiltonian and a vector for specifying the given  $S_z$  space and given particle space. The dimension of these vectors is that of the Hilbert space.

118 5 Algorithm

associated components as  $\mathbf{v}_0 = \mathbf{x}_0/|\mathbf{x}_0|$ ,  $\beta_0 = 0$ ,  $\mathbf{x}_{-1} = 0$ . From this initial condition, we can obtain the normalized orthogonal basis:

$$\alpha_k = (\hat{\mathcal{H}}\boldsymbol{v}_k, \boldsymbol{v}_k), \tag{5.3}$$

$$\boldsymbol{w} = \hat{\mathcal{H}}\boldsymbol{v}_k - \beta_k \boldsymbol{v}_{k-1} - \alpha_k \boldsymbol{v}_k, \tag{5.4}$$

$$\beta_{k+1} = |\boldsymbol{w}|, \tag{5.5}$$

$$\boldsymbol{v}_{k+1} = \frac{\boldsymbol{v}_k}{|\boldsymbol{v}_k|}.\tag{5.6}$$

From these definitions, it it obvious that  $\alpha_k$ ,  $\beta_k$  are real numbers.

In the subspace spanned by these normalized orthogonal basis, the Hamiltonian is transformed as

$$T_n = V_n^{\dagger} \hat{\mathcal{H}} V_n. \tag{5.7}$$

Here,  $V_n$  is a matrix whose column vectors are  $\mathbf{v}_i (i=0,1,\ldots,n-1)$ .  $T_n$  is a tridiagonal matrix and its diagonal elements are  $\alpha_i$  and subdiagonal elements are  $\beta_i$ . It is known that the eigenvalues of  $\hat{\mathcal{H}}$  are well approximated by the eigenvalues of  $T_n$  for sufficiently large n. (We note that  $V^{\dagger}V = I,I$  is an identity matrix). The original eigenvectors of  $\hat{\mathcal{H}}$  are obtained by  $\mathbf{e}_i = V\tilde{\mathbf{e}}_i$ , where  $\tilde{\mathbf{e}}_i$  denotes the eigenvectors of  $T_n$ . From V, we can obtain the eigenvectors of  $\hat{\mathcal{H}}$  by performing the Lanczos method. However, in the actual calculations, it is difficult to keep V, because its dimension is large [dimension of  $V = (\text{dimension of the total Hilbert space}) \times (\text{the number of Lanczos iterations})$ ]. Thus, to obtain the eigenvectors, we again perform the same Lanczos calculations after we obtain the eigenvalues from the Lanczos methods. In the first Lanczos calculation, we keep  $\tilde{\mathbf{e}}_i$ , because its dimension is small\*2. From this procedure, we obtain the eigenvectors from V.

In the Lanczos method, within a few hundred or thousand Lanczos iterations, we obtain accurate eigenvalues near the maximum and minimum eigenvalues. The necessary number of iterations is sufficiently small as compared to the dimensions of the total Hilbert space. We note that it is shown that the errors of the maximum and minimum eigenvalues become exponentially small as a function of Lanczos iterations (for details, see Ref. [13]).

#### 5.1.2 Inverse iteration method

From the approximate value of the eigenvalues  $(E_n)$ , by successively operating  $(\hat{\mathcal{H}} - E_n)^{-1}$  to the initial vector  $\mathbf{y}_0$ , we can obtain the accurate eigenvector for  $E_n$ . From  $(\hat{\mathcal{H}} - E_n)^{-1}\mathbf{y}_0$ , we obtain linear simultaneous equations such as

$$\mathbf{y}_k = (\hat{\mathcal{H}} - E_n)\mathbf{y}_{k+1}. \tag{5.8}$$

By solving this equation using the conjugate gradient method (CG method), we obtain the eigenvector. From the obtained eigenvector, we can calculate the eigenvalues and correlation functions. We note that four additional vectors are necessary to perform the CG method. For a large system size, it may be impossible to allocate memory to the additional vectors.

<sup>\*2</sup>Upper bound of the dimensions of  $\tilde{e_i}$  is # of Lanczos iterations.

#### 5.1.3 Details of implementation

#### Initial vector

For the Lanczos method, an initial vector is specified with  $initial_iv(\equiv r_s)$  defined in an input file for Standard mode or a ModPara file for Expert mode. The type of initial vector can be selected as a real number or complex number by using InitialVecType in a ModPara file.

• For canonical ensemble and initial\_iv  $\geq 0$ 

A component of a target of the Hilbert space is given by

$$(N_{\rm dim}/2 + r_s)\% N_{\rm dim},$$
 (5.9)

where  $N_{\text{dim}}$  is the total number of the Hilbert spaces and  $N_{\text{dim}}/2$  is added to avoid selecting a special Hilbert space for a default value initial\_iv = 1. When the type of initial vector is selected as a real number, the coefficient value is given by 1, while when it is selected as a complex number, the value is given by  $(1+i)/\sqrt{2}$ .

• For a grand canonical ensemble or initial\_iv < 0

The initial vector is given by using a random generator, i.e., the coefficients of all the components for the initial vector are given by random numbers. The seed is calculated as

$$123432 + |r_s|, (5.10)$$

where  $r_s$  is the number given by an input file and  $n_{\text{run}}$  is the number of runs. The maximum value of  $n_{\text{run}}$  is defined by NumAve in an input file for Standard mode or a ModPara file for Expert mode. Random numbers are generated by using SIMD-oriented Fast Mersenne Twister (dSFMT) [14].

#### Convergence condition

In  $\mathcal{H}\Phi$ , we use **dsyev** (routine of LAPACK) for diagonalization of  $T_n$ . We use the energy of the first excited state of  $T_n$  as the criterion of convergence. In the standard setting, after five Lanczos steps, we diagonalize  $T_n$  every two Lanczos steps. If the energy of the first excited states coincides with the previous energy within the specified accuracy, the Lanczos iteration finishes. The accuracy of the convergence can be specified by CDataFileHead (ModPara file in the expert mode).

After obtaining the eigenvalues, we again perform the Lanczos iteration to obtain the eigenvector. From the eigenvectors  $|n\rangle$ , we calculate energy  $E_n = \langle n|\hat{\mathcal{H}}|n\rangle$  and variance  $\Delta = \langle n|\hat{\mathcal{H}}^2|n\rangle - (\langle n|\hat{\mathcal{H}}|n\rangle)^2$ . If  $E_n$  coincides with the eigenvalues obtained by the Lanczos iteration and  $\Delta$  is smaller than the specified value, we finish diagonalization.

If the accuracy of the Lanczos method is not sufficient, we perform the CG method to obtain the eigenvector. As an initial vector of the CG method, we use the eigenvectors obtained by the Lanczos method in the standard setting. This frequently accelerates the convergence.

120 5 Algorithm

## 5.2 Full Diagonalization method

#### 5.2.1 Overview

We generate the matrix of  $\hat{H}$  by using the real space configuration  $|\psi_j\rangle(j=1\cdots d_H)$ , where  $d_H$  is the dimension of the Hilbert space):  $H_{ij}=\langle\psi_i|\hat{H}|\psi_j\rangle$ . By diagonalizing this matrix, we can obtain all the eigenvalues  $E_i$  and eigenvectors  $|\Phi_i\rangle$   $(i=1\cdots d_H)$ . In the diagonalization, we use a LAPACK routine, such as dsyev or zheev. We also calculate and output the expectation values  $\langle A_i\rangle \equiv \langle \Phi_i|\hat{A}|\Phi_i\rangle$ . These values are used for the finite-temperature calculations.

## 5.2.2 Finite-temperature calculations

From  $\langle A_i \rangle \equiv \langle \Phi_i | \hat{A} | \Phi_i \rangle$ , we calculate the finite-temperature properties by using the relation

$$\langle \hat{A} \rangle = \frac{\sum_{i=1}^{N} \langle A_i \rangle e^{-\beta E_i}}{\sum_{i=1}^{N} e^{-\beta E_i}}.$$
 (5.11)

The calculation should be performed by using the own postscripts.

# 5.3 Finite-temperature calculations by the TPQ method

Sugiura and Shimizu showed that it is possible to calculate the finite-temperature properties from a few wavefunctions (in the thermodynamic limit, only one wave function is necessary) [5]. The wavefunction is called the thermal pure quantum (TPQ) state. Because the TPQ state can be generated by operating the Hamiltonian to the random initial wavefunction, we directly use the routine Lanczos method for the TPQ calculations. Here, we explain how to construct the micro canonical TPQ (mTPQ) state, which offers the simplest method for calculating finite-temperature properties.

Let  $|\psi_0\rangle$  be a random initial vector. By operating  $(l - \hat{H}/N_s)^k(l)$  is constant and  $N_s$  represents the number of sites) to  $|\psi_0\rangle$ , we obtain the kth TPQ states as

$$|\psi_k\rangle \equiv \frac{(l - \hat{H}/N_s)|\psi_{k-1}\rangle}{|(l - \hat{H}/N_s)|\psi_{k-1}\rangle|}.$$
(5.12)

From  $|\psi_k\rangle$ , we estimate the corresponding inverse temperature  $\beta_k$  as

$$\beta_k \sim \frac{2k/N_s}{l-u_k}, \quad u_k = \langle \psi_k | \hat{H} | \psi_k \rangle / N_s,$$
 (5.13)

where  $u_k$  is the internal energy. The arbitrary local physical properties at  $\beta_k$  are also estimated as

$$\langle \hat{A} \rangle_{\beta_k} = \langle \psi_k | \hat{A} | \psi_k \rangle / N_s.$$
 (5.14)

In a finite-size system, error is caused by the choice of the initial random vector. To estimate the average value and error of the physical properties, we perform some independent calculations by changing  $|\psi_0\rangle$ .

#### 5.3.1 Details of implementation

#### Initial vector

For the TPQ method, the initial vector is given by using a random generator, i.e., the coefficients of all the components for the initial vector are given by random numbers. The seed is calculated as

$$123432 + (n_{\text{run}} + 1) \times |r_s| + k_{\text{Thread}} + N_{\text{Thread}} \times k_{\text{Process}}, \tag{5.15}$$

where  $r_s$  is the number given by an input file and  $n_{\rm run}$  is the number of runs.  $r_s$  and the maximum value of  $n_{\rm run}$  are defined by initial\_iv and NumAve in an input file for Standard mode or a ModPara file for Expert mode, respectively. Random numbers are generated by using SIMD-oriented Fast Mersenne Twister (dSFMT) [14]. We can select the type of initial vector as a real number or complex number by using InitialVecType in a ModPara file.  $k_{\rm Thread}, N_{\rm Thread}, and k_{\rm Process}$  indicate the thread ID, number of threads, process ID, respectively; the initial vector depends both on initial\_iv and the number of parallelisms.

## 5.4 Bogoliubov representation

In the spin system, the spin indices in the input files of transfer, InterAll, and correlation functions are specified as those of the Bogoliubov representation. The spin operators are written by using creation/annihilation operators:

$$S_{iz} = \sum_{\sigma = -S}^{S} \sigma c_{i\sigma}^{\dagger} c_{i\sigma} \tag{5.16}$$

$$S_{i}^{+} = \sum_{\sigma=-S}^{S-1} \sqrt{S(S+1) - \sigma(\sigma+1)} c_{i\sigma+1}^{\dagger} c_{i\sigma}$$
 (5.17)

$$S_{i}^{-} = \sum_{\sigma=-S}^{S-1} \sqrt{S(S+1) - \sigma(\sigma+1)} c_{i\sigma}^{\dagger} c_{i\sigma+1}$$
 (5.18)

# 6

# Acknowledgement

We would like to express our sincere gratitude to Prof. Hidetoshi Nishimori and Mr. Daisuke Tahara. Implementation of the Lanczos algorithm in  $\mathcal{H}\Phi$  written in C is based on the pioneering diagonalization package TITPACK written in Fortran by Prof. Nishimori. For developing the user interface of  $\mathcal{H}\Phi$ , we followed the design concept of the user interface in the program for variational Monte Carlo developed by Mr. Tahara. A part of the user interface in  $\mathcal{H}\Phi$  is based on his original codes. We would also like to express our thanks for the support of the "Project for advancement of software usability in materials science" of The Institute for Solid State Physics, The University of Tokyo, for the development of  $\mathcal{H}\Phi$  ver. 0.1 - ver. 2.0.4.

Dr. Yuichi Motoyama and Dr. Yasuyuki Kato found and reported bugs. We acknowledge their kindness and carefulness.

We thank Prof. Takeo Hoshi and Prof. Tomohiro Sogabe for the fruitful discussions on the shifted Krylov method and the development of the  $K\omega$  library.

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