

Setup for gauss

2019/10/9

About gauss

1. Information of gauss can be seen in the terminal when you log in to gauss (For details on how to log in, see page 5 of the bookmark.).

```
# System

* Login node : gauss
* Computational nodes : gauss01 - gauss13
* CPU : Xeon E5-2680v4, 28 cores/node
* RAM : 64 GB/node
* Do not login directly the computational nodes.
  Use the queuing system.
* Sample of batch job script : /home/public/sample.sh

# Programs

* Binary : /home/public/bin/
* Source : /home/public/program/

# Scheduled Maintenance

* None
```

13 nodes
28 cores/node

2. Sample script (/home/public/sample.sh)

```
#!/bin/sh
#PBS -l nodes=2:ppn=7
#PBS -n
#
ulimit -m `expr 600000000 / ${PBS_NUM_PPN}`
ulimit -v `expr 600000000 / ${PBS_NUM_PPN}`
source ~/.bashrc
#
export OMP_NUM_THREADS=4
#
cd $PBS_O_WORKDIR
#
mpiexec -hostfile $PBS_NODEFILE /home/public/bin/openmx GaAs.dat -nt $OMP_NUM_THREADS
```

use 2 nodes and 7cores /node

use 4 threads for OpenMP

Submit a job

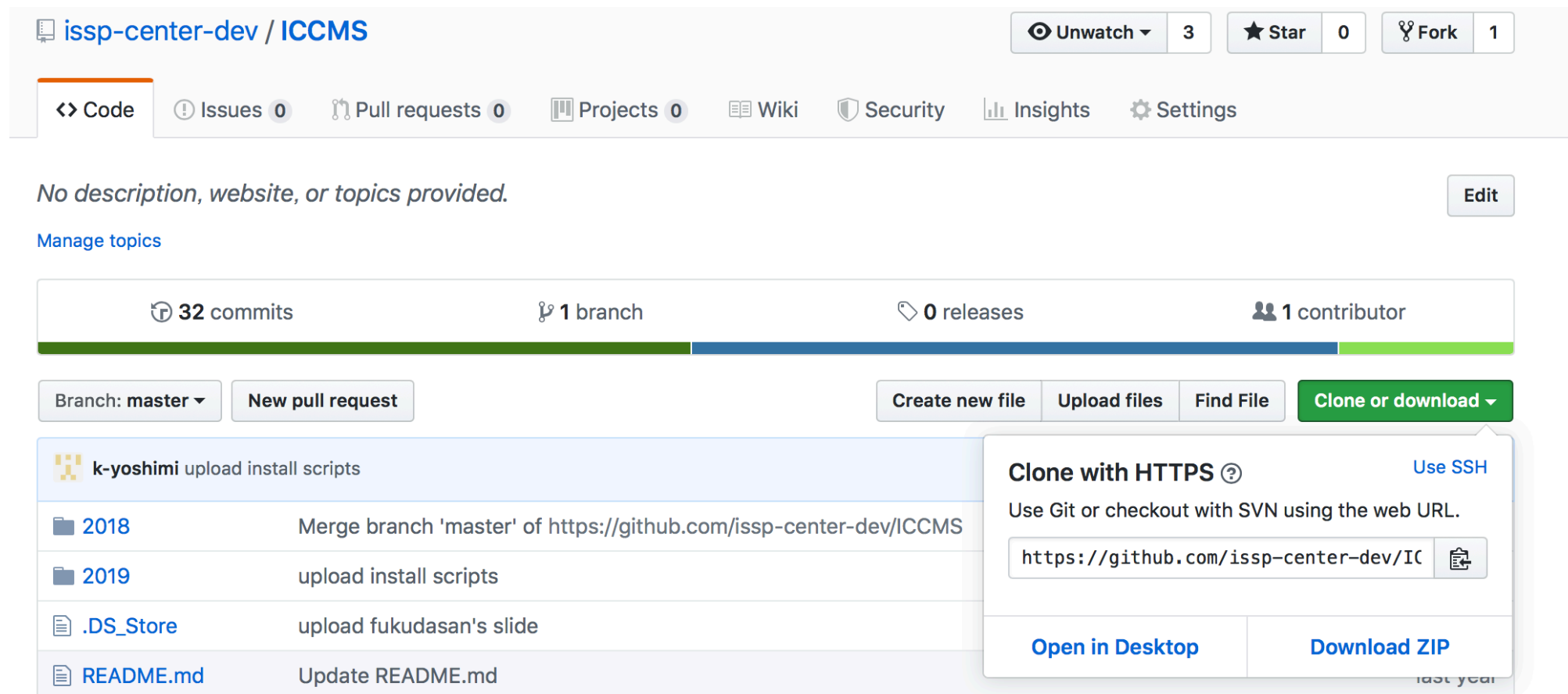
\$ qsub /home/public/sample.sh

If you do not know how to use job scheduler, ask Fukuda-sann (organizer).

Installation of COMBO and HΦ(1)

1. Copy URL of Github repository for ICCMS

- <https://github.com/issp-center-dev/ICCMS>



The screenshot displays the GitHub repository page for `issp-center-dev / ICCMS`. The repository has 32 commits, 1 branch, 0 releases, and 1 contributor. The 'Clone or download' button is highlighted, and a dropdown menu is open, showing the 'Clone with HTTPS' option. The URL `https://github.com/issp-center-dev/ICCMS` is displayed in the dropdown menu. The repository also includes a file list with folders for the years 2018 and 2019, and files for `.DS_Store` and `README.md`.

Click 「Clone or download」 and copy URL

Installation of COMBO and HΦ (2)

2. Git clone in terminal(Below https is the URL you copied).

- \$ git clone <https://github.com/issp-center-dev/ICCMS.git>

```
[guest20@gauss ~]$ git clone https://github.com/issp-center-dev/ICCMS.git
Initialized empty Git repository in /home/guest20/ICCMS/.git/
remote: Enumerating objects: 16, done.
remote: Counting objects: 100% (16/16), done.
remote: Compressing objects: 100% (14/14), done.
remote: Total 195 (delta 0), reused 16 (delta 0), pack-reused 179
Receiving objects: 100% (195/195), 104.99 MiB | 23.78 MiB/s, done.
Resolving deltas: 100% (37/37), done.
```

3. Make and change to tutorial directory and copy scripts.

- \$mkdir tutorial & cd tutorial
- \$cp ~/ICCMS/2019/2019-10-08/script/* .

Installation of COMBO and H Φ (3)

4. Install COMBO

- `$ sh install_combo.sh`

- `$ python`

- `import combo`

Check

```
[guest20@gauss ~]$ python
Python 2.7.15 (default, Sep 23 2019, 01:06:26)
[GCC Intel(R) C++ gcc 5.10 mode] on linux2
Type "help", "copyright", "credits" or "license" for more information.
>>> import combo
>>> exit()
```

5. Install H Φ

- `$ sh install_HPhi.sh`

- `$ cd HPhi.build && make test`

Check all tests are passed.

```
Start 48: te_spin_chain_interall
48/49 Test #48: te_spin_chain_interall ..... Passed    0.70 sec
Start 49: te_kondo_chain_interall
49/49 Test #49: te_kondo_chain_interall ..... Passed    0.71 sec

100% tests passed, 0 tests failed out of 49

Total Test time (real) = 40.93 sec
```

About H Φ , see <http://www.pasums.issp.u-tokyo.ac.jp/hphi/>


About H Φ (1)

From MateriApps

H Φ

Openness:3 ★★★ **Document quality:**2 ★★★

An exact diagonalization package for a wide range of quantum lattice models (e.g. multi-orbital Hubbard model, Heisenberg model, Kondo lattice model). H Φ also supports the massively parallel computations. The Lanczos algorithm for obtaining the ground state and thermal pure quantum state method for finite-temperature calculations are implemented. In addition, dynamical Green's functions can be calculated using [K \$\omega\$](#) , which is a library of the shifted Krylov subspace method. It is possible to perform simulations for real-time evolution from ver. 3.0.



Supported by

Project for Advancement of Software Usability in Materials Science
(<http://www.issp.u-tokyo.ac.jp/supercom/softwaredev>)

About $H\Phi$ (2)

Target substance/model Hubbard model, Heisenberg model, Kondo lattice model, Kitaev model, Kitaev-Heisenberg model, multi-orbital Hubbard model

Physical quantities that can be computed specific heat, susceptibility, ground state energy, free energy, structure factors

Methodology Lanczos algorithm, thermal pure quantum state, full diagonalization

For details, see

<http://www.pasums.issp.u-tokyo.ac.jp/hphi/en/>

Lecture slides :

<http://www.pasums.issp.u-tokyo.ac.jp/hphi/en/doc/presentation>