WIEN2k 24.1 Install with gfortran

December 14, 2024

Linux Operating System Version (lsb_release -a): Ubuntu 24.04.1 LTS Fortran Compiler Version (gfortran --version): gfortran 13.3.0 WIEN2k Version (cat \$WIENROOT/WIEN2k VERSION): 24.1 (Release 1/8/2024)

The Ubuntu 24.04.1 LTS was already installed, but if that is needed it should be available at:

https://ubuntu.com/download/desktop

Operating system packages that were needed were installed before proceeding with step 1 given below. If that hasn't been done, then install them: username@computername:~\$ sudo apt update

. . .

username@computername:~ \$ sudo apt upgrade

. .

username@computername:~ \$ sudo apt install tesh gfortran make autoconf libtool git

. . .

username@computername:~ \$ sudo apt install gnuplot gnuplot-x11 libgd-dev gedit

. .

username@computername:~\$ sudo apt install libquadmath0 build-essential libglu1-mesa-dev

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username@computername:~\$ sudo apt install libtogl-dev tcl-dev tk-dev libfftw3-dev libxmu-dev

. . .

Note: If prompted with "Do you want to continue [Y/n]", enter: Y.

The XCrySDen 1.6.2 was installed using:

username@computername:~ \$ sudo apt install xcrysden

Of note, for XCrySDen to work with Wayland, toglOpt(accum) needs set to false in the /usr/share/xcrysden/Tcl/custom-definitions file:

https://www.mail-archive.com/wien@zeus.theochem.tuwien.ac.at/msg23440.html

Restart the computer.

1. The following was entered into the terminal to install libxc:

username@computername:~/Desktop\$ cd ~ useusername@computername:~\$ wget https://gitlab.com/libxc/libxc/-/archive/7.0.0/libxc-7.0.0.tar.bz2

```
username@computername:~$ tar xvf libxc-7.0.0.tar.bz2
   username@computername:~$ cd libxc-7.0.0/
   username@computername:~/libxc-7.0.0$ autoreconf -i --force
   username@computername:~/libxc-7.0.0$ ./configure FC=gfortran CC=gcc
   --prefix=$HOME/libxc-7.0.0
   username@computername:~/libxc-7.0.0$ make
   username@computername:~/libxc-7.0.0$ make check
   PASS: xc-run testsuite
   Testsuite summary for libxc 7.0.0
   # TOTAL: 1
   # PASS: 1
   # SKIP: 0
   # XFAIL: 0
   # FAIL: 0
   # XPASS: 0
   # ERROR: 0
   username@computername:~/libxc-7.0.0$ make install
   username@computername:~/libxc-7.0.0$ ls ~/libxc-7.0.0/lib
   libxc.a libxcf03.a libxcf03.la libxc.la pkgconfig
2. The following was entered into the terminal to install OpenBLAS:
   username@computername:~/libxc-7.0.0$ cd ~
   username@computername:~$ wget
   https://github.com/OpenMathLib/OpenBLAS/releases/download/v0.3.28/
   OpenBLAS-0.3.28.tar.gz
   username@computername:~$ tar xvf OpenBLAS-0.3.28.tar.gz
   username@computername:~$ cd OpenBLAS-0.3.28/
   username@computername:~/OpenBLAS-0.3.28$ make FC=gfortran CC=gcc
```

OpenBLAS build complete. (BLAS CBLAS LAPACK LAPACKE)

```
... Linux
    OS
    Architecture
                 ... x86 64
                  ... 64bit
    BINARY
                 ... GCC (cmd & version : gcc (Ubuntu 13.3.0-6ubuntu2~24.04) 13.3.0)
    C compiler
    Fortran compiler ... GFORTRAN (cmd & version : GNU Fortran (Ubuntu 13.3.0-
   6ubuntu2~24.04) 13.3.0)
    Library Name ... libopenblas zenp-r0.3.28.a (Multi-threading; Max num-threads is 32)
   username@computername:~/OpenBLAS-0.3.28$ ls ~/OpenBLAS-0.3.28/libopenblas*
   /home/username/OpenBLAS-0.3.28/libopenblas.a
   /home/username/OpenBLAS-0.3.28/libopenblas.so
   /home/username/OpenBLAS-0.3.28/libopenblas.so.0
   /home/username/OpenBLAS-0.3.28/libopenblas zenp-r0.3.28.a
   /home/username/OpenBLAS-0.3.28/libopenblas zenp-r0.3.28.so
   username@computername:~/OpenBLAS-0.3.28$ gedit ~/.bashrc
   XCRYSDEN TOPDIR=/usr/bin
   XCRYSDEN LIB BINDIR=/usr/lib
   export XCRYSDEN TOPDIR XCRYSDEN LIB BINDER
   export LD LIBRARY PATH=$LD LIBRARY PATH:$HOME/OpenBLAS-0.3.28
   username@computername:~/OpenBLAS-0.3.28$ source ~/.bashrc
3. The following was entered into the terminal to install Open MPI:
   username@computername:~/OpenBLAS-0.3.28$ cd ~
   username@computername:~$ wget
   https://download.open-mpi.org/release/open-mpi/v5.0/openmpi-5.0.6.tar.bz2
   username@computername:~\$ tar xvf openmpi-5.0.6.tar.bz2
   username@computername:~\$ cd openmpi-5.0.6/
   username@computername:~/openmpi-5.0.6$ ./configure --prefix=$HOME/openmpi-5.0.6
   Open MPI configuration:
   -----
   Version: 5.0.6
   MPI Standard Version: 3.1
   Build MPI C bindings: yes
   Build MPI Fortran bindings: mpif.h, use mpi, use mpi f08
   Build MPI Java bindings (experimental): no
```

Build Open SHMEM support: false (no spml)

Debug build: no Platform file: (none)

Miscellaneous

Atomics: GCC built-in style atomics

Fault Tolerance support: mpi

HTML docs and man pages: installing packaged docs

hwloc: internal libevent: internal Open UCC: no pmix: internal PRRTE: internal

Threading Package: pthreads

Transports

Cisco usNIC: no

Cray uGNI (Gemini/Aries): no

Intel Omnipath (PSM2): no (not found)

Open UCX: no

OpenFabrics OFI Libfabric: no (not found)

Portals4: no (not found)

Shared memory/copy in+copy out: yes Shared memory/Linux CMA: yes Shared memory/Linux KNEM: no Shared memory/XPMEM: no

TCP: yes

Accelerators

CUDA support: no ROCm support: no

OMPIO File Systems

DDN Infinite Memory Engine: no

Generic Unix FS: yes

IBM Spectrum Scale/GPFS: no (not found)

Lustre: no (not found) PVFS2/OrangeFS: no

username@computername:~/openmpi-5.0.6\$ make all install

..

username@computername:~/openmpi-5.0.6\$ gedit ~/.bashrc

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```
export PATH=$PATH:$HOME/openmpi-5.0.6/bin
```

```
export
   LD LIBRARY PATH=$LD LIBRARY PATH:$HOME/OpenBLAS-0.3.28:$HOME/openmpi
   -5.0.6/lib
   username@computername:~/openmpi-5.0.6$ source ~/.bashrc
   username@computername:~/openmpi-5.0.6$ which mpicc
   /home/username/openmpi-5.0.6/bin/mpicc
4. The following was entered into the terminal to install fftw:
   username@computername:~/openmpi-5.0.6$ cd ~
   username@computername:~\$ wget https://www.fftw.org/fftw-3.3.10.tar.gz
   username@computername:~\$ tar xvf fftw-3.3.10.tar.gz
   username@computername:~$ cd fftw-3.3.10/
   username@computername:~/fftw-3.3.10$ ./configure FCC=gfortran CC=gcc MPICC=mpicc --
   enable-mpi --prefix=$HOME/fftw-3.3.10
   username@computername:~/fftw-3.3.10$ make
   username@computername:~/fftw-3.3.10$ make install
   username@computername:~/fftw-3.3.10\$ ls ~/fftw-3.3.10/include ~/fftw-3.3.10/lib
   /home/username/fftw-3.3.10/include:
   fftw3.f fftw3.h fftw31-mpi.f03 fftw3-mpi.h
   fftw3.f03 fftw31.f03 fftw3-mpi.f03 fftw3q.f03
   /home/username/fftw-3.3.10/lib:
   cmake libfftw3.a libfftw3.la libfftw3 mpi.a libfftw3 mpi.la pkgconfig
5. The following was entered into the terminal to install ScaLAPACK:
   username@computername:~/fftw-3.3.10$ cd ~
   username@computername:~$ wget
   https://github.com/Reference-ScaLAPACK/scalapack/archive/refs/tags/v2.2.0.tar.gz
   username@computername:~\$ tar xvf v2.2.0.tar.gz
   username@computername:~$ cd scalapack-2.2.0/
   username@computername:~/scalapack-2.2.0$ cp SLmake.inc.example SLmake.inc
   username@computername:~/scalapack-2.2.0$ gedit SLmake.inc
   username@computername:~/scalapack-2.2.0$ cat SLmake.inc
```

```
#
# Program:
              ScaLAPACK
#
# Module:
              SLmake.inc
#
# Purpose:
             Top-level Definitions
 Creation date: February 15, 2000
#
 Modified:
              October 13, 2011
#
 Send bug reports, comments or suggestions to scalapack@cs.utk.edu
# C preprocessor definitions: set CDEFS to one of the following:
   -DNoChange (fortran subprogram names are lower case without any suffix)
   -DUpCase (fortran subprogram names are upper case without any suffix)
#
#
   -DAdd
            (fortran subprogram names are lower case with " " appended)
CDEFS
          = -DAdd
 The fortran and C compilers, loaders, and their flags
#
        = mpifort -fallow-argument-mismatch
FC
CC
        = mpicc
NOOPT
           = -00
            = -03
FCFLAGS
            = -O3 -fPIC
CCFLAGS
FCLOADER
             = $(FC)
CCLOADER
             =$(CC)
FCLOADFLAGS = $(FCFLAGS)
CCLOADFLAGS = $(CCFLAGS)
# The archiver and the flag(s) to use when building archive (library)
# Also the ranlib routine. If your system has no ranlib, set RANLIB = echo
#
ARCH
          = ar
ARCHFLAGS
              = cr
RANLIB
           = ranlib
```

```
# The name of the ScaLAPACK library to be created
   SCALAPACKLIB = libscalapack.a
   # BLAS, LAPACK (and possibly other) libraries needed for linking test programs
   BLASLIB
               = -L/$(HOME)/OpenBLAS-0.3.28 -lopenblas
   LAPACKLIB
             = $(LAPACKLIB) $(BLASLIB)
   LIBS
   username@computername:~/scalapack-2.2.0$ make
   username@computername:~/scalapack-2.2.0$ gedit ~/.bashrc
   export
   LD LIBRARY PATH=$LD LIBRARY PATH:$HOME/OpenBLAS-0.3.28:$HOME/openmpi
   -5.0.6/lib:$HOME/scalapack-2.2.0/libscalapack.a
   username@computername:~/scalapack-2.2.0$ source ~/.bashrc
   username@computername:~/scalapack-2.2.0$ ls ~/scalapack-2.2.0/lib*
   /home/username/scalapack-2.2.0/libscalapack.a
6. The following was entered into the terminal to install ELPA:
   username@computername:~/scalapack-2.2.0$ cd ~
   username@computername:~$ wget
   https://gitlab.mpcdf.mpg.de/elpa/elpa/-/archive/release 2024 05 001/elpa-
   release 2024 05 001.tar.bz2
   username@computername:~$ tar xvf elpa-release 2024 05 001.tar.bz2
   username@computername:~$ cd elpa-release 2024 05 001/
   username@computername:~/elpa-release 2024 05 001$ ./autogen.sh
   username@computername:~/elpa-release 2024 05 001$ ./configure CFLAGS="-march=native"
   CPPFLAGS="-I$HOME/openmpi-5.0.6/include"
   SCALAPACK LDFLAGS="-L$HOME/scalapack-2.2.0/"
   LDFLAGS="-L$HOME/OpenBLAS-0.3.28 -L$HOME/openmpi-5.0.6/lib"
   --prefix=$HOME/elpa-release 2024 05 001 --disable-avx512
   The following ELPA2 kernels will be build:
    real generic
    real generic simple
```

```
real generic simple block4
    real generic simple block6
    real sse block2
    real sse block4
    real sse block6
    real sse assembly
    real avx block2
    real avx block4
    real avx block6
    real avx2 block2 (default)
    real avx2 block4
    real avx2 block6
    complex generic
    complex generic simple
    complex sse block1
    complex sse block2
    complex sse assembly
    complex avx block1
    complex avx block2
    complex avx2 block1 (default)
    complex avx2 block2
    This version of ELPA support the minimal API version: 20170403
   The current API version is: 20241105
   This version of ELPA support the minimal autotuning version: 20171201
   The current autotune version is: 20241105
   username@computername:~/elpa-release 2024 05 001$ make
   username@computername:~/elpa-release 2024 05 001$ make install
   username@computername:~/elpa-release 2024 05 001$ ls ~/elpa-release 2024 05 001/lib
   ~/elpa-release 2024 05 001/include/elpa-2024.05.001/elpa
   /home/username/elpa-release 2024 05 001/include/elpa-2024.05.001/elpa:
   elpa configured options.h elpa generated c api.h elpa.h
   elpa constants.h
                         elpa generated.h
                                             elpa simd constants.h
   elpa explicit name.h
                           elpa generic.h
                                              elpa version.h
   /home/username/elpa-release 2024 05 001/lib:
   libelpa.a libelpa.la libelpa.so libelpa.so.19 libelpa.so.19.3.0 pkgconfig
7. Go to:
   http://www.wien2k.at/reg_user/index.html
```

8. Click "Code download (after registration)"

- 9. Enter your username and password that you were given when you purchased WIEN2k.
- 10. Click the link "here" in "You can download the complete source code (about 170 MB) from here" to download "WIEN2k_24.1.tar". You should now have the source code package for WIEN2k 24.1.
- 11. If you have the folder "WIEN2k" in your Linux operating system home directory (for example at /home/username/WIEN2k, where username is replaced by your user name). You can rename it something else, such as "WIEN2k23.2".
- 12. The following was entered into the terminal to install WIEN2k 24.1:

```
username@computername:~/elpa-release 2024 05 001$ cd ~
username@computername:~$ mkdir WIEN2k
username@computername:~$ cd WIEN2k/
username@computername:~/WIEN2k$ ls
WIEN2k 24.1.tar
username@computername:~/WIEN2k$ tar xvf WIEN2k 24.1.tar
username@computername:~/WIEN2k$ gunzip *.gz
username@computername:~/WIEN2k$ chmod +x ./expand lapw
username@computername:~/WIEN2k$ export LC_NUMERIC=en_US.UTF-8
username@computername:~/WIEN2k$ ./expand lapw
continue (y/n)
y
Applied patches at: https://github.com/gsabo/WIEN2k-Patches/tree/master/24.1
username@computername:~/WIEN2k$ ./siteconfig
continue or stop (c/s) c
  Press RETURN to continue
 Selection: LG
  Press RETURN to continue
  Your compiler: gfortran
  Your compiler: gcc
  Press RETURN to continue
```

Hit Enter to continue Would you like to use LIBXC (needed ONLY for self-consistent gKS mGGA calculations, for the stress tensor and experts who want to play with different DFT options. It must have been installed before)? (y,N): y Do you want to automatically search for LIBXC installations? (Y,n): Please specify a comma separated list of directories to search! (If no list is entered, /usr/lib64, /usr/local and /opt will be searched as default): /home/username/libxc-7.0.0 Pick one of the following (enter the number of the line of your choice or 0 to manually specify a path)!: /home/username/libxc-7.0.0/src/.libs/libxc.a /home/username/libxc-7.0.0/lib/libxc.a 2 Press RETURN to continue Do you want to automatically search for FFTW installations? (Y,n): Please specify a comma separated list of directories to search! (If no list is entered, /usr/lib64, /usr/local and /opt will be searched as default): /home/username/fftw-3.3.10 Finding the required fftw3 library-files in /home/username/fftw-3.3.10 /home/username/fftw-3.3.10/.libs/libfftw3.a /home/username/fftw-3.3.10/lib/libfftw3.a Your present FFTW choice is: FFTW3 Present FFTW root directory is: Do you want to use a FFTW version from the list above? (Y,n): Please enter the line number of the chosen version or enter 0 to manually specify your choice!: 2 Please specify the target achitecture of your FFTW library (e.g. lib64) or accept present choice (enter): lib Please specify the name of your FFTW library or accept present choice (enter): fftw3

```
Is this correct? (Y,n): Y
Selection: R
  Real libraries=-L/home/username/OpenBLAS-0.3.28 -lopenblas -lpthread
Selection: O
  Compiler options=-ffree-form -O2 -ftree-vectorize -march=native -ffree-line-length-none
-ffpe-summary=none -fallow-argument-mismatch
Current settings:
 M OpenMP switch:
                           -fopenmp
 O Compiler options:
                          -ffree-form -O2 -ftree-vectorize -march=native -ffree-line-length-
none -ffpe-summary=none -fallow-argument-mismatch
 L Linker Flags:
                       $(FOPT) -L../SRC lib
 P Preprocessor flags
                         '-DParallel'
 R R LIBS (LAPACK+BLAS): -L/home/username/OpenBLAS-0.3.28 -lopenblas -lpthread
 F FFTW options:
                         -DFFTW3 -I/home/username/fftw-3.3.10/include
                        -L/home/username/fftw-3.3.10/lib -lfftw3
   FFTW-LIBS:
 X LIBX options:
                         -DLIBXC -I/home/username/libxc-7.0.0/include
   LIBXC-LIBS:
                         -L/home/username/libxc-7.0.0/lib -lxcf03 -lxc
Selection: S
  Press RETURN to continue
 Shared Memory Architecture? (y/N):y
 Do you know/need a command to bind your jobs to specific nodes?
 (like taskset -c). Enter N / your specific command: N
 Do you have MPI, ScaLAPACK, ELPA, or MPI-parallel FFTW installed and intend
 to run finegrained parallel?
 (y/N) y
 Your compiler: mpifort
Do you want to use a present ScaLAPACK installation? (Y,n): Y
Please specify the target architecture of your ScaLAPACK libraries (e.g. intel64)!:
Please specify the root path of your ScaLAPACK installation!:
ScaLAPACK root: /home/username/scalapack-2.2.0
Please specify the name of your ScaLAPACK library!:
ScaLAPACK library: scalapack
Please specify the root path of your BLACS installation!:
BLACS root:
```

```
Please specify the name of your BLACS library (eg. mkl_blacs_openmpi_lp64)!:
BLACS library:
Is this correct? (Y,n): Y
 Do you want to use ELPA? (y,N):
y
Do you want to automatically search for ELPA installations? (Y,n):
Please specify a comma-separated list of directories to search! (If no list is entered,
/usr/lib64 /usr/local and /opt will be searched as default):
/home/username/elpa-release 2024 05 001
Finding the required ELPA files in /home/username/elpa-release 2024 05 001 ....
More than one version of ELPA found:
/home/username/elpa-release 2024 05 001/.libs
/home/username/elpa-release 2024 05 001/lib
Pick one (enter line number) or enter 0 to manually specify a path!:
Present root directory of ELPA is: /home/username/elpa-release 2024 05 001/
Please specify the ROOT-path of your ELPA installation (like /usr/local/elpa/) or accept present
path (Enter):
Checking for ELPA version ...
More than one set of include files in your ELPA-ROOT directory. Pick one (enter line number)
or enter 0 to manually specify a version!:
/home/username/elpa-release 2024 05 001/test/Fortran/elpa generalized
/home/username/elpa-release 2024 05 001/test/Fortran/elpa2
/home/username/elpa-release 2024 05 001/src/elpa1
/home/username/elpa-release 2024 05 001/src/elpa generalized
/home/username/elpa-release 2024 05 001/src/elpa2
/home/username/elpa-release 2024 05 001/elpa
/home/username/elpa-release 2024 05 001/.fortran dependencies/elpa2 print kernels
/home/username/elpa-release 2024 05 001/include/elpa-2024.05.001
/home/username/elpa-release 2024 05 001/include/elpa-2024.05.001/elpa
/home/username/elpa-release 2024 05 001/share/doc/elpa
8
Is this correct? (Y,n):Y
Please specify the lib-directory of your ELPA installation (e.g. lib or lib64):
lib
Is this correct? (Y,n):Y
```

```
Please specify the name of your installed ELPA library (e.g. elpa or elpa openmp):
elpa
Your current lib-directory of your ELPA installation is: lib
Is this correct? (Y,n):Y
The current library name of your ELPA installation is: elpa
Is this correct? (Y,n):Y
These options derive from your chosen Settings:
 ELPAROOT:
                    /home/username/elpa-release 2024 05 001/
 ELPA VERSION:
                       2024.05.001
 ELPA LIB:
                   lib
 ELPA LIBNAME:
                        elpa
 Is this correct? (Y,n): Y
  Press RETURN to continue
Please enter your choice of additional libraries!:
$(R LIBS) -lmpi
Is this correct? (Y,n): Y
 Please specify your parallel compiler options or accept the recommendations (Enter - default)!:
 Please specify your parallel OMP_SWITCH (type "del" to blank it)
 or accept the recommendations (Enter - default)!:
 Please specify your MPIRUN command or accept the recommendations (Enter - default)!:
  Press RETURN to continue
 Current settings:
     Parallel compiler
                        : mpifort
     SCALAPACK LIBS
                              : -L/home/username/scalapack-2.2.0/ -lscalapack
     FFTW PLIBS
                         : -lfftw3 mpi
     ELPA OPT
                         :-DELPA
-I/home/username/elpa-release 2024 05 001/include/elpa-2024.05.001/elpa
           -I/home/username/elpa-release 2024 05 001/include/elpa-2024.05.001/modules
                         : -lelpa -L/home/username/elpa-release 2024 05 001/lib -Wl,-
     ELPA LIBS
rpath=/home/username/elpa-release 2024 05 001/lib
     FPOPT(par.comp.options): -ffree-form -O2 -ftree-vectorize -march=native -ffree-line-
length-none -ffpe-summary=none -fallow-argument-mismatch
     OMP SWITCH
                           : -fopenmp
     MPIRUN command
                             : mpirun -np NP -machinefile HOSTS EXEC
```

```
parallel execution:
        RP LIBS
                           : $(R LIBS) -lmpi
   Selection: S
      Press RETURN to continue
      Selection: Q
      Selection: A
   Compile time errors (if any) were:
                                            <= It should be blank here if successful.
   Check file compile.msg in the corresponding SRC * directory for the
   compilation log and more info on any compilation problem.
      Press RETURN to continue
    Please enter the full path of the perl program: /usr/bin/perl
      Press RETURN to continue
   Please enter the full path to your temporary directory: /tmp
      Press RETURN to continue
13. If it is the first time installing WIEN2k on the computer, enter in the terminal:
   username@computername:~/WIEN2k$./userconfig lapw
      Specify your prefered editor (default is emacs):
      editor shall be: gedit
      Set editor to gedit (Y/n) Y
      Specify your prefered DATA directory, where your cases should be
      stored (for /home/username/WIEN2k, just enter RETURN key):
      DATA directory: /home/username/wiendata
      Set DATA directory to /home/username/wiendata (Y/n) Y
      Specify your prefered scratch directory, where big case.vector files
```

can be stored (Recommended is a local directory (maybe /scratch), not a

NFS directory. For your working directory, just enter RETURN key): scratch directory:

Set scratch directory to working directory (Y/n) Y

Specify your program to read pdf files (default is okular) (on some Linux systems use xpdf, evince, pdfstudio, ...):evince

Set PDFREADER to evince (Y/n) Y

WIEN2k can use OpenMP parallelization on multicore computers. For details please read the "Parallelization section" of the Usersguide. Your present computer has 6 cores, but more than 4 (8) cores is useless. How many cores do you want to use by default (4):1

Set OMP NUM THREADS to 1 (Y/n) Y

ulimit -s unlimited

!!! The following lines will be added to your .bashrc file if you continue !!! A copy of your current .bashrc will be saved under .bashrc.savelapw!

```
# added by WIEN2k: BEGIN
# -----
alias lsi="ls -aslp *.in*"
alias lso="ls -aslp *.output*"
alias lsd="ls -aslp *.def"
alias lsc="ls -aslp *.clm*"
alias lss="ls -aslp *.scf* */*scf"
alias lse="ls -aslp *.error"
alias LS="ls -alsp |grep /"
alias pslapw="ps -ef |grep "lapw""
alias cdw="cd/home/username/wiendata"
if [ "$OMP NUM THREADS" = "" ]; then export OMP NUM THREADS=1; fi
#export LD LIBRARY PATH=.....
export EDITOR="gedit"
export SCRATCH=./
if [ "$WIENROOT" = "" ]; then export WIENROOT=/home/username/WIEN2k; fi
export W2WEB CASE BASEDIR=/home/username/wiendata
export STRUCTEDIT PATH=$WIENROOT/SRC structeditor/bin
export PDFREADER=evince
export PATH=$WIENROOT:$STRUCTEDIT PATH:$WIENROOT/SRC IRelast/script-
elastic:$PATH:.
export OCTAVE EXEC PATH=${PATH}::
export OCTAVE PATH=${STRUCTEDIT PATH}::
```

```
alias octave="octave -p $OCTAVE PATH"
  # -----
    Do you want to continue (Y/n)? Y
    Edit .rhosts file now? (y/N) N
  username@computername:~/WIEN2k$ source ~/.bashrc
14. To setup or start w2web, enter in the terminal:
  username@computername:~/WIEN2k$ w2web
  # w2web starter
  # Copyright (C) 2001 luitz.at
  w2web installer on host computername
  # w2web installer
  # Copyright (C) 2001 luitz.at
  Checking for Installation in /home/username/.w2web/computername
  Creating /home/username/.w2web
  Creating /home/username/.w2web/computername
  conf directory does not exist - creating it.
  logs directory does not exist - creating it.
  sessions directory does not exist - creating it.
  tmp directory does not exist - creating it.
  Installing w2web files ...
  Please answer these questions for proper installaltion.
  Just press enter for the default value of (in brackets).
  Please enter the username: [admin] username
  Please enter the password: [password] password
  username:password
  Remember these. You will need them when you log in.
  Select the port to run on: [7890]
  Running on port 7890
```

Using localhost

Please enter this system's hostname: [computername] localhost

Is this your master node?: [y] y Installing... Attempting to start now...

Trying to start /home/username/WIEN2k/SRC_w2web/bin/w2web w2web server started, now point your web browser to http://localhost:7890

done.

- 15. The WIEN2k 24.1 installation is now complete. To test if WIEN2k installed okay or not, run the TiC example at [1]. Additional details of the TiC example are given in the WIEN2k usersguide (section "3 Quick Start" starting on page 13) [2].
- [1] https://github.com/gsabo/WIEN2k-Docs/blob/main/WIEN2k24.1%20TiC%20Example.pdf
- [2] http://susi.theochem.tuwien.ac.at/reg_user/textbooks/usersguide.pdf