# User Guide for ffr-LFDFT

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## 1 Introduction

Welcome to ffr-LFDFT documentation. In this document you will find the following information:

- basic information about ffr-LFDFT
- $\bullet\,$  how to compile and use the program
- implementation details of the program

ffr-LFDFT is a poor man's program (or collection of subroutines, as of now) to carry out electronic structure calculations based on density functional theory and Lagrange basis set.

This program is intended for research in implementation of new methods in electronic structure calculations in condensed matter. Currently it is not as stable or have lot of functionalities as more well-known package such as Quantum Espresso, ABINIT, or VASP. However, it can be used to calculate total energy of solids.

This program is written mainly by me, Fadjar Fathurrahman at Research Center of Nanoscience and Nanotechnology, Bandung Institute of Technology, Indonesia.

Some references: [1], [2], [3], [4]

### 2 Installation

There is no need for installation, actually. What is meant by installation here is compiling the library libmain.a and linking the main executable of ffr-LFDFT, namely ffr\_LFDFT.x

A manually written Makefile is provided. On the topmost part of the Makefile you need to specify which make.inc file you want to use. This make.inc file contains definition of compiler executable, compiler flags, and libraries used in the linking process. Several make.inc files that I used can be found in the directory platform. You need to decide which compiler to use if there are more than one compiler in your system. For a typical Linux system, make.inc.gfortran is sufficient. You can manually edit the compiler options in the corresponding make.inc files.

Currently, ffr-LFDFT is tested using the following compilers on Linux system:

- GNU Fortran compiler, executable: gfortran
- G95 Fortran compiler, executable: g95
- Intel Fortran compiler, executable: ifort
- PGI Fortran compiler, executable: pgf90 or pgf95
- Sun (now part of Oracle) Fortran compiler: sunf95

There following external libraries are required to build ffr-LFDFT

- BLAS
- LAPACK
- FFTW3

Typing the command

make

will build the library libmain.a and typing the command

make main

will build the main executable ffr\_LFDFT.x.

### 3 Usage

ffr\_LFDFT.x accepts input file in plain text format. The structure of the input files are very similar to PWSCF input file with minor differences.

As an example, the following input file is for LiH molecule:

```
&CONTROL
  pseudo_dir = '../../HGH'
  etot_conv_thr = 1.0d-6
&SYSTEM
  ibrav = 8
  nat = 2
  ntyp = 2
  A = 8.4668d0
  B = 8.4668d0
  C = 8.4668d0
  nr1 = 45
  nr2 = 45
  nr3 = 45
&ELECTRONS
  KS_Solve = 'Emin_pcg'
  cg_beta = 'DY'
  electron_maxstep = 150
  mixing_beta = 0.1
  diagonalization = 'LOBPCG'
ATOMIC_SPECIES
     3.0 Li_sc.hgh
     1.0 H.hgh
ATOMIC_POSITIONS angstrom
     0.0 0.0 0.0
Η
     1.0 0.0 0.0
```

Other examples can be found under directory works.

The following input variables are special to ffr-LFDFT and not found in PWSCF. These variables are defined in the namelist ELECTRONS.

- KS\_Solve: method to solve Kohn-Sham equation, accepted values:
  - 'SCF': using diagonalization-based self-consistent iterations
  - 'Emin-pcg': using direct minimization based on nonlinear conjugate gradient algorithm
- cg\_beta: method to calculate parameter  $\beta$  in nonlinear CG minimization used in direct Kohn-Sham energy minimization.

```
'FR': using Fletcher-Reeves formula
'PR': using Polak-Ribiere formula
'HS': using Hestenes-Stiefel formula
'DY': using Dai-Yuan formula
```

### 4 Kohn-Sham equation

Within LDA, Kohn-Sham energy functional can be written as:

$$E_{\rm LDA}\left[\{\psi_i(\mathbf{r})\}\right] = E_{\rm kin} + E_{\rm ion} + E_{\rm Ha} + E_{\rm xc} \tag{1}$$

with the following energy terms.

(1) kinetic energy:

$$E_{\rm kin} = -\frac{1}{2} \sum_{i} \int \psi_i^*(\mathbf{r}) \, \nabla^2 \, \psi_i(\mathbf{r}) \, d\mathbf{r}$$
 (2)

(2) ion-electron interaction energy:

$$E_{\rm ion} = \int V_{\rm ion}(\mathbf{r}) \, \rho(\mathbf{r}) \, \mathrm{d}\mathbf{r} \tag{3}$$

(3) Hartree (electrostatic) energy:

$$E_{\text{Ha}} = \int \frac{1}{2} \frac{\rho(\mathbf{r})\rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r} d\mathbf{r}'$$
(4)

(4) Exchange-correlation energy (using LDA):

$$E_{\rm xc} = \int \epsilon_{\rm xc} \left[ \rho(\mathbf{r}) \right] \rho(\mathbf{r}) \, \mathrm{d}\mathbf{r} \tag{5}$$

Central to the density functional theory is the so-called Kohn-Sham equation. This equation can be written as:

$$\left[ -\frac{1}{2} \nabla^2 + V_{KS}(\mathbf{r}) \right] \psi_i(\mathbf{r}) = \epsilon_i \psi_i(\mathbf{r}) \tag{6}$$

where  $\epsilon i$  and  $\psi_i(\mathbf{r})$  is known as Kohn-Sham eigenvalues and eigenvectors (orbitals). Quantity  $V_{\rm KS}$  is called the Kohn-Sham potential, which can be written as sum of several potentials:

$$V_{\rm KS}(\mathbf{r}) = V_{\rm ion}(\mathbf{r}) + V_{\rm Ha}(\mathbf{r}) + V_{\rm xc}(\mathbf{r}) \tag{7}$$

 $V_{\text{ion}}$  denotes attractive potential between ion (or atomic nuclei) with electrons. This potential can be written as:

$$V_{\text{ion}}(\mathbf{r}) = \sum_{I}^{N_{\text{atoms}}} \frac{Z_I}{|\mathbf{r} - \mathbf{R}_I|}$$
(8)

This potential is Coulombic and has singularities at the ionic centers. It is generally difficult to describe this potential fully. It is common to replace the full Coulombic potential with softer potential which is known as pseudopotential. There are various types or flavors of pseudopotentials. In the current implementation, ion-electron potential,  $V_{\rm ion}$  is treated by pseudopotential. HGH-type pseudopotential is employed due to the the availability of analytic forms both in real and reciprocal space.

 $V_{\rm Ha}$  is the classical Hartree potential. It is defined as

$$V_{\rm Ha}(\mathbf{r}) = \int \frac{\rho(\mathbf{r}')}{\mathbf{r} - \mathbf{r}'} \, \mathrm{d}\mathbf{r}', \tag{9}$$

where  $\rho(\mathbf{r})$  denotes electronic density:

$$\rho(\mathbf{r}) = \sum_{i}^{N_{\text{occ}}} \psi_i^*(\mathbf{r}) \psi_i(\mathbf{r})$$
(10)

Alternatively, Hartree potential can also be obtained via solving Poisson equation:

$$\nabla^2 V_{\text{Ha}}(\mathbf{r}) = -4\pi \rho(\mathbf{r}) \tag{11}$$

The last term in Equation (7) is exchange-correlation potential.

### 5 Implementation

ffr-LFDFT is implemented in simple Fortran language. I used global variables heavily, as opposed to using user-defined type to contained them. Currently, only one user-defined type is used in ffr-LFDFT, namely Ps\_HGH\_Params\_T which is mainly used for convenience. I tried to make the code clear for those who are beginners in implementing a density-functional calculations (such as myself).

#### 5.1 Main program

Currently, the calculation flow of the main program of ffr-LFDFT is as follows:

- Getting program argument as input file and reading the input file
- Initializing molecular structure, pseudopotentials, and Lagrange basis functions, including grids
- Setting additional options if necessary based on the input file
- Initializing electronic states variables
- Setting up Hamiltonian: potential and kinetic operators.
- Solving the Kohn-Sham equation via direct minimization or self-consistent field

The appropriate subroutine calls is given below.

```
CALL getarg( 1, filein )
CALL read_input( filein )
CALL setup_from_input()
CALL setup_options()
CALL init_betaNL()
CALL init_states()
CALL init_strfact_shifted()
CALL calc_Ewald_qe()
CALL alloc_hamiltonian()
CALL init_V_ps_loc_G()
CALL init_nabla2_sparse()
CALL init_ilu0_prec()
CALL gen_guess_rho_gaussian()
CALL gen_random_evecs()
CALL gen_gaussian_evecs()
IF( I_KS_SOLVE == 1 ) THEN
  CALL KS_solve_Emin_pcg()
  CALL calc_evals( Nstates, Focc, evecs, evals )
ELSEIF( I_KS_SOLVE == 2 ) THEN
  CALL KS_solve_SCF_v2()
ENDIF
```

Subroutine setup\_from\_input() is a wrapper to three setup calls:

```
CALL setup_atoms()
CALL setup_PsPot()
CALL setup_LF3d()
```

The subroutine names is self-explanatory.

Subroutine  $\mathtt{setup\_options}()$  converts various optional input variables into global variables mainly defined in  $\mathtt{m\_options}$ .

Before going into futher details of the calculation, I will describe first the data structures used for describing grids, basis functions, atomic structure and pseudopotentials.

#### 5.2 Description of LF basis set

Description of LF basis set in 3d is given in module m\_LF3d. All global variables in this module is given prefix LF3d.

```
MODULE m_LF3d
  IMPLICIT NONE
  INTEGER, PARAMETER :: LF3d_PERIODIC = 1
  INTEGER, PARAMETER :: LF3d_CLUSTER = 2
  INTEGER, PARAMETER :: LF3d_SINC
  INTEGER :: LF3d_TYPE
  INTEGER, DIMENSION(3) :: LF3d_NN
  REAL(8), DIMENSION(3) :: LF3d_LL, LF3d_AA, LF3d_BB, LF3d_hh
  INTEGER :: LF3d_Npoints
  REAL(8) :: LF3d_dVol
  REAL(8), ALLOCATABLE :: LF3d_grid_x(:), LF3d_grid_y(:), LF3d_grid_z(:)
  REAL(8), ALLOCATABLE :: LF3d_D1jl_x(:,:), LF3d_D1jl_y(:,:), LF3d_D1jl_z(:,:)
  REAL(8), ALLOCATABLE :: LF3d_D2jl_x(:,:), LF3d_D2jl_y(:,:), LF3d_D2jl_z(:,:)
  REAL(8), ALLOCATABLE :: LF3d_lingrid(:,:)
  INTEGER, ALLOCATABLE :: LF3d_xyz2lin(:,:,:)
  INTEGER, ALLOCATABLE :: LF3d_lin2xyz(:,:)
  REAL(8), ALLOCATABLE :: LF3d_G2(:), LF3d_Gv(:,:)
END MODULE
```

Variables in m\_LF3d is initialized by calling the subroutine init\_LF3d\_XX(), where XX may be one of:

- p: periodic LF
- c: cluster LF
- sinc: sinc L

```
SUBROUTINE init_LF3d_p( NN, AA, BB )
SUBROUTINE init_LF3d_c( NN, AA, BB )
SUBROUTINE init_LF3d_sinc( NN, hh )
```

In the above subroutines:

- NN: an array of 3 integers, specifying sampling points in x, y and z direction.
- AA: an array of 3 floats, specifying left ends of unit cell.
- BB: an array of 3 floats, specifying right ends of unit cell.
- hh: an array of 3 floats, specifying spacing between adjacent sampling points.

Note that for periodic and cluster LF we have to specify NN, AA, and BB while for sinc LF we have to specify NN and hh. Note that for periodic LF NN must be odd numbers.

#### Example:

```
NN = (/ 35, 35, 35 /)
AA = (/ 0.d0, 0.d0, 0.d0 /)
BB = (/ 6.d0, 6.d0, 6.d0 /)
CALL init_LF3d_p( NN, AA, BB )
```

#### 5.3 Description of molecular or crystalline structure

Description of molecular or crystalline structure is given in module m\_atoms. Note that unit cell for crystalline structure (currently only orthorombic structure is possible) is specified by AA and BB in call to init\_LF3d\_p()

```
MODULE m_atoms
IMPLICIT NONE
INTEGER :: Natoms
INTEGER :: Nspecies
REAL(8), ALLOCATABLE :: AtomicCoords(:,:)
INTEGER, ALLOCATABLE :: atm2species(:)
CHARACTER(5), ALLOCATABLE :: SpeciesSymbols(:)
REAL(8), ALLOCATABLE :: AtomicValences(:)
COMPLEX(8), ALLOCATABLE :: StructureFactor(:,:)
END MODULE
```

The global variables in module m\_atoms can initialized from an XYZ file by calling the subroutine init\_atoms\_xyz().

```
SUBROUTINE init_atoms_xyz( fil_xyz )
```

This subroutine takes one argument fil\_xyz which is the path to XYZ file describing the molecular structure or crystalline structure.

In the main program, the variables are initialized by calling the subroutine setup\_atoms(). This subroutine handles conversion from input data read to internal global variables in module m\_atoms.

### 5.4 Pseudopotential

Module m\_PsPot

```
MODULE m_PsPot

USE m_Ps_HGH, ONLY : Ps_HGH_Params_T

IMPLICIT NONE

CHARACTER(128) :: PsPot_Dir = './HGH/'

CHARACTER(128), ALLOCATABLE :: PsPot_FilePath(:)

TYPE(Ps_HGH_Params_T), ALLOCATABLE :: Ps_HGH_Params(:)

INTEGER :: NbetaNL

REAL(8), ALLOCATABLE :: betaNL(:,:)

INTEGER, ALLOCATABLE :: prj2beta(:,:,:,:)

INTEGER :: NprojTotMax

END MODULE
```

We currently support HGH pseudopotential only. The HGH pseudopotential parameter is described by an array of type Ps\_HGH\_Params\_T which is defined in m\_Ps\_HGH:

```
TYPE Ps_HGH_Params_T

CHARACTER(5) :: atom_name

INTEGER :: zval

REAL(8) :: rlocal

REAL(8) :: rc(0:3)

REAL(8) :: c(1:4)

REAL(8) :: h(0:3, 1:3, 1:3)

REAL(8) :: k(0:3, 1:3, 1:3)

INTEGER :: lmax

INTEGER :: Nproj_l(0:3) ! number of projectors for each AM

REAL(8) :: rcut_NL(0:3)

END TYPE
```

The array betall and pri2beta are related to nonlocal pseudopotential calculation.

Except for the array betaNL, most variables in module m\_PsPot are initialized by the call to subroutine init\_PsPot.

```
ALLOCATE( PsPot_FilePath(Nspecies) )
ALLOCATE( Ps_HGH_Params(Nspecies) )
D0 isp = 1,Nspecies
  PsPot_FilePath(isp) = trim(PsPot_Dir) // trim(SpeciesSymbols(isp)) // '.hgh'
  CALL init_Ps_HGH_Params( Ps_HGH_Params(isp), PsPot_FilePath(isp) )
  AtomicValences(isp) = Ps_HGH_Params(isp)%zval
ENDDO
```

Initialization of array prj2beta

```
ALLOCATE( prj2beta(1:3,1:Natoms,0:3,-3:3) )
prj2beta(:,:,:,:) = -1
NbetaNL = 0
D0 ia = 1,Natoms
  isp = atm2species(ia)
  D0 1 = 0,Ps_HGH_Params(isp)%lmax
    D0 iprj = 1,Ps_HGH_Params(isp)%Nproj_l(1)
    D0 m = -1,1
        NbetaNL = NbetaNL + 1
        prj2beta(iprj,ia,1,m) = NbetaNL
        ENDDO ! m
    ENDDO ! iprj
ENDDO ! ia
```

#### 5.5 Kinetic operator and energy

Using LF basis:

$$T_{\alpha,\beta,\gamma} = -\frac{1}{2} \sum_{i} f_{i} \left\langle \psi_{i} \middle| \nabla^{2} \middle| \psi_{i} \right\rangle = -\frac{1}{2} \sum_{i} f_{i} \sum_{\alpha\alpha'} \sum_{\beta\beta'} \sum_{\gamma\gamma'} C_{i,\alpha\beta\gamma} \mathbb{L}_{\alpha\beta\gamma}^{\alpha'\beta'\gamma'} C_{i,\alpha'\beta'\gamma'}$$
(12)

The Laplacian matrix has the following form:

$$\mathbb{L}_{\alpha\beta\gamma}^{\alpha'\beta'\gamma'} = D_{\alpha\alpha'}^{(2)}\delta_{\beta\beta'}\delta_{\gamma\gamma'} + D_{\beta\beta'}^{(2)}\delta_{\alpha\alpha'}\delta_{\gamma\gamma'} + D_{\gamma\gamma'}^{(2)}\delta_{\alpha\alpha'}\delta_{\beta\beta'}$$
(13)

For periodic LF:

$$D_{ij}^{(2)} = -\left(\frac{2\pi}{L}\right)^2 \frac{N'}{3} \left(N'+1\right) \delta_{ij} + \frac{\left(\frac{2\pi}{L}\right)^2 (-1)^{i-j} \cos\left[\frac{\pi(i-j)}{N}\right]}{2\sin^2\left[\frac{\pi(i-j)}{N}\right]} (1 - \delta_{nn'})$$
(14)

where N' = (N - 1)/2.

An implementation of the equation (14) can be found in the file init\_deriv\_matrix\_p.

```
! Diagonal elements
NPRIMED = (N-1)/2
D0 jj = 1, N
   D1j1(jj,jj) = 0d0
   D2j1(jj,jj) = -( 2.d0*PI/L )**2.d0 * NPRIMED * (NPRIMED+1)/3.d0
ENDD0
! Off diagonal elements
D0 jj = 1, N
```

```
DO 11 = jj+1, N

nn = jj - 11

tt1 = PI/L * (-1.d0)**nn

tt2 = sin(PI*nn/N)

tt3 = (2.d0*PI/L)**2d0 * (-1.d0)**nn * cos(PI*nn/N)

tt4 = 2.d0*sin(PI*nn/N)**2d0

D1j1(jj,11) = tt1/tt2

D1j1(11,jj) = -tt1/tt2

D2j1(jj,11) = -tt3/tt4

D2j1(11,jj) = -tt3/tt4

ENDDO

ENDDO
```

Code for calculating kinetic term contribution to total energy:

There two ways to implement kinetic term contribution to wave function gradient:

- By building the matrix representation of kinetic operator in the sparse form
- Using matrix-free method

The following code build the matrix representation of kinetic operator:

```
! Number of nonzeros per column
nnzc = Nx + Ny + Nz - 2
! Total number of nonzeros
NNZ = nnzc*Npoints
! Allocate arrays for CSC format
ALLOCATE( rowval(NNZ) )
ALLOCATE( nzval(NNZ) )
ALLOCATE( colptr(Npoints+1) )
! Initialize rowGbl pattern for x, y, and z components
ALLOCATE( rowGbl_x_orig(Nx) )
ALLOCATE( rowGbl_y_orig(Ny) )
ALLOCATE( rowGbl_z_orig(Nz) )
rowGbl_x_orig(1) = 1
DO ix = 2, Nx
 rowGbl_x_orig(ix) = rowGbl_x_orig(ix-1) + Ny*Nz
ENDDO
rowGbl_y_orig(1) = 1
DO iy = 2,Ny
 rowGbl_y_orig(iy) = rowGbl_y_orig(iy-1) + Nz
ENDDO
DO iz = 1,Nz
 rowGbl_z_orig(iz) = iz
ENDDO
ip = 0
DO colGbl = 1, Npoints
  ! Determine local column index for x, y, and z components
  colLoc_x = ceiling( real(colGbl)/(Ny*Nz) )
  yy = colGbl - (colLoc_x - 1)*Ny*Nz
  colLoc_y = ceiling( real(yy)/Nz )
```

```
izz = ceiling( real(colGbl)/Nz )
  colLoc_z = colGbl - (izz-1)*Nz
  ! Add diagonal element (only one element in any column)
  ip = ip + 1
  rowval(ip) = colGbl
 nzval(ip) = D2jl_x(colLoc_x,colLoc_x) + D2jl_y(colLoc_y,colLoc_y) + &
              D2j1_z(colLoc_z,colLoc_z)
  ! Add non-diagonal elements
  DO ix = 1, Nx
    IF ( ix /= colLoc_x ) THEN
      ip = ip + 1
      rowval(ip) = rowGbl_x_orig(ix) + colGbl - 1 - (colLoc_x - 1)*Ny*Nz
      nzval(ip) = D2jl_x(ix,colLoc_x)
    ENDIF
  ENDDO
  DO iy = 1,Ny
    IF ( iy \neq colLoc_y ) THEN
      ip = ip + 1
      rowval(ip) = rowGbl_y_orig(iy) + colGbl - 1 - (izz-1)*Nz + (colLoc_x - 1)*Ny*Nz
      nzval(ip) = D2jl_y(iy,colLoc_y)
   ENDIF
  ENDDO
  D0 iz = 1,Nz
    IF ( iz /= colLoc_z ) THEN
      ip = ip + 1
      rowval(ip) = rowGbl_z_orig(iz) + (izz-1)*Nz
      nzval(ip) = D2jl_z(iz,colLoc_z)
    ENDIF
  ENDDO
ENDDO
! Now colptr
colptr(1) = 1
DO ii = 2, Npoints+1
  colptr(ii) = colptr(ii-1) + nnzc
ENDDO
! Sort using subroutine csort from SPARSKIT
nwork = max( Npoints+1, 2*(colptr(Npoints+1)-colptr(1)) )
ALLOCATE( iwork( nwork ) )
CALL csort( Npoints, nzval, rowval, colptr, iwork, .TRUE. )
Multiplication of kinetic matrix with wave function can be done using SPARSKIT's sparse matrix-
vector multiplication subroutine amux:
CALL amux( Npoints, v(:,ic), Hv(:,ic), nzval, rowval, colptr )
Alternatively, one can use matrix-free method (without building 3D Laplacian matrix)
DO ip = 1, Npoints
 i = lin2xyz(1,ip)
  j = lin2xyz(2,ip)
 k = lin2xyz(3,ip)
 Lv(ip) = 0.d0
  DO ii = 1, Nx
   Lv(ip) = Lv(ip) + D2jl_x(ii,i) * v(xyz2lin(ii,j,k))
  ENDDO
  DO jj = 1, Ny
   Lv(ip) = Lv(ip) + D2jl_y(jj,j) * v(xyz2lin(i,jj,k))
  ENDDO
  DO kk = 1, Nz
    Lv(ip) = Lv(ip) + D2jl_z(kk,k) * v(xyz2lin(i,j,kk))
```

```
ENDDO
ENDDO
```

ILU0 preconditioner based on kinetic matrix:

```
ALLOCATE( alu_ilu0(Npoints*(Nx+Ny+Nz-2)) )
ALLOCATE( jlu_ilu0(Npoints*(Nx+Ny+Nz-2)) )
ALLOCATE( ju_ilu0(Npoints) )
ALLOCATE( iw_ilu0(Npoints) )
CALL ilu0( Npoints, -0.5d0*nzval, rowval, colptr, alu_ilu0, jlu_ilu0, & ju_ilu0, iw_ilu0, ierr )
Apply preconditioner:
CALL lusol( Npoints, v, Kv, alu_ilu0, jlu_ilu0, ju_ilu0)
```

#### 5.6 Local pseudopotential

Local pseudopotential term is very simple because it is diagonal in real space. This term is represented by the global array V\_ps\_loc found in file m\_hamiltonian. Despite its name, it can however be used to represent any local external potential such as harmonic potential.

It contribution to total energy is simply sum over grid points:

```
E_ps_loc = sum( Rhoe(:) * V_ps_loc(:) )*dVol
```

Its contribution to wave function gradient is simply obtained by point-wise multiplication with wave function expansion coefficients.

#### 5.7 Hartree term: solution of Poisson equation

Hartree potential can be calculated by solving Poisson equation once the charge density has been calculated. There are several methods to solve Poisson equation. For periodic system, the most popular method is via FFT. In this method charge density is first transformed to reciprocal space or  $\mathbf{G}$ -space. In this space, Hartree potential can be calculated by simply dividing charge density with magnitude of non-zero reciprocal vectors  $\mathbf{G}$ .

To implement this method we need to generate reciprocal Gectors **G**. In the current implementation **G**-vectors are declared in module m\_LF3d, namely LF3d\_Gv and LF3d\_Gv2 for **G**-vectors and their magnitudes, respectively. The subroutine which is responsible to generate **G**-vectors is init\_gvec().

```
ALLOCATE( G2(Npoints) )
ALLOCATE( Gv(3, Npoints) )
ig = 0
DO k = 0, NN(3)-1
DO j = 0, NN(2)-1
D0 i = 0, NN(1)-1
  ig = ig + 1
  ii = mm_to_nn( i, NN(1) )
  jj = mm_{to_nn}(j, NN(2))
  kk = mm_to_nn(k, NN(3))
  Gv(1,ig) = ii * 2.d0*PI/LL(1)
  Gv(2,ig) = jj * 2.d0*PI/LL(2)
  Gv(3,ig) = kk * 2.d0*PI/LL(3)
  G2(ig) = Gv(1,ig)**2 + Gv(2,ig)**2 + Gv(3,ig)**2
ENDDO
ENDDO
ENDDO
```

The function mm\_to\_nn describes mapping between real space grid and Fourier grid.

```
FUNCTION mm_to_nn( mm, S ) RESULT(idx)
   IMPLICIT NONE
   INTEGER :: idx
   INTEGER :: mm
   INTEGER :: S
   IF(mm > S/2) THEN
      idx = mm - S
   ELSE
      idx = mm
   ENDIF
END FUNCTION
```

Driver for solving Poisson equation via FFT is implemented in subroutine Poisson\_solve\_fft()

```
ALLOCATE( tmp_fft(Npoints) )
D0 ip = 1, Npoints
   tmp_fft(ip) = cmplx( rho(ip), 0.d0, kind=8 )
ENDDO
! forward FFT
CALL fft_fftw3( tmp_fft, Nx, Ny, Nz, .false. ) ! now 'tmp_fft = rho(G)'
tmp_fft(1) = (0.d0,0.d0) ! zero-G component
D0 ip = 2, Npoints
   tmp_fft(ip) = 4.d0*PI*tmp_fft(ip) / G2(ip)
ENDDO ! now 'tmp_fft' = phi(G)
! Inverse FFT
CALL fft_fftw3( tmp_fft, Nx, Ny, Nz, .true. )
! Transform back to real space
D0 ip = 1, Npoints
   phi(ip) = real( tmp_fft(ip), kind=8 )
ENDDO
```

Hartree energy is calculated according to the following equation:

$$E_{\text{Ha}} = \frac{1}{2} \int \rho(\mathbf{r}) V_{\text{Ha}}(\mathbf{r}) \, d\mathbf{r}$$
 (15)

This is done by simple summation over grid points:

```
E_Hartree = 0.5d0*sum( Rhoe(:) * V_Hartree(:) )*dVol
```

### 5.8 XC energy and potential

Currently the only supported form of XC functional is of Volko-Wilk-Nusair which is implemented in the file LDA\_VWN.f90. In the future version, LibXC implementation will be implemented hopefully. XC contribution to total energy is implemented as follows.

```
CALL excVWN( Npoints, Rhoe, epsxc )
E_xc = sum( Rhoe(:) * epsxc(:) )*dVol
```

**TODO**: Equation for  $V_{XC}$  XC contribution to wavefunction gradient is treated the same way as local potential.

```
CALL excVWN( Npoints, Rhoe, epsxc )
CALL excpVWN( Npoints, Rhoe, depsxc )
V_xc(:) = epsxc(:) + Rhoe(:)*depsxc(:)
```

#### 5.9 Nonlocal pseudopotential

Nonlocal pseudopotential contribution to total energy using HGH pseudopotential can be written as.

$$E_{\mathrm{ps,NL}} = \Omega \sum_{i_{st}}^{N_{\mathrm{occ}}} f_{i_{st}} \sum_{I_{a}}^{N_{\mathrm{atom}}} \sum_{l=0}^{l_{\mathrm{max}}} \sum_{m=-l}^{l} \sum_{i_{p},j_{p}} h_{i_{p},j_{p}} \left[ \sum_{\alpha\beta\gamma}^{N_{\mathrm{points}}} C_{\alpha\beta\gamma}^{i_{st}} B_{i_{st}I_{a}lm}^{\mathrm{NL}} \left( \mathbf{r}_{\alpha\beta\gamma} - \mathbf{R}_{I_{a}} \right) \right]$$

$$\left[ \sum_{\alpha'\beta'\gamma'}^{N_{\mathrm{points}}} C_{\alpha'\beta'\gamma'}^{i_{st}} B_{i_{st}I_{a}lm}^{\mathrm{NL}} \left( \mathbf{r}_{\alpha'\beta'\gamma'} - \mathbf{R}_{I_{a}} \right) \right]$$

$$(16)$$

This equation can be implemented as follows.

```
E_ps_NL = 0.d0
DO ist = 1, Nstates_occ
  enl1 = 0.d0
  DO ia = 1, Natoms
    isp = atm2species(ia)
    DO 1 = 0, Ps(isp)%lmax
    D0 m = -1,1
      DO iprj = 1,Ps(isp)%Nproj_1(1)
      DO jprj = 1,Ps(isp)%Nproj_1(1)
        ibeta = prj2beta(iprj,ia,1,m)
        jbeta = prj2beta(jprj,ia,1,m)
        hij = Ps(isp)%h(1,iprj,jprj)
        enl1 = enl1 + hij*betaNL_psi(ia,ist,ibeta)*betaNL_psi(ia,ist,jbeta)
      ENDDO ! jprj
      ENDDO ! iprj
    ENDDO ! m
    ENDDO ! l
 E_ps_NL = E_ps_NL + Focc(ist)*enl1
ENDDO
```

Nonlocal HGH pseudopotential action can be written as follows.

$$\hat{V}_{\text{ps,NL}}\psi_{i_{st}}(\mathbf{r}_{\alpha\beta\gamma}) = \sum_{i_{st}}^{N_{\text{occ}}} \sum_{I_{a}}^{N_{\text{atom}}} \sum_{l=0}^{+l} \sum_{m=-l}^{+l} \sum_{i_{p},j_{p}} h_{i_{p},j_{p}} \mathbf{B}_{i_{st}I_{a}lm}^{\text{NL}} \left(\mathbf{r}_{\alpha\beta\gamma} - \mathbf{R}_{I_{a}}\right) \left[ \sum_{\alpha'\beta'\gamma'}^{N_{\text{points}}} C_{\alpha'\beta'\gamma'}^{i} \mathbf{B}_{i_{st}I_{a}lm}^{\text{NL}} \left(\mathbf{r}_{\alpha'\beta'\gamma'} - \mathbf{R}_{I_{a}}\right) \right]$$
(17)

This equation is implemented as follows.

```
ENDDO ! iprj
    ENDDO ! m
    ENDDO ! l
  ENDDO
ENDDO
The array betaNL is defined initialized in subroutine init_betaNL:
ALLOCATE( betaNL(Npoints,NbetaNL) )
! loop structure must be the same as in init_PsPot
ibeta = 0
DO ia = 1, Natoms
  isp = atm2species(ia)
  DO 1 = 0, Ps(isp)%lmax
    DO iprj = 1,Ps(isp)%Nproj_l(1)
      D0 m = -1,1
        ibeta = ibeta + 1
        Np_beta = 0
        DO ip = 1, Npoints
          CALL calc_dr_periodic_1pnt( LL, atpos(:,ia), lingrid(:,ip), dr_vec )
          dr = sqrt(dr_vec(1)**2 + dr_vec(2)**2 + dr_vec(3)**2)
          IF( dr <= Ps(isp)%rcut_NL(1) ) THEN</pre>
            Np_beta = Np_beta + 1
            betaNL(ip,ibeta) = hgh_eval_proj_R( Ps(isp), 1, iprj, dr ) * &
                                Ylm_real( 1, m, dr_vec )
          ENDIF
        ENDDO
        ! Test normalization of projectors
        nrm = sum(betaNL(:,ibeta)**2)*dVol
        WRITE(*,'(1x,A,I5,I8,F18.10)') 'ibeta, Np_beta, integ = ', ibeta, Np_beta, nrm
      ENDDO ! iprj
    ENDDO ! m
  ENDDO ! l
ENDDO
and betaNL_psi is calculated in calc_betaNL_psi:
! immediate return if no projectors are available
IF( NbetaNL <= 0 ) THEN</pre>
 RETURN
ENDIF
betaNL_psi(:,:,:) = 0.d0
DO ia = 1, Natoms
 DO ist = 1, Nstates
    DO ibeta = 1,NbetaNL
     betaNL_psi(ia,ist,ibeta) = ddot( Npoints, betaNL(:,ibeta),1, psi(:,ist),1 ) * dVol
    ENDDO
 ENDDO
ENDDO
       Ion-Ion interaction
Via Ewald summation: Using the code included in PWSCF
INTEGER, PARAMETER :: mxr = 50
! setup at and bg
at(:,:) = 0.d0
```

ENDDO ! jprj

at(1,1) = LL(1)at(2,2) = LL(2)

```
at(3,3) = LL(3)
omega = LL(1)*LL(2)*LL(3)
bg(:,:) = 0.d0
bg(1,1) = TPI/LL(1)
bg(2,2) = TPI/LL(2)
bg(3,3) = TPI/LL(3)
gcutm = maxval( NN )*TPI !! ???? FIXME
alat = 1.d0
gamma_only = .FALSE.
gstart = 2
charge = 0.d0
DO na = 1, nat
   charge = charge + zv( ityp(na) )
ENDDO
alpha = 2.9d0
100 \text{ alpha} = \text{alpha} - 0.1d0
! choose alpha in order to have convergence in the sum over G
! upperbound is a safe upper bound for the error in the sum over {\it G}
IF( alpha <= 0.d0) THEN
 WRITE(*,*) 'ERROR in calculating Ewald energy:'
 WRITE(*,*) 'optimal alpha not found'
 STOP
ENDIF
! beware of unit of gcutm
upperbound = 2.d0*charge**2*sqrt(2.d0*alpha/tpi) * erfc(sqrt(gcutm/4.d0/alpha))
IF(upperbound > 1.0d-7) GOTO 100
! G-space sum here.
! Determine if this processor contains G=0 and set the constant term
IF(gstart==2) THEN
  ewaldg = - charge**2 / alpha / 4.0d0
ELSE
  ewaldg = 0.0d0
ENDIF
! gamma_only should be .FALSE. for our case
IF(gamma_only) THEN
 fact = 2.d0
ELSE
 fact = 1.d0
ENDIF
DO ng = gstart, ngm
 rhon = (0.d0, 0.d0)
 DO nt = 1, ntyp
    rhon = rhon + zv(nt)*CONJG(strf(ng, nt))
 ENDDO
  ewaldg = ewaldg + fact*abs(rhon) **2 * exp( -gg(ng)/alpha/4.d0 )/ gg(ng)
ewaldg = 2.d0 * tpi / omega * ewaldg
! Here add the other constant term
IF (gstart==2) THEN
  DO na = 1, nat
    ewaldg = ewaldg - zv(ityp(na))**2 * sqrt(8.d0/tpi*alpha)
 ENDDO
ENDIF
! R-space sum here (only for the processor that contains G=0)
ewaldr = 0.d0
IF( gstart==2 ) THEN
  rmax = 4.d0 / sqrt(alpha) / alat
  ! with this choice terms up to ZiZj*erfc(4) are counted (erfc(4)=2x10^-8)
 DO na = 1, nat
    DO nb = 1, nat
      dtau(:) = tau(:,na) - tau(:,nb)
```

```
! generates nearest-neighbors shells
      CALL rgen( dtau, rmax, mxr, at, bg, r, r2, nrm )
      ! and sum to the real space part
      DO nr = 1, nrm
       rr = sqrt (r2 (nr)) * alat
        ewaldr = ewaldr + zv (ityp (na) ) * zv (ityp (nb) ) * erfc( sqrt (alpha) * rr) / rr
      ENDDO
    ENDDO
 ENDDO
ENDIF
E_nn = 0.5d0*(ewaldg + ewaldr)
Listing of rgen.f90:
SUBROUTINE rgen ( dtau, rmax, mxr, at, bg, r, r2, nrm )
      generates neighbours shells (cartesian, in units of lattice parameter)
     with length < rmax, and returns them in order of increasing length:
       r(:) = i*a1(:) + j*a2(:) + k*a3(:) - dtau(:), r2 = r^2
     where a1, a2, a3 are primitive lattice vectors. Other input variables:
      mxr = maximum number of vectors
       at = lattice \ vectors \ (a1=at(:,1), a2=at(:,2), a3=at(:,3))
       bg = reciprocal\ lattice\ vectors\ (\ b1=bg(:,1),\ b2=bg(:,2),\ b3=bg(:,3)\ )
     Other output variables:
       nrm = the number of vectors with r^2 < rmax^2
  IMPLICIT NONE
  INTEGER, PARAMETER :: DP=8
  INTEGER, INTENT(in) :: mxr
  INTEGER, INTENT(out):: nrm
  REAL(DP), INTENT(in) :: at(3,3), bg(3,3), dtau(3), rmax
 REAL(DP), INTENT(out):: r(3,mxr), r2(mxr)
  !
       and here the local variables
  INTEGER, ALLOCATABLE :: irr (:)
  INTEGER :: nm1, nm2, nm3, i, j, k, ipol, ir, indsw, iswap
  real(DP) :: ds(3), dtau0(3)
  real(DP) :: t (3), tt, swap
  real(DP), EXTERNAL :: dnrm2
  !
  nrm = 0
  IF (rmax==0.d0) RETURN
  ! bring dtau into the unit cell centered on the origin - prevents trouble
  ! if atomic positions are not centered around the origin but displaced
  ! far away (remember that translational invariance allows this!)
  ds(:) = matmul( dtau(:), bg(:,:) )
  ds(:) = ds(:) - anint(ds(:))
  dtau0(:) = matmul( at(:,:), ds(:) )
  ALLOCATE (irr( mxr))
  ! these are estimates of the maximum values of needed integer indices
  nm1 = int (dnrm2 (3, bg (1, 1), 1) * rmax) + 2
  nm2 = int (dnrm2 (3, bg (1, 2), 1) * rmax) + 2
  nm3 = int (dnrm2 (3, bg (1, 3), 1) * rmax) + 2
  !
```

```
DO i = -nm1, nm1
    DO j = -nm2, nm2
        DO k = -nm3, nm3
           tt = 0.d0
           DO ipol = 1, 3
              t (ipol) = i*at (ipol, 1) + j*at (ipol, 2) + k*at (ipol, 3) &
                       - dtau0(ipol)
              tt = tt + t (ipol) * t (ipol)
           IF (tt<=rmax**2.and.abs (tt) >1.d-10) THEN
              nrm = nrm + 1
              IF (nrm>mxr) then
                WRITE(*,*) 'ERROR in rgen: too many r-vectors', nrm
                STOP
              ENDIF
              DO ipol = 1, 3
                 r (ipol, nrm) = t (ipol)
              ENDDO
              r2 (nrm) = tt
           ENDIF
        ENDDO
    ENDDO
 ENDDO
     reorder the vectors in order of increasing magnitude
      initialize the index inside sorting routine
  irr (1) = 0
  IF (nrm>1) CALL hpsort (nrm, r2, irr)
 D0 ir = 1, nrm - 1
    indsw = irr (ir)
    IF (indsw/=ir) THEN
        DO ipol = 1, 3
           swap = r (ipol, indsw)
           r (ipol, indsw) = r (ipol, irr (indsw) )
           r (ipol, irr (indsw) ) = swap
        ENDDO
        iswap = irr (ir)
        irr (ir) = irr (indsw)
        irr (indsw) = iswap
        GOTO 20
    ENDIF
  ENDDO
 DEALLOCATE(irr)
 RETURN
END SUBROUTINE rgen
```

#### 5.11 Energy minimization

Using nonlinear conjugate gradient algorithm:

```
ALLOCATE( g(Npoints,Nstates) )
ALLOCATE( g_old(Npoints,Nstates) )
ALLOCATE( g_t(Npoints,Nstates) )
ALLOCATE( d(Npoints,Nstates) )
ALLOCATE( d_old(Npoints,Nstates) )
ALLOCATE( Kg(Npoints,Nstates) )
```

```
ALLOCATE( Kg_old(Npoints, Nstates) )
ALLOCATE( tv(Npoints, Nstates) )
! Read starting eigenvectors from file
IF( restart ) THEN
 READ(112) v ! FIXME Need to use file name
ENDIF
CALL calc_Rhoe( Focc, v )
CALL update_potentials()
CALL calc_betaNL_psi( Nstates, v )
CALL calc_energies( v )
Etot_old = Etot
alpha = 0.d0
beta = 0.d0
g(:,:)
          = 0.d0
g_t(:,:) = 0.d0
d(:,:)
        = 0.d0
d_old(:,:) = 0.d0
         = 0.d0
Kg(:,:)
Kg_old(:,:) = 0.d0
D0 iter = 1, Emin_NiterMax
  ! Evaluate gradient at current trial vectors
 CALL calc_grad( Nstates, v, g )
  ! Precondition
  DO ist = 1, Nstates
    CALL prec_ilu0( g(:,ist), Kg(:,ist) )
  ENDDO
  ! set search direction
  IF( iter /= 1 ) THEN
    SELECT CASE ( I_CG_BETA )
    CASE(1)
      ! Fletcher-Reeves
      beta = sum(g * Kg) / sum(g_old * Kg_old)
    CASE(2)
      ! Polak-Ribiere
      beta = sum((g-g_old)*Kg) / sum(g_old * Kg_old)
      ! Hestenes-Stiefel
     beta = sum((g-g_old)*Kg) / sum((g-g_old)*d_old)
    CASE(4)
      ! Dai-Yuan
     beta = sum(g * Kg) / sum((g-g_old)*d_old)
    END SELECT
  ENDIF
  IF( beta < 0 ) THEN</pre>
    WRITE(*,'(1x,A,F18.10,A)') 'beta is smaller than zero: ', beta, ': setting it to zero'
  beta = max(0.d0, beta)
  d(:,:) = -Kg(:,:) + beta*d_old(:,:)
  ! Evaluate gradient at trial step
  tv(:,:) = v(:,:) + alpha_t * d(:,:)
  CALL orthonormalize( Nstates, tv )
  CALL calc_Rhoe( Focc, tv )
  CALL update_potentials() ! Now global vars on m_hamiltonian are changed
  CALL calc_betaNL_psi( Nstates, tv )
  CALL calc_grad( Nstates, tv, g_t)
  ! Compute estimate of best step and update current trial vectors
```

```
denum = sum((g - g_t) * d)
  IF( denum /= 0.d0 ) THEN  ! FIXME: use abs ?
    alpha = abs( alpha_t * sum( g * d )/denum )
  ELSE
    alpha = 0.d0
 ENDIF
  v(:,:) = v(:,:) + alpha * d(:,:)
  CALL orthonormalize (Nstates, v)
  CALL calc_Rhoe( Focc, v )
  CALL update_potentials()
  CALL calc_betaNL_psi( Nstates, v )
  CALL calc_energies( v )
  WRITE(*,'(1x,I5,F18.10,ES18.10)') iter, Etot, Etot_old-Etot
  IF( abs(Etot - Etot_old) < Emin_ETOT_CONV_THR ) THEN</pre>
    WRITE(*,*) 'KS_solve_Emin_pcg converged in iter', iter
    EXIT
 ENDIF
 Etot_old = Etot
  g_old(:,:) = g(:,:)
 d_old(:,:) = d(:,:)
 Kg_old(:,:) = Kg(:,:)
  flush(6)
ENDDO
Calculation of gradient:
ALLOCATE( Hv(Npoints) )
DO ic = 1, Ncols
  CALL op_H_1col( ic, v(:,ic), Hv(:) )
  grad(:,ic) = Hv(:)
 DO icc = 1, Ncols
    grad(:,ic) = grad(:,ic) - ddot( Npoints, v(:,icc),1, Hv(:),1 )*v(:,icc)*dVol
 grad(:,ic) = Focc(ic)*grad(:,ic)
ENDDO
5.12
       Self-consistent field
ALLOCATE( Rhoe_old(Npoints) )
beta0 = SCF_betamix
betamax = 1.d0
mixsdb = 4
broydpm(1) = 0.4d0
broydpm(2) = 0.15d0
ALLOCATE( beta_work(Npoints) )
ALLOCATE( f_work(Npoints) )
Etot_old = 0.d0
Rhoe_old(:) = Rhoe(:)
! Allocate memory for ELK mixing subroutines
! Linear mixing
IF( MIXTYPE == 0 ) THEN
 nwork = Npoints
 ALLOCATE( workmix(nwork) )
! Adaptive linear mixing
ELSEIF( MIXTYPE == 1 ) THEN
```

```
nwork = 3*Npoints
 ALLOCATE( workmix(nwork) )
! Broyden mixing
ELSEIF( MIXTYPE == 3 ) THEN
 nwork = (4+2*mixsdb)*Npoints + mixsdb**2
 ALLOCATE( workmix(nwork) )
ELSE
 WRITE(*,*) 'Unknown MIXTYPE: ', MIXTYPE
 WRITE(*,*) 'Switching to default: adaptive linear mixing'
 MIXTYPE = 1
 nwork = 3*Npoints
 ALLOCATE( workmix(nwork) )
ENDIF
flush(6)
dr2 = 1.d0
DO iterSCF = 1, SCF_NiterMax
  IF( iterSCF==1 ) THEN
    ethr = 1.d-1
 ELSE
    IF( iterSCF == 2 ) ethr = 1.d-2
    ethr = ethr/5.d0
    ethr = max( ethr, ETHR_EVALS_LAST )
  ENDIF
  CALL Sch_solve_diag()
  CALL calc_rhoe( Focc, evecs )
  CALL mixerifc( iterSCF, MIXTYPE, Npoints, Rhoe, dr2, nwork, workmix )
  CALL normalize_rhoe( Npoints, Rhoe ) ! make sure no negative or very small rhoe
  integRho = sum(Rhoe)*dVol
  IF( abs(integRho - Nelectrons) > 1.0d-6 ) THEN
    WRITE(*,'(1x,A,ES18.10)') 'WARNING: diff after mix rho = ', abs(integRho-Nelectrons)
    WRITE(*,*) 'Rescaling Rho'
    Rhoe(:) = Nelectrons/integRho * Rhoe(:)
    integRho = sum(Rhoe)*dVol
    WRITE(*,'(1x,A,F18.10)') 'After rescaling: integRho = ', integRho
    WRITE(*,*)
  ENDIF
  CALL update_potentials()
  CALL calc_betaNL_psi( Nstates, evecs )
  CALL calc_energies( evecs ) ! update the potentials or not ?
  dEtot = abs(Etot - Etot_old)
  IF( dEtot < 1d-7) THEN
    WRITE(*,*)
    WRITE(*,'(1x,A,I5,A)') 'SCF converged at ', iterSCF, ' iterations.'
   EXIT
  ENDIF
  WRITE(*,'(1x,A,I5,F18.10,2ES18.10)') 'SCF iter', iterSCF, Etot, dEtot, dr2
  Etot_old = Etot
  Rhoe_old(:) = Rhoe(:)
  flush(6)
ENDDO
```

### A Lagrange basis function

#### A.1 Periodic Lagrange function

For a given interval [0, L], with L > 0, the grid points  $x_i$  appropriate for periodic Lagrange function are given by:

$$x_i = \frac{L}{2} \frac{2i - 1}{N} \tag{18}$$

with i = 1, ..., N. Number of points N should be an odd number.

The periodic cardinal functions  $L_i^{\text{per}}(x)$ , defined at grid point i are given by:

$$L_i^{\text{per}}(x) = \frac{1}{\sqrt{NL}} \sum_{n=1}^{N} \cos\left(\frac{\pi}{L} (2n - N - 1)(x - x_i)\right). \tag{19}$$

The expansion of periodic function in terms of Lagrange functions:

$$f(x) = \sum_{i=1}^{N} c_i L_i^{\text{per}}(x)$$
(20)

with expansion coefficients  $c_i = \sqrt{L/N} f(x_i)$ . When doing variational calculation, the cofficients  $c_i$  are the variational parameters. The actual function values  $f(x_i)$  at grid points  $x_i$  is obtained by  $f(x_i) = \sqrt{N/L}c_i$ . The prefactor is sometimes abbreviated by h = L/N and is also referred to as scaling factor.

Consider periodic potential in one dimension:

$$V(x+L) = V(x). (21)$$

Floquet-Bloch theorem states that the wave function solution for periodic potentials can be written in the form:

$$\psi_k(x) = e^{ikx}\phi_k(x) \tag{22}$$

where function  $\phi_k(x)$  and its first derivative  $\phi'_k(x)$  have the same periodicity as V(x) and k is a constant called the crystal momentum. Substituting this expression to Schrödinger equation we obtain:

$$\left[ -\frac{\hbar^2}{2m} \left( \frac{\mathrm{d}^2}{\mathrm{d}x^2} + 2ik \frac{\mathrm{d}}{\mathrm{d}x} - k^2 \right) + V(x) \right] \phi_k(x) = E\phi_k(k). \tag{23}$$

An alternative way of enforcing periodicity of the wave function is to require that:

$$\psi_k(x+L) = e^{ikL}\psi_k(x). \tag{24}$$

This condition follows from:

$$\psi_k(x+L) = e^{ik(x+L)}\phi_k(x+L)$$

$$= e^{ik(x+L)}\phi_k(x)$$

$$= e^{ikL}e^{ikx}\phi_k(x)$$

$$= e^{ikL}\psi_k(x)$$

Using periodic cardinal the Schrodinger equation for periodic potential can be written as:

$$\sum_{j=1}^{N} \left[ -\frac{\hbar^2}{2m} \left( D_{jl}^{(2)} + 2\imath k D_{jl}^{(1)} - k^2 \delta_{jl} \right) + V(j) \delta_{jl} \right] \phi(j) = E\phi(l)$$
(25)

with l = 1, ..., N.  $D_{il}^{(1)}$  are matrix elements of the first derivatives:

$$D_{jl}^{(1)} = \begin{cases} 0 & j = l \\ -\frac{2\pi}{L} (-1)^{j-l} \left( 2\sin\frac{\pi(j-l)}{N} \right)^{-1} & j \neq l \end{cases}$$
 (26)

and  $D_{il}^{(2)}$  are matrix elements of the second derivatives, N' = (N-1)/2:

$$D_{jl}^{(2)} = \begin{cases} -\left(\frac{2\pi}{L}\right)^2 \frac{N'(N'+1)}{3} & j=l\\ -\left(\frac{2\pi}{L}\right)^2 (-1)^{j-l} \frac{\cos(\pi(j-l)/N)}{2\sin^2[\pi(j-l)/N]} & j \neq l \end{cases}$$
(27)

Note that,  $D_{jl}^{(1)}$  is not symmetric, but  $D_{jl}^{(1)} = -D_{lj}^{(1)}$ . Meanwhile, the second derivative matrix  $D_{jl}^{(2)}$  is symetric, i.e.  $D_{jl}^{(2)} = D_{lj}^{(2)}$ . With the above expressions, first and second derivative of periodic cardinals can be expressed as

$$\frac{\mathrm{d}}{\mathrm{d}x}L_i^{\mathrm{per}}(x) = \sum_{j=1}^N D_{ji}^{(1)} L_j^{\mathrm{per}}(x)$$
 (28)

$$\frac{\mathrm{d}^2}{\mathrm{d}x^2} L_i^{\text{per}}(x) = \sum_{j=1}^N D_{ji}^{(2)} L_j^{\text{per}}(x)$$
(29)

The previous approach also can be extended to periodic potential in 3D:

$$V(\mathbf{r}) = V(x, y, z) = V(x + L_x, y + L_y, z + L_z)$$

Using periodic LF, Schrodinger equation can be casted into the following form:

$$\left[ -\frac{\hbar^2}{2m} \left( \nabla^2 + 2i \mathbf{k} \cdot \nabla - \mathbf{k}^2 \right) + V(\mathbf{r}) \right] \phi_{\mathbf{k}}(\mathbf{r}) = E \ \phi_{\mathbf{k}}(\mathbf{r})$$
(30)

#### A.2 Cluster Lagrange function

For a given interval [A, B], with B > A, the grid points  $x_i$  appropriate for cluster Lagrange function are given by:

$$x_i = A + \frac{B - A}{N + 1}i$$

where i = 1, ..., N. Number of points N can be either odd or even number.

The cluster Lagrange functions  $L_i^{\text{clu}}(x)$ , defined at grid point i are given by:

$$L_i^{\text{clu}}(x) = \frac{2}{\sqrt{(N+1)(B-A)}} \sum_{n=1}^N \sin(k_n(x_i - A)) \sin(k_n(x - A)).$$
 (31)

where  $k_n = \pi n/(B-A)$ . The expansion of a function f(x) in terms of cluster Lagrange functions:

$$f(x) = \sum_{i=1}^{N} c_i L_i^{\text{clu}}(x)$$
(32)

with expansion coefficients  $c_i = \sqrt{(B-A)/(N+1)}f(x_i)$ . When doing variational calculation, the coefficients  $c_i$  are the variational parameters. The actual function values  $f(x_i)$  at grid points  $x_i$  is obtained by  $f(x_i) = \sqrt{(N+1)/(B-A)}c_i$ .

Matrix elements  $D_{jl}^{(2)}$  of the second derivatives for cluster Lagrange functions are

$$D_{jl}^{(2)} = \begin{cases} -\frac{1}{2} \left(\frac{\pi}{B-A}\right)^2 \frac{2(N+1)^2 + 1}{3} - \frac{1}{\sin^2\left[\pi j/(N+1)\right]} & j = l\\ -\frac{1}{2} \left(\frac{\pi}{B-A}\right)^2 (-1)^{j-l} \left[\frac{1}{\sin^2\left[\frac{\pi(j-l)}{2(N+1)}\right]} - \frac{1}{\sin^2\left[\frac{\pi(j+l)}{2(N+1)}\right]}\right] & j \neq l \end{cases}$$
(33)

For free or cluster boundary condition, we don't need  $D_{jl}^{(1)}$ .

### B HGH pseudopotential

HGH pseudopotential has analytic forms both in real space and reciprocal space. Local component of pseudopotential in real space

$$V_{\text{loc}}(\mathbf{r}) = -\frac{Z_{\text{ion}}}{r} \operatorname{erf}\left(\frac{r}{\sqrt{2}r_{\text{loc}}}\right) + \exp\left[-\frac{1}{2}\left(\frac{r}{r_{\text{loc}}}\right)^{2}\right] \times \left[C_{1} + C_{2}\left(\frac{r}{r_{\text{loc}}}\right)^{2} + C_{3}\left(\frac{r}{r_{\text{loc}}}\right)^{4} + C_{4}\left(\frac{r}{r_{\text{loc}}}\right)^{6}\right]$$
(34)

with parameters:  $r_{loc}$ ,  $C_1$ ,  $C_2$ ,  $C_3$ , and  $C_4$ .

Local component of local pseudopotential in **G**-space:

$$V_{\text{loc}}(\mathbf{G}) = -\frac{1}{\Omega} \frac{4\pi Z_{\text{ion}}}{G^2} \exp\left[-\frac{1}{2} (Gr_{\text{loc}})^2\right] + \sqrt{8\pi^3} \frac{r_{\text{loc}}}{\Omega} \exp\left[-\frac{1}{2} (Gr_{\text{loc}})^2\right] \times \left\{C_1 + C_2 \left[3 - (Gr_{\text{loc}})^2\right] + C_3 \left[15 - 10 (Gr_{\text{loc}})^2 (Gr_{\text{loc}})^4\right] + C_4 \left[105 - 105 (Gr_{\text{loc}})^2 + 21 (Gr_{\text{loc}})^4 - (Gr_{\text{loc}})^6\right]\right\}$$
(35)

Nonlocal component of pseudopotential can be written as

$$V_{l}(\mathbf{r}, \mathbf{r}') = \sum_{i=1}^{3} \sum_{j=1}^{3} \sum_{m=-l}^{l} \beta_{ilm}(\mathbf{r}) h_{ij}^{l} \beta_{jlm}^{*}(\mathbf{r}')$$
(36)

with atomic-centered functions projector functions as

$$\beta_{ilm}(\mathbf{r}) = p_i^l(r)Y_{lm}(\hat{\mathbf{r}}) \tag{37}$$

The radial projector functions have the following form in real space

$$p_i^l(r) = \frac{\sqrt{2}r^{l+2(i-1)} \exp\left(-\frac{r^2}{2r_l^2}\right)}{r_l^{l+(4i-1)/2} \sqrt{\Gamma\left(l + \frac{4i-1}{2}\right)}}$$
(38)

The radial projector functions satisfy the following normalization condition

$$\int_0^\infty p_i^l(r)p_i^l(r)\,r^2\,\mathrm{d}r = 1$$
(39)

For l = 0, the Fourier transform of radial projector functions can be written as:

$$p_1^{l=0}(G) = \frac{4\sqrt{2r_0^3}\pi^{5/4}}{\sqrt{\Omega}\exp\left[(Gr_0)^2/2\right]}$$
(40)

$$p_2^{l=0}(G) = \frac{\sqrt{8\frac{2r_0^3}{15}}\pi^{5/4} \left(3 - (Gr_0)^2\right)}{\sqrt{\Omega}\exp\left[(Gr_0)^2/2\right]}$$
(41)

$$p_3^{l=0}(G) = \frac{16\sqrt{\frac{2r_0^3}{105}}\pi^{5/4} \left(15 - 10(Gr_0)^2 - (Gr_0)^4\right)}{3\sqrt{\Omega}\exp\left[(Gr_0)^2/2\right]}$$
(42)

For l = 1, the Fourier transform of radial projector functions can be written as

$$p_1^{l=1}(G) = \frac{8\sqrt{\frac{r_1^5}{3}}\pi^{5/4}G}{\sqrt{\Omega}\exp\left[(Gr_1)^2/2\right]}$$
(43)

$$p_2^{l=1}(G) = \frac{16\sqrt{\frac{r_1^5}{105}}\pi^{5/4} \left(5 - (Gr_1)^2\right) G}{\sqrt{\Omega} \exp\left[(Gr_1)^2/2\right]}$$
(44)

$$p_3^{l=1}(G) = \frac{32\sqrt{\frac{r_1^5}{1155}}\pi^{5/4} \left(35 - 14(Gr_1)^2 + (Gr_1)^4\right)G}{3\sqrt{\Omega}\exp\left[(Gr_1)^2/2\right]}$$
(45)

For l=2, the Fourier transform of radial projector functions can be written as

$$p_1^{l=2}(G) = \frac{8\sqrt{\frac{2r_2^7}{15}}\pi^{5/4}G^2}{\sqrt{\Omega}\exp\left[(Gr_2)^2/2\right]}$$
(46)

$$p_2^{l=2}(G) = \frac{16\sqrt{\frac{2r_2^7}{105}}\pi^{5/4} \left(7 - (Gr_2)^2\right) G^2}{3\sqrt{\Omega} \exp\left[(Gr_2)^2/2\right]}$$
(47)

For l=3, the Fourier transform of radial projector function can be written as

$$p_1^{l=3}(G) = \frac{16\sqrt{\frac{2r_3^9}{105}}\pi^{5/4}G^3}{\sqrt{\Omega}\exp\left[(Gr_3)^2/2\right]}$$
(48)

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