# **PWDFT.jl Documentation**

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## Part I. User Guide

### 1 Overview

PWDFT.jl is a Julia package to carry out (electronic) density functional theory calculations on materials. It uses plane wave basis sets to discretize Kohn-Sham equations. It also uses pseudopotentials to replace strong Coulombic nuclei-electron interactions  $^1$ .

I assumes that the readers already have some experience with similar electronic structure programs such as Quantum Espresso, ABINIT, VASP, and others.

PWDFT.jl uses Hartree atomic units internally.

#### 2 Installation

You need the following to use PWDFT.jl:

- Julia version 0.7 or higher
- LibXC

<sup>&</sup>lt;sup>1</sup> You can carry out the calculation with full potential, however this will requires a lot of plane waves (large cutoff energy)

3 Getting started 2

- SPGLIB
- C compiler to compiler LibXC and SPGLIB

PWDFT.jl is a Julia package, so you need to copy (or clone) under Julia's package depot so that it can be used within Julia. Presently, it is not yet a registered package. There are at least two ways of doing this.

The first way is using Julia's package manager. The following command can be issued at the Julia's console.

```
Pkg.add(PackageSpec(url="https://github.com/f-fathurrahman/PWDFT.jl"))
```

The second way is to use Julia's development directory located at \$HOME/.julia/dev. I usually do this by sym-linking my original PWDFT.jl directory to \$HOME/.julia/dev.

To test the installation, you can run the following:

```
using PWDFT
```

at the Julia's console. If there are no error messages then the installation is OK. <sup>2</sup>

## 3 Getting started

We need to prepare several things before using PWDFT.jl:

- Atomic structures: coordinates and unit cell
- Pseudopotentials

This code will initialize a hydrogen molecule (H2) inside a box of  $16 \times 16 \times 16$  bohr.

#### 4 Status

28 May 2018 The following features are working now:

- LDA and GGA, spin-paired and spin polarized calculations
- Calculation with k-points (for periodic solids). SPGLIB is used to reduce the Monkhorst-Pack grid points for integration over Brillouin zone.

Band structure calculation is possible in principle as this can be done by simply solving Schrodinger equation with converged Kohn-Sham potentials, however there is currently no tidy script or function to do that.

Total energy result for isolated systems (atoms and molecules) agrees quite well with ABINIT and PWSCF results.

Total energy result for periodic solid is quite different from ABINIT and PWSCF. I suspect that this is related to treatment of electrostatic terms in periodic system.

SCF is rather shaky for several systems, however it is working in quite well in nonmetallic system.

## Part II. Implementation Notes

In this note, I will describe several user-defined types that are used throughout PWDFT.jl. Beginners are not expected to know them all, however they are needed to know the internal of PWDFT.jl.

When referring to a file, I always meant to the file contained in the src/ directory.

Even though it is not strictly required in Julia, I tried to always use type annotations.

 $<sup>^{2}</sup>$  This does not check the LibXC or SPGLIB installation.

5 Atoms 3

#### 5 Atoms

Atoms can be used to represent molecular or crystalline structures. This type is implemented in the file src/Atoms.jl. It has the following fields:

- Natoms::Int64: number of atoms present in the system
- Nspecies::Int64: number of atomic species present in the system
- positions::Array{Float64,2}: An array containing coordinates of atoms in bohr units. Its shape is (3, Natoms)
- atm2species::Array{Int64,1}: An array containing mapping between atom index to species index. Its shape is (Natoms,)
- atsymbs::Array{String,1}: An array containing atomic symbols for each atoms. Its shape is (Natoms,).
- SpeciesSymbols::Array{String,1}: An array containing unique symbols for each atomic species present in the system. Its shape is (Nspecies,).
- LatVecs::Array{Float64,2}: A 3 by 3 matrix describing lattice vectors for unit cell of the system.
- Zvals::Array{Float64,1}: An array containing number of (valence) electrons of each atomic species. Its shape is (Nspecies,).

An instance of Atoms can be initialized using any of the following ways:

• Using Atoms constructor, which has the following signature

• The above mentioned Atoms constructor is actually a wrapper for the functions init\_atoms\_xyz and init\_atoms\_xyz\_stri
which takes either a string containing path to xyz file or the content of the xyz file itself. When using this function,
one must set LatVecs. field manually.

A note about Zvals: Both Atoms and init\_atoms\_\* function set Zvals to zeros(Nspecies). After passed to Hamiltonian constructor Zvals will be set according to the pseudopotentials used.

TODO: examples

#### 6 Hamiltonian

An instance Hamiltonian is a central object in PWDFT.jl. It is used to store various instances of other important types such as atoms, plane wave grids, pseudopotentials, etc. It is implemented in the file src/Hamiltonian.jl.

To create an instance of Hamiltonian, we normaly need to provide at least three arguments to the Hamiltonian constructor:

- atoms::Atoms: an instance of Atoms
- pspfiles::Array{String,1}: a list of strings specifying the locations of pseudopotentials used in the calculations. Note that, the order should be the same as species ordering of atoms, i.e. pspfiles[isp] is the path of pseudopotentials of species with symbols atoms.SpeciesSymbols[isp].
- ecutwfc::Float64: cutoff energy for wave function expansion using plane wave basis set.

A simplified version of Hamiltonian constructor only needs two arguments: atoms::Atoms and ecutwfc::Float64. In this case, full Coulomb potential will be used. We usually need very large cutoff energy in this case (probably in the order or 10<sup>2</sup> Hartree to obtain good convergence).

The structure of Hamiltonian is designed such that we can perform application or multiplication of Hamiltonian to wave function:

```
Hpsi = op_H(H, psi)
or, (by overloading the * operator 3)
Hpsi = H*psi
```

 $<sup>^3</sup>$  The operator  $^\ast$  is actually implemented as function in Julia

7 Plane wave basis 4

#### 7 Plane wave basis

The plane wave basis is described by the type PWGrid. This type is defined in the file PWGrid.jl. It has the following fields:

- ecutwfc::Float64: cutoff for wave function expansion
- ecutrho::Float64: cutoff for electron density expansion, for norm-converving pseudopotential: ecutrho = 4\*ecutwfc.
- Ns::Tuple{Int64, Int64, Int64}: parameters defining real-space grid points.
- LatVecs::Array{Float64,2}: lattice vectors of unit cell  $(3 \times 3 \text{ matrix})$
- RecVecs::Array{Float64,2}: reciprocal lattice vectors (3 × 3 matrix)
- CellVolume::Float64: the volume of real-space unit cell
- r::Array{Float64,2}: real-space grid points. Its shape is (3, Npoints)
- gvec::GVectors: an instace of GVectors: for potentials and density expansion
- gvecw::GVectorsW: an instace of GVectorsW, for wave function expansion
- planfw::FFTW.cFFTWPlan{Complex{Float64},-1,false,3}: FFTW forward plan
- planbw::AbstractFFTs.
  ScaledPlan{ComplexF64,FFTW.cFFTWPlan{ComplexF64,1,false,3-Float64}}: FFTW backward plan

The following constructor can be used to create an instance of 'PWGrid':

## 7.1 G-vectors for potentials and densities expansion

G-vectors are described by type 'GVectors'. It is defined in file 'PWGrid.jl'. It has the following fields:

- Ng::Int64: total number of G-vectors
- G::Array{Float64,2} The array containing the actual G-vectors. Its shape is (3,Ng).
- G2::Array{Float64,1}: The array containing magnitude of G-vectors. Its shape is (Ng,).
- idx\_g2r::Array{Int64,1} The array containing mapping between G-vectors to real space grid points. Its shape is (Ng,).

The following function is used as the constructor:

```
function init_gvec( Ns, RecVecs, ecutrho )
```

This function takes the following arguments

- Ns: a tuple of three Int64's specifying sampling points along the 1st, 2nd, and 3rd lattice vector directions.
- RecVecs: 3 by 3 matrix describing reciprocal lattice vectors
- ecutrho: cutoff energy (in Hartree). For norm-conserving PP, it is 4 times 'ecutwfc'.

#### 7.2 G-vectorsW

The **G**-vectors for wave function expansion is described by the type **GVectorsW**. They are a subset of **GVectors**. It has the following fields:

- Ngwx::Int64: maximum number of G-vectors for all kpoints.
- Ngw::Array{Int64,1}: number of G-vectors for each kpoints
- idx\_gw2g::Array{Array{Int64,1},1}: mapping between indices of GVectorsW to indices of GVectors.
- idx\_gw2r::Array{Array{Int64,1},1}: mapping between indices of 'GVectorsW' to indices of real space grid points PWGrid.r
- kpoints:: KPoints: an instance of KPoints

Constructor: TODO

8 Wave functions

#### 7.3 Bloch wave vector

The type describing Bloch wave vectors, or commonly referred to as k-points, is KPoints. It is defined in the file KPoints.jl. It has the following fields

- Nkpt::Int64: total number of k-points.
- k::Array{Float64,2}: the actual k-points. Its shape is (3,Nkpt).
- wk::Array{Float64,1}: the weight of each k-points needed for integration over Brillouin zone
- RecVecs::Array{Float64,2}: a copy of PWGrid.RecVecs.

Monkhorst-Pack grid points:

$$k_i = \frac{2n_i - N_i - 1}{2N_i} \mathbf{b}_i. \quad i = 1, 2, 3$$
 (1)

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where  $n_i = 1, 2, ..., N_i$  size =  $(N_1, N_2, N_3)$  and the  $\mathbf{b}_i$ 's are reciprocal lattice vectors.

Constructor:

SPGLIB is used internally to generate reduce number of k-points.

#### 8 Wave functions

Using 'Array{ComplexF64,2}.
General wavefunction on kpoints

## 9 Total energy components

Total energy components are stored in the type Energies. Currently its fields are as follows.

- 'Kinetic': kinetic energy
- Ps\_loc: local pseudopotential energy
- Ps\_nloc: nonlocal pseudopotential energy
- Hartree: classical electrostatic energy
- XC: exchange correlation energy
- NN: nuclear-nuclear (repulsive) interaction energy
- PspCore: core (screened) pseudopotential energy
- mTS: electronic entropy contribution (with minus sign). This is used for calculation with partial occupations such as metals.

#### 10 Local potentials

Various local potentials are stored in the type Potentials. Currently its fields are as follows.

- Ps\_loc local pseudopotential components. Its shape is (Npoints,).
- Hartree: classical electrostatic potential. Its shape is (Npoints,).
- XC: exchange-correlation potential. This potential can be spin dependent so its shape is (Npoints, Nspin).

All of these potentials are in the real space representation.

11 Pseudopotentials

6

## 11 Pseudopotentials

Currently, only GTH pseudopotentials with no core-correction are supported. The type for handling GTH pseudopotential for a species is PsPot\_GTH. It is declared as follows.

```
struct PsPot_GTH
   pspfile::String
   atsymb::String
  zval::Int64
  rlocal::Float64
  rc::Array{Float64,1}
  c::Array{Float64,3}
  lmax::Int64
  Nproj_l::Array{Int64,1}
  rcut_NL::Array{Float64,1}
end
```

## 12 Solving Kohn-Sham problems

Two main algorithms: SCF and direct minimization Function names: KS\_solve\_\*

## 12.1 Self-consistent field

```
KS_solve_SCF
Solves Kohn-Sham problem using traditional self-consistent field (SCF)
iterations with density mixing.
function KS_solve_SCF!( Ham::Hamiltonian ;
                        startingwfc=nothing, savewfc=false,
                        betamix = 0.5, NiterMax=100, verbose=true,
                        print_final_ebands=true,
                        print_final_energies=true,
                        check_rhoe_after_mix=false,
                        use_smearing = false, kT=1e-3,
                        update_psi="LOBPCG", cheby_degree=8,
                        mix_method="simple", MIXDIM=4,
                        ETOT_CONV_THR=1e-6 )
    pw = Ham.pw
    Ngw = pw.gvecw.Ngw
    wk = Ham.pw.gvecw.kpoints.wk
    kpoints = pw.gvecw.kpoints
    Nkpt = kpoints.Nkpt
    Ns = pw.Ns
    Npoints = prod(Ns)
    CellVolume = pw.CellVolume
    dVol = CellVolume/Npoints
    electrons = Ham.electrons
    Nelectrons = electrons.Nelectrons
    Focc = copy(electrons.Focc) # make sure to use the copy
    Nstates = electrons.Nstates
    Nspin = electrons.Nspin
    Nkspin = Nkpt*Nspin
```

```
Nstates_occ = electrons.Nstates_occ
# Random guess of wave function
if startingwfc==nothing
    psiks = rand_BlochWavefunc(pw, electrons)
    psiks = startingwfc
end
# Calculated electron density from this wave function and update Hamiltonian
Rhoe = zeros(Float64, Npoints, Nspin)
for ispin = 1:Nspin
    idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
    Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
end
if Nspin == 2
    @printf("\nInitial integ magn_den = %18.10f\n", sum(Rhoe[:,1] - Rhoe[:,2])*dVol)
end
update! (Ham, Rhoe)
Etot_old = 0.0
Rhoe_new = zeros(Float64, Npoints, Nspin)
diffRhoe = zeros(Nspin)
evals = zeros(Float64, Nstates, Nkspin)
ETHR\_EVALS\_LAST = 1e-6
ethr = 0.1
if mix_method == "anderson"
    df = zeros(Float64, Npoints*Nspin, MIXDIM)
    dv = zeros(Float64, Npoints*Nspin, MIXDIM)
elseif mix_method in ("rpulay", "rpulay_kerker")
    XX = zeros(Float64, Npoints*Nspin, MIXDIM)
    FF = zeros(Float64, Npoints*Nspin, MIXDIM)
    x_old = zeros(Float64, Npoints*Nspin)
    f_old = zeros(Float64, Npoints*Nspin)
end
E_GAP_INFO = false
Nstates_occ = electrons.Nstates_occ
if Nstates_occ < Nstates
    E_GAP_INFO = true
    if Nspin == 2
        idx_HOMO = max(round(Int64, Nstates_occ/2),1)
        idx_LUMO = idx_HOMO + 1
    else
        idx_HOMO = Nstates_occ
        idx_LUMO = idx_HOMO + 1
    end
end
```

```
@printf("\n")
@printf("Self-consistent iteration begins ...\n")
@printf("update_psi = %s\n", update_psi)
@printf("\n")
@printf("mix_method = %s\n", mix_method)
if mix_method in ("rpulay", "rpulay_kerker", "anderson")
    @printf("MIXDIM = %d\n", MIXDIM)
Oprintf("Density mixing with betamix = 10.5f\n", betamix)
if use_smearing
    Qprintf("Smearing = %f\n", kT)
println("") # blank line before SCF iteration info
# calculate E_NN
Ham.energies.NN = calc_E_NN( Ham.atoms )
# calculate PspCore energy
Ham.energies.PspCore = calc_PspCore_ene( Ham.atoms, Ham.pspots, CellVolume )
CONVERGED = 0
for iter = 1:NiterMax
    # determine convergence criteria for diagonalization
    if iter == 1
        ethr = 0.1
    elseif iter == 2
        ethr = 0.01
    else
        ethr = ethr/5.0
        ethr = max( ethr, ETHR_EVALS_LAST )
    end
    if update_psi == "LOBPCG"
        evals =
        diag_LOBPCG!( Ham, psiks, verbose=false, verbose_last=false,
                      Nstates_conv=Nstates_occ )
    elseif update_psi == "davidson"
        evals =
        diag_davidson!( Ham, psiks, verbose=false, verbose_last=false,
                        Nstates_conv=Nstates_occ )
    elseif update_psi == "PCG"
        evals =
        diag_Emin_PCG!( Ham, psiks, verbose=false, verbose_last=false,
                        Nstates_conv=Nstates_occ )
    elseif update_psi == "CheFSI"
        # evals will be calculated later
        diag_CheFSI!( Ham, psiks, cheby_degree )
    else
        @printf("ERROR: Unknown method for update_psi = %s\n", update_psi)
        error("STOPPED")
```

end

```
if E_GAP_INFO && verbose
        println("E gap = ", minimum(evals[idx_LUMO,:] - evals[idx_HOMO,:]))
    end
    if use_smearing
        Focc, E_fermi = calc_Focc( evals, wk, Nelectrons, kT, Nspin=Nspin )
        Entropy = calc_entropy( Focc, wk, kT, Nspin=Nspin )
        Ham.electrons.Focc = copy(Focc)
    end
    for ispin = 1:Nspin
        idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
        Rhoe_new[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
        diffRhoe[ispin] = norm(Rhoe_new[:,ispin] - Rhoe[:,ispin])
    end
    if mix_method == "simple"
        for ispin = 1:Nspin
            Rhoe[:,ispin] = betamix*Rhoe_new[:,ispin] + (1-betamix)*Rhoe[:,ispin]
        end
    elseif mix_method == "simple_kerker"
        for ispin = 1:Nspin
            Rhoe[:,ispin] = Rhoe[:,ispin] + betamix*precKerker(pw, Rhoe_new[:,ispin] -
Rhoe[:,ispin])
        end
    elseif mix_method == "rpulay"
        Rhoe = reshape( mix_rpulay!(
            reshape(Rhoe,(Npoints*Nspin)),
            reshape(Rhoe_new,(Npoints*Nspin)), betamix, XX, FF, iter, MIXDIM, x_old, f_old
            ), (Npoints, Nspin) )
        if Nspin == 2
            magn_den = Rhoe[:,1] - Rhoe[:,2]
        end
    elseif mix_method == "rpulay_kerker"
        Rhoe = reshape( mix_rpulay_kerker!( pw,
            reshape(Rhoe,(Npoints*Nspin)),
            reshape(Rhoe_new,(Npoints*Nspin)), betamix, XX, FF, iter, MIXDIM, x_old, f_old
            ), (Npoints, Nspin) )
        if Nspin == 2
            magn_den = Rhoe[:,1] - Rhoe[:,2]
        end
    elseif mix_method == "anderson"
        Rhoe[:,:] = mix_anderson!( Nspin, Rhoe, Rhoe_new, betamix, df, dv, iter, MIXDIM )
    else
        @printf("ERROR: Unknown mix_method = %s\n", mix_method)
        error("STOPPED")
    end
    for rho in Rhoe
        if rho < eps()
            rho = 0.0
        end
    end
```

```
# renormalize
    if check_rhoe_after_mix
        integRhoe = sum(Rhoe)*dVol
        @printf("After mixing: integRhoe = %18.10f\n", integRhoe)
        Rhoe = Nelectrons/integRhoe * Rhoe
        integRhoe = sum(Rhoe)*dVol
        @printf("After renormalize Rhoe: = %18.10f\n", integRhoe)
    end
   update! ( Ham, Rhoe )
    # Calculate energies
   Ham.energies = calc_energies( Ham, psiks )
    if use_smearing
        Ham.energies.mTS = Entropy
   end
   Etot = sum(Ham.energies)
   diffE = abs( Etot - Etot_old )
   if verbose
        if Nspin == 1
            @printf("SCF: %8d %18.10f %18.10e %18.10e\n",
                    iter, Etot, diffE, diffRhoe[1] )
        else
            @printf("SCF: %8d %18.10f %18.10e %18.10e \n",
                    iter, Etot, diffE, diffRhoe[1], diffRhoe[2] )
            magn_den = Rhoe[:,1] - Rhoe[:,2]
            Oprintf("integ magn_den = %18.10f\n", sum(magn_den)*dVol)
        end
    end
    if diffE < ETOT_CONV_THR
        CONVERGED = CONVERGED + 1
    else # reset CONVERGED
        CONVERGED = 0
    end
    if CONVERGED >= 2
        if verbose
            @printf("SCF is converged: iter: %d , diffE = %10.7e\n", iter, diffE)
        end
        break
    end
   Etot_old = Etot
   flush(stdout)
end
# Eigenvalues are not calculated if using CheFSI.
# We calculate them here.
if update_psi == "CheFSI"
   for ispin = 1:Nspin
   for ik = 1:Nkpt
       Ham.ik = ik
       Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        Hr = psiks[ikspin] ' * op_H( Ham, psiks[ikspin] )
        evals[:,ikspin] = eigvals(Hermitian(Hr))
   end
    end
end
```

```
Ham.electrons.ebands = evals
if verbose && print_final_ebands
   @printf("\n")
   @printf("----\n")
   @printf("Final Kohn-Sham eigenvalues:\n")
   @printf("----\n")
   @printf("\n")
   print_ebands(Ham.electrons, Ham.pw.gvecw.kpoints)
end
if verbose && print_final_energies
   @printf("\n")
   @printf("----\n")
   @printf("Final Kohn-Sham energies:\n")
   @printf("----\n")
   @printf("\n")
   println(Ham.energies, use_smearing=use_smearing)
end
if savewfc
   for ikspin = 1:Nkpt*Nspin
       wfc_file = open("WFC_ikspin_"*string(ikspin)*".data","w")
       write( wfc_file, psiks[ikspin] )
       close( wfc_file )
   end
end
return
```

## 12.2 Eigensolver

end

Davidson, LOBPCG, and PCG

### 12.3 Direct energy minimization via nonlinear conjugate gradient method

```
KS_solve_Emin_PCG
This method is described by IsmailBeigi-Arias
11 11 11
Solves Kohn-Sham problem using direct energy minimization as described
by Ismail-Beigi and Arias.
function KS_solve_Emin_PCG!( Ham::Hamiltonian;
                               \verb|startingwfc=| nothing, savewfc=| false, \\
                               _t=3e-5, NiterMax=200, verbose=true,
                               I_CG_BETA=2, ETOT_CONV_THR=1e-6 )
    pw = Ham.pw
    electrons = Ham.electrons
    Focc = electrons.Focc
    Nstates = electrons.Nstates
    Ns = pw.Ns
    Npoints = prod(Ns)
    CellVolume = pw.CellVolume
    Ngw = pw.gvecw.Ngw
    Ngwx = pw.gvecw.Ngwx
    Nkpt = pw.gvecw.kpoints.Nkpt
    Nspin = electrons.Nspin
```

```
Nkspin = Nkpt*Nspin
# Initial wave function
if startingwfc == nothing
   psiks = rand_BlochWavefunc(pw, electrons)
    psiks = startingwfc
end
# Calculated electron density from this wave function and
# update Hamiltonian (calculate Hartree and XC potential).
Rhoe = zeros(Float64, Npoints, Nspin)
for ispin = 1:Nspin
    idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
    Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
end
update! (Ham, Rhoe)
# Variables for PCG
g = Array{Array{ComplexF64,2},1}(undef,Nkspin)
d = Array{Array{ComplexF64,2},1}(undef,Nkspin)
g_old = Array{Array{ComplexF64,2},1}(undef,Nkspin)
d_old = Array{Array{ComplexF64,2},1}(undef,Nkspin)
Kg = Array{Array{ComplexF64,2},1}(undef,Nkspin)
Kg_old = Array{Array{ComplexF64,2},1}(undef,Nkspin)
psic = Array{Array{ComplexF64,2},1}(undef,Nkspin)
gt = Array{Array{ComplexF64,2},1}(undef,Nkspin)
for ispin = 1:Nspin
for ik = 1:Nkpt
    ikspin = ik + (ispin - 1)*Nkpt
    g[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
    d[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
    g_old[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
    d_old[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
    Kg[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
    Kg_old[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
    psic[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
    gt[ikspin] = zeros(ComplexF64, Ngw[ik], Nstates)
end
end
 = zeros(Nkspin)
 = zeros(Nkspin)
Etot_old = 0.0
# calculate E_NN
Ham.energies.NN = calc_E_NN( Ham.atoms )
# calculate PspCore energy
Ham.energies.PspCore = calc_PspCore_ene( Ham.atoms, Ham.pspots, CellVolume )
# Calculate energy at this psi
energies = calc_energies(Ham, psiks)
Ham.energies = energies
```

```
Etot = sum(energies)
CONVERGED = 0
if verbose
    @printf("\n")
    @printf("Minimizing Kohn-Sham energy using PCG\n")
    @printf("------
    Qprintf("NiterMax = %d\n", NiterMax)
    @printf("_t
                    = \%e\n", _t)
    @printf("conv_trh = %e\n", ETOT_CONV_THR)
    if I_CG_BETA == 1
        @printf("Using Fletcher-Reeves formula for CG_BETA\n")
    elseif I_CG_BETA == 2
        @printf("Using Polak-Ribiere formula for CG_BETA\n")
    elseif I_CG_BETA == 3
        @printf("Using Hestenes-Stiefeld formula for CG_BETA\n")
    else
        @printf("Using Dai-Yuan formula for CG_BETA\n")
    end
    @printf("\n")
end
for iter = 1:NiterMax
    for ispin = 1:Nspin
    for ik = 1:Nkpt
        Ham.ik = ik
        Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        g[ikspin] = calc_grad( Ham, psiks[ikspin] )
        Kg[ikspin] = Kprec( Ham.ik, pw, g[ikspin] )
        # XXX: define function trace for real(sum(conj(...)))
        if iter != 1
            if I_CG_BETA == 1
                [ikspin] =
real(sum(conj(g[ikspin]).*Kg[ikspin]))/real(sum(conj(g_old[ikspin]).*Kg_old[ikspin]))
            elseif I_CG_BETA == 2
                [ikspin] =
                real(sum(conj(g[ik-
spin]-g_old[ikspin]).*Kg[ikspin]))/real(sum(conj(g_old[ikspin]).*Kg_old[ikspin]))
            elseif I_CG_BETA == 3
                [ikspin] =
                real(sum(conj(g[ik-
spin]-g_old[ikspin]).*Kg[ikspin]))/real(sum(conj(g[ikspin]-g_old[ikspin]).*d[ikspin]))
            else
                [ikspin] =
                real(sum(conj(g[ikspin]).*Kg[ikspin]))/real(sum((g[ik-
spin]-g_old[ikspin]).*conj(d_old[ikspin])))
            end
        end
        if [ikspin] < 0.0
            [ikspin] = 0.0
        end
        d[ikspin] = -Kg[ikspin] + [ikspin] * d_old[ikspin]
        psic[ikspin] = ortho_sqrt(psiks[ikspin] + _t*d[ikspin])
```

```
end # ik
end # ispin
for ispin = 1:Nspin
    idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
    Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
end
update!(Ham, Rhoe)
for ispin = 1:Nspin
for ik = 1:Nkpt
    Ham.ik = ik
    Ham.ispin = ispin
    ikspin = ik + (ispin - 1)*Nkpt
    gt[ikspin] = calc_grad(Ham, psic[ikspin])
    denum = real(sum(conj(g[ikspin]-gt[ikspin]).*d[ikspin]))
    if denum != 0.0
        [ikspin] = abs( _t*real(sum(conj(g[ikspin]).*d[ikspin]))/denum )
        [ikspin] = 0.0
    end
    # Update wavefunction
    psiks[ikspin] = psiks[ikspin] + [ikspin]*d[ikspin]
    \# Update potentials
    psiks[ikspin] = ortho_sqrt(psiks[ikspin])
end
end
for ispin = 1:Nspin
    idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
    Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
end
update!(Ham, Rhoe)
Ham.energies = calc_energies( Ham, psiks )
Etot = sum(Ham.energies)
diffE = abs(Etot-Etot_old)
if verbose
    Qprintf("CG step %8d = %18.10f %10.7e\n", iter, Etot, diffE)
end
if diffE < ETOT_CONV_THR
    CONVERGED = CONVERGED + 1
else
    CONVERGED = 0
end
if CONVERGED >= 2
    if verbose
        @printf("CONVERGENCE ACHIEVED\n")
    end
    break
end
g_old = copy(g)
d_old = copy(d)
Kg\_old = copy(Kg)
```

```
Etot_old = Etot
        flush(stdout)
    end
    # Calculate eigenvalues
    for ispin = 1:Nspin
    for ik = 1:Nkpt
        Ham.ik = ik
        Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        psiks[ikspin] = ortho_sqrt(psiks[ikspin])
        Hr = psiks[ikspin] ' * op_H( Ham, psiks[ikspin] )
        evals, evecs = eigen(Hr)
        evals = real(evals[:])
        # We need to sort this
        idx_sorted = sortperm(evals)
        Ham.electrons.ebands[:,ikspin] = evals[idx_sorted]
        psiks[ikspin] = psiks[ikspin]*evecs[:,idx_sorted]
    end
    end
    if savewfc
        for ikspin = 1:Nkpt*Nspin
            wfc_file = open("WFC_ikspin_"*string(ikspin)*".data","w")
            write( wfc_file, psiks[ikspin] )
            close( wfc_file )
        end
    end
    return
end
```

### 12.4 Direct constrained minimization method of Yang

```
KS_solve_DCM
Solves Kohn-Sham problem using direct constrined minimization (DCM) as described
by Yang.
function KS_solve_DCM!( Ham::Hamiltonian;
                        NiterMax = 100, startingwfc=nothing,
                        savewfc=false, ETOT_CONV_THR=1e-6 )
        pw = Ham.pw
    Ngw = pw.gvecw.Ngw
    Ns = pw.Ns
    Npoints = prod(Ns)
    CellVolume = pw.CellVolume
    V = CellVolume/Npoints
    electrons = Ham.electrons
    Focc = electrons.Focc
    Nocc = electrons.Nstates_occ
    Nstates = electrons.Nstates
    Nkpt = Ham.pw.gvecw.kpoints.Nkpt
    Nspin = electrons.Nspin
    Nkspin = Nkpt*Nspin
```

```
psiks = Array{Array{ComplexF64,2},1}(undef,Nkspin)
# Initial wave function
if startingwfc == nothing
   psiks = rand_BlochWavefunc(pw, electrons)
    psiks = startingwfc
end
# Calculated electron density from this wave function and update Hamiltonian
Rhoe = zeros(Float64, Npoints, Nspin)
for ispin = 1:Nspin
    idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
    Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
update! (Ham, Rhoe)
Rhoe_old = copy(Rhoe)
evals = zeros(Float64, Nstates, Nkspin)
# calculate E_NN
Ham.energies.NN = calc_E_NN( Ham.atoms )
# calculate PspCore energy
Ham.energies.PspCore = calc_PspCore_ene( Ham.atoms, Ham.pspots, CellVolume)
# Starting eigenvalues and psi
for ispin = 1:Nspin
for ik = 1:Nkpt
   Ham.ik = ik
   Ham.ispin = ispin
    ikspin = ik + (ispin - 1)*Nkpt
    evals[:,ikspin], psiks[ikspin] =
    diag_LOBPCG( Ham, psiks[ikspin], verbose_last=false, NiterMax=10 )
end
end
Ham.energies = calc_energies( Ham, psiks )
Etot_old = sum(Ham.energies)
# subspace
Y = Array{Array{ComplexF64,2},1}(undef,Nkspin)
R = Array{Array{ComplexF64,2},1}(undef,Nkspin)
P = Array{Array{ComplexF64,2},1}(undef,Nkspin)
G = Array{Array{ComplexF64,2},1}(undef,Nkspin)
T = Array{Array{Float64,2},1}(undef,Nkspin)
B = Array{Array{Float64,2},1}(undef,Nkspin)
A = Array{Array{Float64,2},1}(undef,Nkspin)
C = Array{Array{Float64,2},1}(undef,Nkspin)
for ispin = 1:Nspin
for ik = 1:Nkpt
    ikspin = ik + (ispin - 1)*Nkpt
    Y[ikspin] = zeros( ComplexF64, Ngw[ik], 3*Nstates )
    R[ikspin] = zeros( ComplexF64, Ngw[ik], Nstates )
    P[ikspin] = zeros( ComplexF64, Ngw[ik], Nstates )
    G[ikspin] = zeros( ComplexF64, 3*Nstates, 3*Nstates )
    T[ikspin] = zeros( Float64, 3*Nstates, 3*Nstates )
```

```
B[ikspin] = zeros( Float64, 3*Nstates, 3*Nstates )
    A[ikspin] = zeros(Float64, 3*Nstates, 3*Nstates)
    C[ikspin] = zeros( Float64, 3*Nstates, 3*Nstates )
end
end
D = zeros(Float64,3*Nstates,Nkspin) # array for saving eigenvalues of subspace problem
set1 = 1:Nstates
set2 = Nstates+1:2*Nstates
set3 = 2*Nstates+1:3*Nstates
set4 = Nstates+1:3*Nstates
set5 = 1:2*Nstates
MaxInnerSCF = 3
for iter = 1:NiterMax
    for ispin = 1:Nspin
    for ik = 1:Nkpt
        Ham.ik = ik
        Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        Hpsi = op_H( Ham, psiks[ikspin] )
        psiHpsi = psiks[ikspin], * Hpsi
        psiHpsi = 0.5*( psiHpsi + psiHpsi' )
        # Calculate residual
        R[ikspin] = Hpsi - psiks[ikspin]*psiHpsi
        R[ikspin] = Kprec( ik, pw, R[ikspin] )
        # Construct subspace
        Y[ikspin][:,set1] = psiks[ikspin]
        Y[ikspin][:,set2] = R[ikspin]
        #
        if iter > 1
            Y[ikspin][:,set3] = P[ikspin]
        end
        # Project kinetic and ionic potential
        if iter > 1
            KY = op_K( Ham, Y[ikspin] ) + op_V_Ps_loc( Ham, Y[ikspin] )
            T[ikspin] = real(Y[ikspin]'*KY)
            B[ikspin] = real(Y[ikspin],*Y[ikspin])
            B[ikspin] = 0.5*(B[ikspin] + B[ikspin],)
        else
            \# only set5=1:2*Nstates is active for iter=1
            KY = op_K( Ham, Y[ikspin][:,set5] ) + op_V_Ps_loc( Ham, Y[ikspin][:,set5] )
            T[ikspin][set5,set5] = real(Y[ikspin][:,set5]'*KY)
            bb = real(Y[ikspin][set5,set5],*Y[ikspin][set5,set5])
            B[ikspin][set5, set5] = 0.5*(bb + bb')
        end
        if iter > 1
            G[ikspin] = Matrix(1.0I, 3*Nstates, 3*Nstates) #eye(3*Nstates)
            G[ikspin] = Matrix(1.0I, 2*Nstates, 2*Nstates)
        end
    end
    end
```

```
@printf("DCM iter: %3d\n", iter)
for iterscf = 1:MaxInnerSCF
   for ispin = 1:Nspin
   for ik = 1:Nkpt
        Ham.ik = ik
        Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        V_loc = Ham.potentials.Hartree + Ham.potentials.XC[:,ispin]
        if iter > 1
            yy = Y[ikspin]
        else
            yy = Y[ikspin][:,set5]
        end
        #
        if Ham.pspotNL.NbetaNL > 0
            VY = op_V_Ps_nloc( Ham, yy ) + op_V_loc( ik, pw, V_loc, yy )
        else
            VY = op_V_loc( ik, pw, V_loc, yy )
        end
        if iter > 1
            A[ikspin] = real( T[ikspin] + yy'*VY )
            A[ikspin] = 0.5*( A[ikspin] + A[ikspin], )
            aa = real( T[ikspin] [set5,set5] + yy'*VY )
            A[ikspin] = 0.5*(aa + aa')
        end
        if iter > 1
            BG = B[ikspin]*G[ikspin][:,1:Nocc]
            C[ikspin] = real( BG*BG' )
            C[ikspin] = 0.5*( C[ikspin] + C[ikspin], )
        else
            BG = B[ikspin][set5,set5]*G[ikspin][set5,1:Nocc]
            cc = real( BG*BG' )
            C[ikspin][set5, set5] = 0.5*(cc + cc')
        end
        if iter > 1
            D[:,ikspin], G[ikspin] = eigen( A[ikspin], B[ikspin] )
        else
            D[set5,ikspin], G[ikspin][set5,set5] =
            eigen( A[ikspin][set5,set5], B[ikspin][set5,set5] )
        end
        # update wavefunction
        if iter > 1
            psiks[ikspin] = Y[ikspin]*G[ikspin][:,set1]
            ortho_sqrt!(psiks[ikspin]) # is this necessary ?
        else
            psiks[ikspin] = Y[ikspin][:,set5]*G[ikspin][set5,set1]
            ortho_sqrt!(psiks[ikspin])
        end
    end
   end
    for ispin = 1:Nspin
        idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
```

end

end

```
Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
            update! ( Ham, Rhoe )
            Rhoe_old = copy(Rhoe)
            # Calculate energies once again
            Ham.energies = calc_energies( Ham, psiks )
            Etot = sum(Ham.energies)
            diffE = -(Etot - Etot_old)
            @printf("innerSCF: %5d %18.10f %18.10e", iterscf, Etot, diffE)
            # positive value of diffE is taken as reducing
            if diffE < 0.0
                Oprintf(" : Energy is not reducing !\n")
            else
                @printf("\n")
            end
        end
        # Calculate energies once again
        Ham.energies = calc_energies( Ham, psiks )
        Etot = sum(Ham.energies)
        diffE = abs( Etot - Etot_old )
        @printf("DCM: %5d %18.10f %18.10e\n", iter, Etot, diffE)
        if abs(diffE) < ETOT_CONV_THR
            @printf("DCM is converged: iter: %d , diffE = %10.7e\n", iter, diffE)
            break
        end
        Etot_old = Etot
        # No need to update potential, it is already updated in inner SCF loop
        for ispin = 1:Nspin
        for ik = 1:Nkpt
            ikspin = ik + (ispin - 1)*Nkpt
            if iter > 1
                P[ikspin] = Y[ikspin][:,set4]*G[ikspin][set4,set1]
                P[ikspin] = Y[ikspin][:,set2]*G[ikspin][set2,set1]
            end
        end
        end
        flush(stdout)
    end # end of DCM iteration
    Ham.electrons.ebands = evals[:,:]
    if savewfc
        for ikspin = 1:Nkpt*Nspin
            wfc_file = open("WFC_ikspin_"*string(ikspin)*".data","w")
            write( wfc_file, psiks[ikspin] )
            close( wfc_file )
        end
    return
function KS_solve_TRDCM!( Ham::Hamiltonian;
```

```
savewfc=false, ETOT_CONV_THR=1e-6 )
   pw = Ham.pw
Ngw = pw.gvecw.Ngw
Ns = pw.Ns
Npoints = prod(Ns)
CellVolume = pw.CellVolume
V = CellVolume/Npoints
electrons = Ham.electrons
Focc = electrons.Focc
Nstates = electrons.Nstates
Nocc = electrons.Nstates_occ
Nkpt = Ham.pw.gvecw.kpoints.Nkpt
Nspin = electrons.Nspin
Nkspin = Nkpt*Nspin
psiks = Array{Array{ComplexF64,2},1}(undef,Nkspin)
# Initial wave function
if startingwfc == nothing
   psiks = rand_BlochWavefunc(pw, electrons)
    psiks = startingwfc
end
# Calculated electron density from this wave function and update Hamiltonian
Rhoe = zeros(Float64, Npoints, Nspin)
for ispin = 1:Nspin
    idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
    Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
end
update! (Ham, Rhoe)
evals = zeros(Float64, Nstates, Nkspin)
# Starting eigenvalues and psi
for ispin = 1:Nspin
for ik = 1:Nkpt
   Ham.ik = ik
    Ham.ispin = ispin
    ikspin = ik + (ispin - 1)*Nkpt
    evals[:,ikspin], psiks[ikspin] =
    diag_LOBPCG( Ham, psiks[ikspin], verbose_last=true, NiterMax=100 )
end
end
# calculate E_NN
Ham.energies.NN = calc_E_NN( Ham.atoms )
# calculate PspCore energy
Ham.energies.PspCore = calc_PspCore_ene( Ham.atoms, Ham.pspots, CellVolume )
Ham.energies = calc_energies( Ham, psiks )
Etot = sum(Ham.energies)
Etot_old = Etot
```

NiterMax = 100, startingwfc=nothing,

```
# subspace
Y = Array{Array{ComplexF64,2},1}(undef,Nkspin)
R = Array{Array{ComplexF64,2},1}(undef,Nkspin)
P = Array{Array{ComplexF64,2},1}(undef,Nkspin)
G = Array{Array{ComplexF64,2},1}(undef,Nkspin)
T = Array{Array{Float64,2},1}(undef,Nkspin)
B = Array{Array{Float64,2},1}(undef,Nkspin)
A = Array{Array{Float64,2},1}(undef,Nkspin)
C = Array{Array{Float64,2},1}(undef,Nkspin)
for ispin = 1:Nspin
for ik = 1:Nkpt
    ikspin = ik + (ispin - 1)*Nkpt
    Y[ikspin] = zeros( ComplexF64, Ngw[ik], 3*Nstates )
    R[ikspin] = zeros( ComplexF64, Ngw[ik], Nstates )
    P[ikspin] = zeros( ComplexF64, Ngw[ik], Nstates )
    G[ikspin] = zeros( ComplexF64, 3*Nstates, 3*Nstates )
    T[ikspin] = zeros( Float64, 3*Nstates, 3*Nstates )
    B[ikspin] = zeros( Float64, 3*Nstates, 3*Nstates )
    A[ikspin] = zeros(Float64, 3*Nstates, 3*Nstates)
    C[ikspin] = zeros( Float64, 3*Nstates, 3*Nstates )
end
end
D = zeros(Float64,3*Nstates,Nkspin) # array for saving eigenvalues of subspace problem
#XXX use plain 3d-array for G, T, and B ?
set1 = 1:Nstates
set2 = Nstates+1:2*Nstates
set3 = 2*Nstates+1:3*Nstates
set4 = Nstates+1:3*Nstates
set5 = 1:2*Nstates
MaxInnerSCF = 3
MAXTRY = 10
FUDGE = 1e-12
SMALL = 1e-12
sigma = zeros(Float64,Nkspin)
gapmax = zeros(Float64,Nkspin)
CONVERGED = 0
for iter = 1:NiterMax
    for ispin = 1:Nspin
    for ik = 1:Nkpt
        Ham.ik = ik
        Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        Hpsi = op_H( Ham, psiks[ikspin] )
        psiHpsi = psiks[ikspin], * Hpsi
        psiHpsi = 0.5*( psiHpsi + psiHpsi' )
        # Calculate residual
        R[ikspin] = Hpsi - psiks[ikspin]*psiHpsi
        R[ikspin] = Kprec( ik, pw, R[ikspin] )
        # Construct subspace
        Y[ikspin][:,set1] = psiks[ikspin]
        Y[ikspin][:,set2] = R[ikspin]
```

```
if iter > 1
        Y[ikspin][:,set3] = P[ikspin]
    end
    # Project kinetic and ionic potential
    if iter > 1
        KY = op_K( Ham, Y[ikspin] ) + op_V_Ps_loc( Ham, Y[ikspin] )
        T[ikspin] = real(Y[ikspin]'*KY)
        B[ikspin] = real(Y[ikspin]'*Y[ikspin])
        B[ikspin] = 0.5*(B[ikspin] + B[ikspin],)
    else
        # only set5=1:2*Nstates is active for iter=1
        KY = op_K( Ham, Y[ikspin][:,set5] ) + op_V_Ps_loc( Ham, Y[ikspin][:,set5] )
        T[ikspin][set5,set5] = real(Y[ikspin][:,set5]'*KY)
        bb = real(Y[ikspin][set5,set5],*Y[ikspin][set5,set5])
        B[ikspin][set5, set5] = 0.5*(bb + bb')
    end
    if iter > 1
        G[ikspin] = Matrix(1.0I, 3*Nstates, 3*Nstates) #eye(3*Nstates)
        G[ikspin][set5,set5] = Matrix(1.0I, 2*Nstates, 2*Nstates)
    end
end
end
@printf("TRDCM iter: %3d\n", iter)
sigma[:] .= 0.0 # reset sigma to zero at the beginning of inner SCF iteration
numtry = 0
Etot0 = sum(Ham.energies)
println("Etot0 = ", Etot0)
for iterscf = 1:MaxInnerSCF
    for ispin = 1:Nspin
    for ik = 1:Nkpt
        Ham.ik = ik
        Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        # Project Hartree, XC potential, and nonlocal pspot if any
        V_loc = Ham.potentials.Hartree + Ham.potentials.XC[:,ispin]
        if iter > 1
            yy = Y[ikspin]
            yy = Y[ikspin][:,set5]
        end
        if Ham.pspotNL.NbetaNL > 0
            VY = op_V_Ps_nloc( Ham, yy ) + op_V_loc( ik, pw, V_loc, yy )
            VY = op_V_{loc}(ik, pw, V_{loc}, yy)
        end
        if iter > 1
            A[ikspin] = real( T[ikspin] + yy'*VY )
```

```
A[ikspin] = 0.5*( A[ikspin] + A[ikspin], )
            else
                 aa = real( T[ikspin][set5,set5] + yy'*VY )
                A[ikspin] = 0.5*(aa + aa')
            end
             #
            if iter > 1
                BG = B[ikspin]*G[ikspin][:,1:Nocc]
                C[ikspin] = real( BG*BG' )
                C[ikspin] = 0.5*( C[ikspin] + C[ikspin], )
            else
                BG = B[ikspin][set5,set5]*G[ikspin][set5,1:Nocc]
                 cc = real( BG*BG' )
                 C[ikspin][set5, set5] = 0.5*(cc + cc')
            end
             # apply trust region if necessary
            if abs(sigma[ikspin]) > SMALL # sigma is not zero
                println("Trust region is imposed")
                if iter > 1
                    D[:,ikspin], G[ikspin] =
                     eigen( A[ikspin] - sigma[ikspin] *C[ikspin], B[ikspin] )
                 else
                    D[set5,ikspin], G[ikspin][set5,set5] =
                     eigen( A[ikspin][set5,set5] - sigma[ikspin]*C[ikspin][set5,set5],
B[ikspin][set5,set5] )
                 end
            else
                 if iter > 1
                     D[:,ikspin], G[ikspin] = eigen( A[ikspin], B[ikspin] )
                 else
                     D[set5,ikspin], G[ikspin][set5,set5] = eigen( A[ikspin][set5,set5],
B[ikspin][set5,set5] )
                 end
            end
            evals[:,ikspin] = D[1:Nstates,ikspin] .+ sigma[ikspin]
             # update wavefunction
            if iter > 1
                psiks[ikspin] = Y[ikspin]*G[ikspin][:,set1]
                ortho_sqrt!(psiks[ikspin]) # is this necessary ?
            else
                psiks[ikspin] = Y[ikspin][:,set5]*G[ikspin][set5,set1]
                 ortho_sqrt!(psiks[ikspin])
            end
        end
        end
        for ispin = 1:Nspin
            idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
            Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
        end
        update! ( Ham, Rhoe )
        # Calculate energies once again
        Ham.energies = calc_energies( Ham, psiks )
        Etot = sum(Ham.energies)
        println("Etot = ", Etot)
        if Etot > Etot0
```

```
@printf("TRDCM: %f > %f: Trust region will be imposed\n", Etot, Etot0)
             # Total energy is increased, impose trust region
             # Do this for all kspin
             for ikspin = 1:Nkspin
                 if iter == 1
                     gaps = D[2:2*Nstates,ikspin] - D[1:2*Nstates-1,ikspin]
                     gapmax[ikspin] = maximum(gaps)
                 else
                     gaps = D[2:3*Nstates] - D[1:3*Nstates-1]
                     gapmax[ikspin] = maximum(gaps)
                 end
                 gap0 = D[Nocc+1,ikspin] - D[Nocc,ikspin]
                 while (gap0 < 0.9*gapmax[ikspin]) & (numtry < MAXTRY)
                     println("Increase sigma to fix gap0:")
                     @printf("gap0 : %f < %f\n", gap0, 0.9*gapmax[ikspin])</pre>
                     if abs(sigma[ikspin]) < SMALL # approx for sigma == 0.0
                          # initial value for sigma
                          sigma[ikspin] = 2*gapmax[ikspin]
                     else
                          sigma[ikspin] = 2*sigma[ikspin]
                     @printf("fix gap0: ikspin = %d, sigma = %f\n", ikspin, sigma[ikspin])
                     if iter > 1
                          D[:,ikspin], G[ikspin] = eigen( A[ikspin] - sigma[ikspin]*C[ikspin],
B[ikspin] )
                          gaps = D[2:2*Nstates,ikspin] - D[1:2*Nstates-1,ikspin]
                     else
                          D[set5,ikspin], G[ikspin][set5,set5] =
                          eigen( A[ikspin][set5,set5] - sigma[ikspin]*C[ikspin][set5,set5],
B[ikspin][set5,set5] )
                          gaps = D[2:3*Nstates] - D[1:3*Nstates-1]
                     gapmax[ikspin] = maximum(gaps)
                     gap0 = D[Nocc+1,ikspin] - D[Nocc,ikspin]
                 end
                 numtry = numtry + 1
             end # Nkspin
         end # if Etot > Etot0
         println("sigma = ", sigma)
         while (Etot > Etot0) &
               \#(abs(Etot-Etot0) > FUDGE*abs(Etot0)) &
               (numtry < MAXTRY)</pre>
             @printf("Increase sigma part 2: %f > %f ?\n", Etot, Etot0)
             # update wavefunction
             for ikspin = 1:Nkspin
                 if iter > 1
                     psiks[ikspin] = Y[ikspin]*G[ikspin][:,set1]
                     ortho_gram_schmidt!(psiks[ikspin])
                 else
                     psiks[ikspin] = Y[ikspin][:,set5]*G[ikspin][set5,set1]
                     ortho_gram_schmidt!(psiks[ikspin])
                 end
             end
```

```
for ispin = 1:Nspin
                idxset = (Nkpt*(ispin-1)+1):(Nkpt*ispin)
                Rhoe[:,ispin] = calc_rhoe( pw, Focc[:,idxset], psiks[idxset] )
            end
            update! ( Ham, Rhoe )
            # Calculate energies once again
            Ham.energies = calc_energies( Ham, psiks )
            Etot = sum(Ham.energies)
            if Etot > Etot0
                println("Increase sigma part 2")
                for ikspin = 1:Nkspin
                    if abs(sigma[ikspin]) > SMALL # sigma is not 0
                        sigma[ikspin] = 2*sigma[ikspin]
                    else
                        sigma[ikspin] = 1.2*gapmax[ikspin]
                    @printf("ikspin = %d sigma = %f\n", ikspin, sigma[ikspin])
                        D[:,ikspin], G[ikspin] = eigen( A[ikspin] - sigma[ikspin]*C[ikspin],
B[ikspin] )
                    else
                        D[set5,ikspin], G[ikspin][set5,set5] = eigen(A[ikspin][set5,set5] -
sigma[ikspin]*C[ikspin][set5,set5], B[ikspin][set5,set5] )
                    end
                end
            end
            numtry = numtry + 1 # outside ikspin loop
        end # while
        Etot0 = Etot
    end # end of inner SCF iteration
    # Calculate energies once again
    Ham.energies = calc_energies( Ham, psiks )
    Etot = sum(Ham.energies)
    diffE = abs( Etot - Etot_old )
    @printf("TRDCM: %5d %18.10f %18.10e\n", iter, Etot, diffE)
    if diffE < ETOT_CONV_THR
        CONVERGED = CONVERGED + 1
    else # reset CONVERGED
        CONVERGED = 0
    end
    if CONVERGED >= 2
        @printf("TRDCM is converged: iter: %d , diffE = %10.7e\n", iter, diffE)
        break
    end
    Etot_old = Etot
    # No need to update potential, it is already updated in inner SCF loop
    for ispin = 1:Nspin
    for ik = 1:Nkpt
        ikspin = ik + (ispin - 1)*Nkpt
        if iter > 1
            P[ikspin] = Y[ikspin][:,set4]*G[ikspin][set4,set1]
        else
            P[ikspin] = Y[ikspin][:,set2]*G[ikspin][set2,set1]
```

```
end
end
end
flush(stdout)
end # end of DCM iteration

Ham.electrons.ebands = evals[:,:]

if savewfc
   for ikspin = 1:Nkpt*Nspin
        wfc_file = open("WFC_ikspin_"*string(ikspin)*".data","w")
        write( wfc_file, psiks[ikspin] )
        close( wfc_file )
        end
end
return
end
```

## 12.5 Chebyshev filtered subspace iteration SCF

```
update_psi="CheFSI"
function diag_CheFSI! ( Ham::Hamiltonian, psiks::BlochWavefunc, cheby_degree::Int64)
    Nspin = Ham.electrons.Nspin
    Nkpt = Ham.pw.gvecw.kpoints.Nkpt
    Nstates = Ham.electrons.Nstates
    for ispin = 1:Nspin
    for ik = 1:Nkpt
        Ham.ik = ik
        Ham.ispin = ispin
        ikspin = ik + (ispin - 1)*Nkpt
        ub, lb = get_ub_lb_lanczos( Ham, Nstates*2 )
        psiks[ikspin] = chebyfilt( Ham, psiks[ikspin], cheby_degree, lb, ub)
        psiks[ikspin] = ortho_sqrt( psiks[ik] )
    end
    end
end
function chebyfilt( Ham::Hamiltonian, X, degree, lb, ub)
    Ngw_ik = size(X)[1]
    Nstates = size(X)[2]
    ee = (ub - 1b)/2
    c = (ub + 1b)/2
    sigma = ee/(lb-ub)
    sigma1 = sigma
    Y = zeros(ComplexF64, Ngw_ik, Nstates)
    Y1 = zeros(ComplexF64,Ngw_ik,Nstates)
    Y = op_H(Ham, X) - X*c
    Y = Y*sigma1/ee
    #
    for i = 2:degree
        sigma2 = 1/(2/sigma1 - sigma)
        Y1 = (op_H(Ham, Y) - Y*c)*2 * sigma2/ee - X*(sigma*sigma2)
        X = Y
        Y = Y1
        sigma = sigma2
```

```
end
    return Y
end
function get_ub_lb_lanczos( Ham::Hamiltonian, nlancz::Int64 )
    pw = Ham.pw
    ik = Ham.ik
    Ngw_ik = pw.gvecw.Ngw[ik]
    V = zeros(ComplexF64,Ngw_ik,nlancz)
    HV = zeros(ComplexF64,Ngw_ik,nlancz)
    T = zeros(Float64,nlancz,nlancz)
    f = zeros(ComplexF64,Ngw_ik)
    s = zeros(ComplexF64,nlancz)
    h = zeros(ComplexF64,nlancz)
    V[:,1] = randn(Ngw_ik) + im*randn(Ngw_ik)
    beta = norm(V[:,1])
    V[:,1] = V[:,1] ./ beta
    HV[:,1] = op_H(Ham, V[:,1])
   h[1] = real( V[:,1], * HV[:,1] )
    T[1,1] = h[1]
    # One-step of reorthogonalization
    f[:] = HV[:,1] - V[:,1]*h[1]
    s[1] = V[:,1], * f[:]
    h[1] = h[1] + s[1]
    f[:] = f[:] - V[:,1]*s[1]
    # MAIN LOOP
    for j = 2:nlancz
        \#Oprintf("iter lanczos = %d \ n", j)
        beta = norm(f)
        T[j,j-1] = beta
        V[:,j] = f[:]/beta
        HV[:,j] = op_H(Ham, V[:,j])
        for jj = 1:j
            h[jj] = V[:,jj], * HV[:,j]
        end
        f[:] = HV[:,j] - V[:,1:j]*h[1:j]
        # One-step of reorthogonalization
        \#s = V' * f
        #h = h + s
        for jj=1:j
            s[jj] = V[:,jj], * f[:]
            h[jj] = h[jj] + s[jj]
        f[:] = f[:] - V[:,1:j]*s[1:j]
        T[1:j,j] = real(h[1:j])
    end
    evalsT = eigvals(T)
    \#lb = evalsT[Nstates+2]
    #ub = evalsT[2*Nstates]
    lb = evalsT[Int64(nlancz/2)]
    ub = norm_matrix_induced(T) + norm(f)
    return lb, ub
```

end

```
function norm_matrix_induced(A::Array{Float64,2})
   N = size(A)[1]
   # FIXME no check for matrix form

# unit-norm vector
   d = 1/sqrt(N)
   v1 = ones(N)*d
   #
   v = A*v1
   return norm(v)
end
```