A detailed description of the application of the finite difference method to the electronic structure has been given by Laaksonen et al in the case of an Hartree-Fock approach[1].

1 Method

1.1 Conjugate gradient method

The solution of a linear systems of equation A|x>=-|b>, where A is symmetric and positive definite, is equivalent to minimizing the quadratic function

$$f(|x>) = \frac{1}{2} < x|A|x> + < b|x> + c, \tag{1}$$

with respect to the vector |x>.

2 The mesh

We deal with the problem by using finite differences on a cubic uniform grid, \mathbf{r}_i . In such ar representation, the Hamiltonian is sparse. We used second-order discretization for the Laplacian. In the framework of a finite differenc approach, the only nonvanishing matrix elements, $H_{ij} = \langle \mathbf{r}_i | \hat{H} | \mathbf{r}_j \rangle$, are on the diagonal and between neighbouring points (2 in 1D, 4 in 2D and 6 in 3D):

$$H_{ij} = \begin{cases} \frac{1}{(\Delta x)^2 + \Delta y)^2 + \Delta z)^2}, & \text{if } i = j, \\ -\frac{1}{2(\Delta)^2}, & \text{if} |\boldsymbol{r}_i - \boldsymbol{r}_j| = \Delta, \\ 0, & \text{otherwise,} \end{cases}$$
 (2)

where Δx , Δy , et Δz are the spacing between grid points.

Using Hartree atomic units, the Schrödinger we want to solve is

$$-\frac{1}{2}\left(\frac{\mathrm{d}^2}{\mathrm{d}x^2} + \frac{\mathrm{d}^2}{\mathrm{d}y^2} + \frac{\mathrm{d}^2}{\mathrm{d}z^2}\right)\phi(x, y, z) + V(x, y, z)\phi(x, y, z) = \epsilon\phi(x, y, z) \tag{3}$$

For solving this equation, we resort to the finite difference method, a mesh-type approach[2, 3]. It consists to discretize the space in a finite set of locations in space so that to form a mesh:

$$(x, y, z) \rightarrow (i, j, k),$$
 (4)

$$\phi(x, y, z) \rightarrow \phi_{i,j,k},$$
 (5)

(6)

with $x = i\Delta x$, $y = j\Delta y$, and $z = k\Delta z$.

Then, each node of the space obeys the equation

$$-\frac{1}{2}\phi_{i,j,k}'' + V_{i,j,k}\phi_{i,j,k} = \epsilon\phi_{i,j,k}.$$
 (7)

The principle of the finite difference method is to replace the derivatives in differential equations by approximations made up of a weighted sums of function values, derived using Taylor series expansions. The number of function values necessary to approximate a derivative at any node of the space depends on both the desired accuracy and the dimension. For example, at the lowest accurate approximation (three-point finite difference), in 3D, the second derivative of $\phi_{i,j,k}$ can be written as

$$\phi_{i,j,k}'' = \frac{\phi_{i-1,j,k} - 2\phi_{i,j,k} + \phi_{i+1,j,k}}{(\Delta x)^2} + \frac{\phi_{i,j-1,k} - 2\phi_{i,j,k} + \phi_{i,j+1,k}}{(\Delta y)^2} + \frac{\phi_{i,j,k-1} - 2\phi_{i,j,k} + \phi_{i,j,k-1}}{(\Delta z)^2}$$
(8)

For each node, the replacement of the derivatives by a sum of function value leads to a linear system of algebraic equations coupling the different values $\phi_{i,j,k}$:

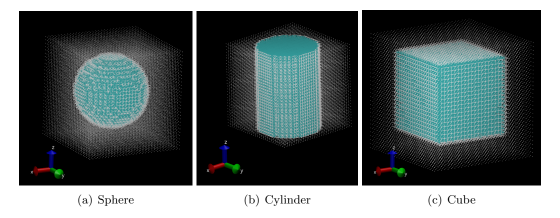


Figure 1: Definition of the "active" (blue) and "unactive" (white) domains. 3D Box $30 \times 30 \times 30$ bohr radius; $30 \times 30 \times 30$ nodes; radius of the computational domaine: 15 bohr radius.

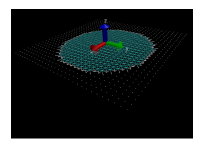


Figure 2: Definition of the "active" (blue) and "unactive" (white) domains. 2D Box 30×30 bohr radius; 30×30 nodes; radius of the computational domaine: 15 bohr radius.

$$-\frac{1}{2} \left(\frac{\phi_{i-1,j,k} - 2\phi_{i,j,k} + \phi_{i+1,j,k}}{(\Delta x)^2} + \frac{\phi_{i,j-1,k} - 2\phi_{i,j,k} + \phi_{i,j+1,k}}{(\Delta y)^2} + \frac{\phi_{i,j,k-1} - 2\phi_{i,j,k} + \phi_{i,j,k-1}}{(\Delta z)^2} \right) + V_{i,j,k}\phi_{i,j,k} = \epsilon \phi_{i,j,k}$$
(9)

In the code, all informations about the mesh are stocked the data type t_mesh (defined in global.f90):

```
type t_mesh
  integer :: Nx,Ny,Nz,N,nactive,nunactive
  integer,allocatable :: list_bound(:,:),n_bound(:) ! list_bound is linked with bound(:)
  double precision :: dx,dy,dz,dv
  type(t_box)::box
  integer :: dim
  integer::nbound
  type(t_point),allocatable::bound(:)
  type(t_ijk_to_idx),allocatable::ijk_to_idx(:,:,:) ! from (i,j,k) -> n
  type(t_node),allocatable::node(:)
end type t_mesh
```

Different kinds of parameters are necessary to define the mesh: the dimension (1D, 2D or 3D), shape of the box(sphere, cube, ...), size of the box(L_x , L_y , L_z), number of nodes (N_x , N_y , N_z), ...

The mesh spacings, Δx , Δy and Δz , are then given by $\Delta \alpha = L_{\alpha}/(N_{\alpha}+1)$.

The mesh is defined by calling the subroutine new_mesh(mesh,param) (in mesh.f90). The mesh, containing $N = N_x \times N_y \times N_z$ nodes, consists in two kind of nodes: the "active" nodes, belonging to the computational domain, and the "unactive" ones, belonging to the boundaries domain (figs. 1 and 2). The "active" nodes are the nodes where we will compute quantities such as wavefunctions, potentials, ... while the "unactive" nodes are the nodes where all these quantities are constant. Example of "unactive" nodes

is the nodes located in the outer edge of the box defining the shape of the space we consider. At these locations, the value of the wavefunction is zero in the case of isolated systems. The type of any node of the space is determined by the subroutine set_nodes(mesh):

• if the node (i, j, k) belongs to the computational domain, an single positive index n is assigned to the node; each index n corresponds to an single set (i, j, k) of the space.

$$(x, y, z) \rightarrow (i, j, k) \rightarrow n,$$
 (10)

$$\phi(x, y, z) \rightarrow \phi_{i,j,k} \rightarrow \phi_n,$$
 (11)

The total number of nodes inside the computational domain is n_{active} .

• if the node (i, j, k) belongs to the boundaries domain, its index n is set to -1.

The computational and boundaries domains are defined thanks to parameters decribing the shape and the size of the computational box (mesh%box%shape, mesh%box%center, mesh%box%radius)

From equations (9) and by using the application $(x, y, z) \to (i, j, k) \to n$, solving the Schrödinger equation (3) is equivalent to solve an eigenvalues problem $H|\phi>=\epsilon|\phi>$; the vector $|\phi>$ is an eigenvector to be determined, corresponding to the eigenvalue ϵ .

For each node (i, j, k), the list of neighbors $((i \pm 1, j, k), (i, j \pm 1, k), (i, j, k \pm 1))$ in the case given here) involved in equations (9) is given by the matrix mesh%node(nactive)%list_neighbors(:) (mesh%node(nactive)%n_neighbors contains the number of neighbors for each node (i, j, k)). These lists are set in the subroutine compute_list_neighbors(mesh).

3 Electrostatic potential

There two ways to compute the electrostatic potential of the charge distribution, $\rho(\mathbf{r})$, of the electron in the system:

- by solving the Poisson equation, $\nabla U(\mathbf{r}) = -4\pi \rho(\mathbf{r})$ (Hartree a. u.),
- by solving the integral equation, $U(\mathbf{r}) = \int d\mathbf{r}' \rho(\mathbf{r}') \frac{1}{|\mathbf{r} \mathbf{r}'|}$.

Solving the integral equation is the simplest way to proceed but it is also the most inefficient because it needs to integrate the equation for each point belonging to the grid.

Using the Poisson equation is a more efficient way to proceed. In the framework of the finite difference method, in three dimensions (3D), the Poisson equation has the form

$$\frac{U_{i-1,j,k} - 2U_{i,j,k} + U_{i+1,j,k}}{(\Delta x)^2} + \frac{U_{i,j-1,k} - 2U_{i,j,k} + U_{i,j+1,k}}{(\Delta y)^2} + \frac{U_{i,j,k-1} - 2U_{i,j,k} + U_{i,j,k+1}}{(\Delta z)^2} = -4\pi\rho_{i,j,k}. \quad (12)$$

For each active node, the replacement of the second derivative of the electrostatic potential by a sum of functions leads to a linear systems of algebrais equations coupling the different value $U_{i,j,k}$,

$$L|U\rangle = -4\pi|\rho\rangle. \tag{13}$$

L is the Laplace matrix.

Note that for the active nodes located at the edge of the active domain, the replacement of the second derivative by a sum of function needs to take into account the unactive nodes located at the edge of the unactive. For example, for the node (i, j, k),

$$\frac{U_{i-1,j,k}^* - 2U_{i,j,k} + U_{i+2,j,k}}{(\Delta x)^2} + \frac{U_{i,j-1,k} - 2U_{i,j,k} + U_{i,j+1,k}}{(\Delta y)^2} + \frac{U_{i,j,k-1} - 2U_{i,j,k} + U_{i,j,k+1}}{(\Delta z)^2} = -4\pi \rho_{i,j,k} \quad (14)$$

the node $U_{i-1,j,k}^*$ belongs to the unactive domain and imposes a boundary condition. Note that only a part of the unactive domain is involved in the boundary conditions. This part of the unactive domain is noted Ω in the following. The unactive nodes involved in the boundary conditions are listed in the data type

t_node: for each active node, the integer mesh%node(:)%n_bound gives the number of neigbors unactive nodes contributing to the boundary conditions (their index is given in mesh%node(:)%list_bound(:)).

Then

$$\frac{-2U_{i,j,k} + U_{i+2,j,k}}{(\Delta x)^2} + \frac{U_{i,j-1,k} - 2U_{i,j,k} + U_{i,j+1,k}}{(\Delta y)^2} + \frac{U_{i,j,k-1} - 2U_{i,j,k} + U_{i,j,k+1}}{(\Delta z)^2} = -4\pi\rho_{i,j,k} - \frac{U_{i-1,j,k}^*}{(\Delta x)^2}$$
(15)

$$-2\left(\frac{1}{(\Delta x)^{2}} + \frac{1}{(\Delta y)^{2}} + \frac{1}{(\Delta z)^{2}}\right)U_{i,j,k} + \frac{U_{i+2,j,k}}{(\Delta x)^{2}} + \frac{U_{i,j-1,k}}{(\Delta y)^{2}} + \frac{U_{i,j+1,k}}{(\Delta y)^{2}} + \frac{U_{i,j,k-1}}{(\Delta z)^{2}} + \frac{U_{i,j,k+1}}{(\Delta z)^{2}} = -4\pi\rho_{i,j,k} - \frac{U_{i-1,j,k}^{*}}{(\Delta x)^{2}}$$
(16)

$$L|U\rangle = -4\pi|\rho\rangle - |b\rangle. \tag{17}$$

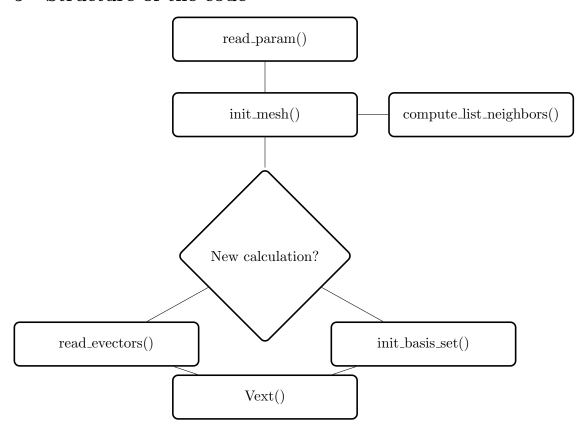
|b> is the vector containing the boundary conditions, i. e, the values of the electrostatic potential at the edge of the active domain.

Now the question is: how to get the values of the electrostatic potential at the unactive point edge of the unactive domain in

4 Structure of the input file

The input file is provided directly from the command line: ./Hbinitio.x inp_davidson for example. It must be ended by the line cmd=end to indicate the end of the file.

5 Structure of the code



6 Dealing with the perturbation

6.1 Real-space representation of the resolvent of the Hamiltonian, \hat{H}

A way to compute the electron density, n(r), is to resort to the identity relating the density matrix of a system, $\rho(r, r')$, with its one-electron Green's function:

$$G(\mathbf{r}, \mathbf{r}'; \epsilon) = \langle \mathbf{r} | \hat{G}(\epsilon) | \mathbf{r}' \rangle = \langle \mathbf{r} | \frac{1}{\epsilon - \hat{H}} | \mathbf{r}' \rangle$$
 (18)

$$n(\mathbf{r}) = \rho(\mathbf{r}, \mathbf{r})$$

Baroni and Giannozzi[4] introduced a Green's function method which employs the finite difference and a real-space grid.

Using the Green's method needs to calculate the inverse of $(\epsilon - \hat{H})$ which may be an huge task. Iterative algorithms, such as the Davidson's one, used to solve elliptic partial differential equations could in principle be used to calculate the inverse of $(\epsilon - \hat{H})$ (in the Davidson's method they are used to compyte de |delta> vector from the residual). However, for the Green's method, we don't need to get the *full* inverse matrix but only the diagonal element of the inverse.

Haydock, Heine and Kelly[5, 6] developped a convenient way to compute a *single* diagonal element of the Green's function: the recursion method. In the recursion method, diagonal elements of the Green's function, $\langle \phi_0 | \hat{G} | \phi_0 \rangle$, are expressed in terms of a continuous fraction whose coefficients are calculated from a chain of orthogonal states recursively generated from $|\phi_0\rangle$:

7 Sources

#! /bin/bash
####SBATCH -p public
#SBATCH -p grant -A g2018a7

```
###SBATCH -p grantgpu -Ag1018a7
#SBATCH -n 64 -tasks-per-node=64
#SBATCH -t 2:00:00
                       # Le job sera tue au bout de 1h
####SBATCH --mail-user=herve.bulou@ipcms.unistra.fr
#####SBATCH --mem=4096
                          # Quantite memoire demandee par noeud en Mo (unite obligat
export OMP_NUM_THREADS=64
time ./Hbinitio.x
module time_tracking
 implicit none
 type t_time
   real :: start,end,start_loc,end_loc
 end type t_time
contains
 1 -----
 subroutine time_tracking_init(time_spent)
  implicit none
  type(t_time)::time_spent
  call cpu_time(time_spent%start)
  open(unit=1,file="dbg.dat",form='formatted',status='unknown')
  write(1,*)
  close(1)
 end subroutine time_tracking_init
 subroutine time_tracking_write(iloop,time_spent,text)
  integer :: iloop
  type(t_time)::time_spent
  character (len=*) :: text
  open(unit=1,file="dbg.dat",form='formatted',status='unknown',access='append')
  write(1,'(A50,I4,F12.6,F12.6,F12.6)') text,iloop,time_spent%end_loc,&
      time_spent%start_loc,time_spent%end_loc-time_spent%start_loc
  close(1)
 end subroutine time_tracking_write
end module time_tracking
 program Hbinitio
 !$ use OMP_LIB
 use time_tracking
 implicit none
! include 'mpif.h'
 type t_GramSchmidt
   integer :: nindep
   integer :: ndep ! number of linear dependencies discovered
 end type t_GramSchmidt
 type t_mesh
```

```
integer :: Nx,Ny,Nz,N
    integer,allocatable :: list_neighbors(:,:),n_neighbors(:)
    double precision :: dx,dy,dz,dv
    double precision :: center(3)
 end type t_mesh
 type(t_mesh) :: mesh
 type t_cvg
    integer,allocatable:: list_cvg(:)
    integer :: ncvg
    double precision :: ETA
    integer :: nvec_to_cvg
 end type t_cvg
 type(t_cvg) :: cvg
 type(t_time) :: time_spent
 type t_param
    logical::restart
    integer::ieof
    integer::loopmax
    integer::nvecini
    integer::nvecmax
    integer::Nx
    integer::nvec_to_cvg
    double precision :: ETA
    double precision::box_width
 end type t_param
 type(t_param)::param
 integer :: nvec
 integer,parameter :: seed = 86456
 double precision, allocatable :: V(:,:) ! wavefunctions
 double precision,allocatable :: Sprev(:),dS(:) ! eigenvalues
 double precision,allocatable :: pot_ext(:) ! external potential
 integer :: iloop
 character (len=1024) :: filecube
 character (len=1024)::line
 integer :: i
! integer::ierr,my_id,num_procs
 call time_tracking_init(time_spent)
! call mpi_init(ierr )
! call MPI_COMM_RANK (MPI_COMM_WORLD, my_id, ierr)
! call MPI_COMM_SIZE (MPI_COMM_WORLD, num_procs, ierr)
 call read_param(param)
 call init_mesh(mesh,param)
! nvecini=2
 nvec=param%nvecini
 allocate(V(mesh%N,nvec))
! param%restart=.TRUE.
! param%restart=.FALSE.
```

```
if (.not.(param%restart))
                   then
   print *,"new calculation"
   call init_basis_set(V,nvec,seed,mesh)
 else
   print *,'restart an old calculation'
   call read_evectors(V,mesh,nvec)
 allocate(pot_ext(mesh%N))
 call Vext(mesh,pot_ext)
 open(unit=1,file="eigenvalues.dat",form='formatted',status='unknown'); write(1,*); close(1)
 iloop=1
 cvg%ncvg=0
 cvg%nvec_to_cvg=param%nvec_to_cvg
 cvg%ETA=param%ETA
 allocate(Sprev(param%nvecini))
 allocate(dS(param%nvecini))
 Sprev(:)=0.0
 dS(:)=0.0
 do while((iloop.le.param%loopmax).and.(cvg%ncvg.lt.cvg%nvec_to_cvg))
   write(*,'(A,I4,A)') 'Main > ######### scf loop=',iloop,' ###########
   call davidson(nvec, V, mesh, param%nvecini, iloop, cvg, pot_ext, time_spent)
 end do
 call save_evectors(V,mesh,param%nvecini)
 do i=1,param%nvecini
   write(filecube,'(a,i0,a)') 'evec',i,'.cube'
   call norm(mesh, V(:,i))
   call save_cube(V(:,i),filecube,mesh)
 end do
 deallocate(V)
 deallocate(Sprev)
 deallocate(dS)
 deallocate(pot_ext)
 call free_mesh(mesh)
 call cpu_time(time_spent%end)
 if (cvg%ncvg.ge.cvg%nvec_to_cvg) print *,'Main > Job DONE !'
 print '("Main > Total Time = ",e16.6," seconds.")',time_spent%end-time_spent%start
! call mpi_finalize(ierr)
 contains
          read_param()
```

```
subroutine read_param(param)
  implicit none
  type(t_param)::param
  integer::lline,eqidx
  double precision, parameter :: pi=3.1415927
  param%ieof=0
  param%loopmax=1000
  param%restart=.FALSE.
  param%nvecini=20
  param%nvecmax=41
  param%Nx=30
  param%nvec_to_cvg=20
  param%ETA=1.0e-3
  param%box_width=pi/sqrt(2.0)
  open(unit=1,file='inp',form='formatted')
  do while(.not.(is_iostat_end(param%ieof)))
    read(1,*,iostat=param%ieof) line
     lline=len_trim(line)
     eqidx=index(line,"=")
    print *,'###',eqidx,lline
     if(line(1:eqidx-1).eq."restart") then
        if(line(eqidx+1:lline).eq.'.TRUE.') then
           param%restart=.TRUE.
        else
           param%restart=.FALSE.
        end if
     end if
     if(line(1:eqidx-1).eq."loopmax") then
        read(line(eqidx+1:lline),*) param%loopmax
     end if
     if(line(1:eqidx-1).eq."nvecini") then
        read(line(eqidx+1:lline),*) param%nvecini
     end if
     if(line(1:eqidx-1).eq."nvecmax") then
        read(line(eqidx+1:lline),*) param%nvecmax
     if(line(1:eqidx-1).eq."Nx") then
        read(line(eqidx+1:lline),*) param%nx
     if(line(1:eqidx-1).eq."ETA") then
        read(line(eqidx+1:lline),*) param%ETA
     if(line(1:eqidx-1).eq."nvec_to_cvg") then
        read(line(eqidx+1:lline),*) param%nvec_to_cvg
     end if
     if(line(1:eqidx-1).eq."box_width") then
        read(line(eqidx+1:lline),*) param%box_width
     end if
     line=',
  end do
  close(1)
```

```
print *,'#restart=',param%restart
 print *,'#loopmax=',param%loopmax
 print *,'#nvecini=',param%nvecini
 print *,'#nvecmax=',param%nvecmax
 print *,'#ETA=',param%ETA
 print *,'#nvec_to_cvg=',param%nvec_to_cvg
 print *,'#box_width=',param%box_width
 print *,'#Nx=',param%nx
 print *,'#dh=',param%box_width/(param%Nx+1)
end subroutine read_param
                         -----
             save_evectors()
subroutine save_evectors(V,m,nvecini)
 implicit none
 type(t_mesh)::m
 double precision :: V(:,:)
 integer::nvecini,i,j
 open(unit=1,file="evectors.dat",form='formatted',status='unknown')
 do i=1,m%N
    write(1,*) (V(i,j),j=1,nvecini)
 end do
 close(1)
end subroutine save_evectors
             read_evectors()
subroutine read_evectors(V,m,nvecini)
 implicit none
 type(t_mesh)::m
 double precision :: V(:,:)
 integer::nvecini,i,j
 open(unit=1,file="evectors.dat",form='formatted',status='unknown')
 do i=1,m%N
    read(1,*) (V(i,j),j=1,nvecini)
 end do
 close(1)
end subroutine read_evectors
             norm()
! -----
subroutine norm(m, evec)
 implicit none
 double precision :: evec(:),normloc
 double precision, external :: ddot
```

```
type(t_mesh)::m
    normloc=sqrt(m%dv*ddot(m%N,evec(:),1,evec(:),1))
     call dscal(m%N,normloc,evec(:),1)
end subroutine norm
                                   Vext()
! ------
subroutine Vext(m,pot_ext)
     implicit none
    type(t_mesh) :: m
    double precision :: pot_ext(:)
    double precision :: pts(3),rsqr
     character (len=1024) :: filename
     integer :: i,j,k,nn
    do k=1,m\%Nz
           pts(3)=k*m%dz
           do i=1,m\%Nx
                  pts(1)=i*m%dx
                   do j=1,m\%Ny
                         pts(2)=j*m%dy
                         rsqr=(pts(1)-m%center(1))**2+(pts(2)-m%center(2))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2+(pts(3)-m%center(3))**2
                         nn=j+(i-1)*m\%Ny+(k-1)*m\%Ny*m\%Nx
                          pot_ext(nn)=10*rsqr
                   end do
            end do
     end do
    filename='pot_ext.cube'
     call save_cube(pot_ext,filename,m)
     !stop
end subroutine Vext
                                   DAVIDSON()
     ______
subroutine davidson(nvec, V, m, nvecini, iloop, cvg, pot_ext, time_spent)
     implicit none
     integer :: nvec,nvecini,iloop
    double precision,allocatable :: V(:,:),pot_ext(:)
     type(t_mesh) :: m
     type(t_cvg)::cvg
    type(t_time)::time_spent
    double precision, allocatable :: S(:) ! eigenvalues
    double precision, allocatable :: T(:,:) ! reduced matrix T
     double precision, allocatable :: VRitz(:,:) ! Ritz's vectors
    double precision,allocatable :: residual(:,:) ! residual
    double precision,allocatable :: delta(:,:) ! delta vectors
     double precision, allocatable :: Vnew(:,:) ! Vnew
     integer :: i
```

```
integer :: ndelta
integer :: newnvec
type(t_GramSchmidt)::GS
! T (reduced matrix) computing
allocate(T(nvec,nvec))
call cpu_time(time_spent%start_loc)
call compute_T(T,V,nvec,m,pot_ext)
call cpu_time(time_spent%end_loc)
call time_tracking_write(iloop,time_spent,'Davidson -> compute_T')
! Diagonatilzation of T
allocate(S(nvec))
call cpu_time(time_spent%start_loc)
call diagonalization(S,T,nvec)
call cpu_time(time_spent%end_loc)
call time_tracking_write(iloop,time_spent,'Davidson -> Diagonalization')
dS(:)=S(1:nvecini)-Sprev(:)
Sprev(:)=S(1:nvecini)
do i=1,nvecini
   write(*,'(A,I6,A,F12.6,A,E12.2,A)') 'Main > Eigenvalue(',i,'): ',S(i),'(',dS(i),')'
end do
!call cpu_time(inter)
open(unit=1,file="eigenvalues.dat",form='formatted',status='unknown',access='append')
write(1,*) iloop,S(1:nvecini)
close(1)
! computation of the Ritz's vectors
allocate(VRitz(m%N,nvec))
call cpu_time(time_spent%start_loc)
call Ritz(VRitz,V,T,nvec)
call cpu_time(time_spent%end_loc)
call time_tracking_write(iloop,time_spent,'Davidson -> Diagonalization')
! computation of residual
allocate(residual(m%N,nvec))
allocate(cvg%list_cvg(nvec))
cvg%list_cvg(:)=0
cvg%ncvg=0
call cpu_time(time_spent%start_loc)
call compute_residual(residual, VRitz, S, nvec, cvg, m, pot_ext)
call cpu_time(time_spent%end_loc)
call time_tracking_write(iloop,time_spent,'Davidson -> Residual')
! computation of delta
allocate(delta(m%N,nvec))
delta(:,:)=0.0
!call cpu_time(inter)
```

```
call cpu_time(time_spent%start_loc)
  call compute_delta(delta,residual,S,nvec,cvg,m,ndelta,pot_ext)
  call cpu_time(time_spent%end_loc)
  call time_tracking_write(iloop,time_spent,'Davidson -> Delta')
 deallocate(V)
  allocate(V(m%N,nvec+ndelta))
 V(:,1:nvec)=VRitz(:,:)
 print *,'ndelta=',ndelta
 print *,'nvec=',nvec
 V(:,nvec+1:nvec+ndelta)=delta(:,:ndelta)
  allocate(Vnew(m%N,nvec+ndelta))
 newnvec=nvec+ndelta
 GS%nindep=nvec
  !call cpu_time(inter)
  call GramSchmidt(Vnew, V, newnvec, m, GS)
  ! call cpu_time(inter2); call dbg(iloop,inter,inter2,'GS')
 print *,'Main > ',GS%ndep,newnvec
 deallocate(V)
  if(newnvec.le.param%nvecmax) then
    nvec=newnvec
  else
    print *,'Main > restart from nvecini'
    nvec=nvecini
  end if
  allocate(V(m%N,nvec))
 V(:,:)=Vnew(:,1:nvec)
  !call check_ortho(V,nvec,m)
 print *,'Main > New size of the basis ',nvec
  iloop=iloop+1
 deallocate(S)
 deallocate(T)
 deallocate(VRitz)
 deallocate(residual)
 deallocate(delta)
 deallocate(Vnew)
 deallocate(cvg%list_cvg)
end subroutine davidson
1 -----
              SAVE_CUBE()
subroutine save_cube(data,filename,m)
  implicit none
 double precision :: data(:)
  integer :: idxmin,idxmax
 type(t_mesh) :: m
```

```
character (len=1024) :: filename
  character(len=*),parameter :: FMT1='(I5,3F12.6)'
  integer :: i,j,k,nn,ifield
  open(unit=1,file=filename,form='formatted',status='unknown')
  write(1,*) 'Cubefile created from Hbinitio.f90 calculation'
 write(1,*) ' H. Bulou, November 2018'
 write(1,FMT1) 1,0.0,0.0,0.0
 write(1,FMT1) m%Nx,m%dx,0.0,0.0
 write(1,FMT1) m%Ny,0.0,m%dy,0.0
 write(1,FMT1) m%Nz,0.0,0.0,m%dz
 write(1,'(I5,4F12.6)') 1,1.0,0.0,0.0,0.0
  do k=1,m%Nz
    ifield=0
    do i=1,m\%Nx
       do j=1,m\%Ny
          nn=j+(i-1)*m\%Ny+(k-1)*m\%Ny*m\%Nx
          write(1,'(E13.5)',advance='no') data(nn)
          ifield=ifield+1
          if (mod(ifield,6).eq.0) then
             ifield=0
             write(1,*)
          end if
       end do
    end do
    write(1,*)
  end do
  close(1)
end subroutine save_cube
 ______
              COMPUTE_DELTA()
subroutine compute_delta(delta,r,lambda,nvec,cvg,m,ndelta,pot_ext)
  ! INPUT: the residual |r>, the Ritz's vectors | VRitz>, the eigenvalues lambda
  ! OUTPUT : the correction |delta> to improve the Ritz's vectors so that to
            minimize the residual
  implicit none
 type(t_mesh)::m
  double precision :: lambda(:),r(:,:),delta(:,:),pot_ext(:)
  integer :: nvec,ndelta
 type(t_cvg)::cvg
 double precision, external :: ddot
 double precision, parameter::alpha=1.0,beta=0.0
 double precision, allocatable :: normloc
  double precision, allocatable :: Dinv(:,:)
  integer :: i,j
  double precision :: deltasqr
```

```
deltasqr=m%dx**2
 print *,'Delta > -----'
 print *,'Delta > --- compute_delta ---'
 print *,'Delta > -----'
 !delta(:,:)=0.0
 ! allocate(Dinv(m%N,m%N))
! Dinv(:,:)=0.0
 ndelta=0
 do i=1,nvec
     if(cvg%list_cvg(i).eq.0) then
       ndelta=ndelta+1
       do j=1,m%N
          delta(j,ndelta)=r(j,ndelta)/((3.0/deltasqr+pot_ext(j))-lambda(i))
           Dinv(j,j)=1.0/((3.0/deltasqr+pot_ext(j))-lambda(i))
       ! see Victor Eijkhout in "Introduction to scientific and technical computing" edited b
       ! Chap 15 Libraries for Linear Algebra
       ! to get a comprehensive way to use dgemv
        call dgemv('N', m%N, m%N, alpha, Dinv, m%N, r(:, ndelta), 1, beta, delta(:, ndelta), 1)
       !norm=sqrt(ddot(m%N,delta(:,i),1,delta(:,i),1))
        !write(*, '(A10,I4,A2,E12.6)',advance='no') ' Delta > delta(',i,')=',norm
       !delta(:,ndelta)=delta(:,ndelta)+VRitz(:,ndelta)
       normloc=1.0/sqrt(ddot(m%N,delta(:,ndelta),1,delta(:,ndelta),1))
       call dscal(m%N,normloc,delta(:,ndelta),1)
     end if
  end do
  !deallocate(Dinv)
 print *,'Delta > ',ndelta,' new vector(s)'
end subroutine compute_delta
              COMPUTE_RESIDUAL()
subroutine compute_residual(r,VRitz,S,nvec,cvg,m,pot_ext)
  implicit none
 type(t_mesh)::m
  integer :: nvec
 double precision :: r(:,:),VRitz(:,:),S(:),pot_ext(:)
 type(t_cvg) :: cvg
  integer :: i,j,k
 double precision :: normloc
 double precision, external :: ddot
 double precision :: deltasqr
 print *,'Residual > -----'
 print *,'Residual > --- compute residual ---'
 print *,'Residual > -----'
 deltasqr=m%dx**2
 r(:,:)=0.0
```

```
cvg%ncvg=0
 do j=1,nvec
    do i=1,m%N
       r(i,j)=(3.0/deltasqr+pot_ext(i))*VRitz(i,j)
       do k=1,m%n_neighbors(i)
          r(i,j)=r(i,j)-0.5*VRitz(m%list_neighbors(i,k),j)/deltasqr
       end do
       r(i,j)=r(i,j)-S(j)*VRitz(i,j)
    end do
    normloc=ddot(m\%N,r(:,j),1,r(:,j),1)
    write(*,'(A,I4,A,E12.4,A,E12.4)',advance='no') 'Residual > r(',j,')= ',normloc,'/',cvg%ET
    if (normloc.lt.cvg%ETA) then
       cvg%ncvg=cvg%ncvg+1
       cvg%list_cvg(j)=1
       write(*,*) '--> converged'
    else
       write(*,*)
    end if
 end do
end subroutine compute_residual
             RITZ()
! ------
subroutine Ritz(Vout, Vin, y, nvec)
 implicit none
 double precision :: Vin(:,:),Vout(:,:),y(:,:)
 integer :: nvec
 integer :: i,j
 print *,'Ritz > -----'
 print *,'Ritz > --- Ritz() ---'
 print *,'Ritz > -----'
 do i=1,nvec
    Vout(:,i)=y(1,i)*Vin(:,1)
    do j=2,nvec
       Vout(:,i)=Vout(:,i)+y(j,i)*Vin(:,j)
    end do
 end do
end subroutine Ritz
             DIAGONALIZATION()
subroutine diagonalization(S,H,N)
 implicit none
 integer :: N
```

```
double precision :: H(:,:),S(:)
  integer :: lwork,info
  integer :: lwmax
  double precision,allocatable::work(:)
 parameter(lwmax=100000)
 allocate(work(lwmax))
 lwork=-1
  call dsyev('vectors', 'Upper', N, H, N, S, work, lwork, info)
  lwork=min(lwmax,int(work(1)))
  if (lwork.ge.lwmax) then
    write(*,*) 'Diagonalization > WARNING info = ',info
    write(*,*) 'Diagonalization > WARNING lwork=',lwork
    write(*,*) 'Diagonalization > WARNING size of work(1)',int(work(1))
  end if
  call dsyev('vectors','Upper',N,H,N,S,work,lwork,info)
  if(info.gt.0) then
    write(*,*) "Diagonalization > WARNING The algorithm computing failed"
    stop
  end if
 deallocate(work)
end subroutine diagonalization
l ------
!
              COMPUTE_T()
subroutine compute_T(T,V,nvec,m,pot_ext)
  implicit none
 double precision,allocatable :: V(:,:),T(:,:),pot_ext(:)
  integer :: nvec
 type(t_mesh)::m
  integer :: i,j,k,l
 double precision :: deltasqr,acc
 double precision, parameter::alpha=0.0
  double precision::beta
 deltasqr=m%dx**2
  !$OMP PARALLEL private(acc)
  !$OMP DO
 do j=1,nvec
    do i=1,nvec ! Tij
       T(i,j)=0.0
       do k=1,m%N
          acc=(3.0/deltasqr+pot_ext(k))*V(k,j)! the potential will be added here
          do l=1,m%n_neighbors(k)
             acc=acc-0.5*V(m%list_neighbors(k,1),j)/deltasqr
              print *, omp_get_thread_num(),i,j,k,l
          end do
```

```
T(i,j)=T(i,j)+V(k,i)*acc
!
             print *,' -> ',omp_get_thread_num(),i,j,T(i,j)
          print *,omp_get_thread_num(),i,j,T(i,j)
ļ
      end do
   end do
   !$OMP END DO
   !$OMP END PARALLEL
    stop
 end subroutine compute_T
                INIT_BASIS_SET()
 subroutine init_basis_set(V,nvec,seed,m)
   implicit none
   integer :: nvec,seed
   double precision,allocatable :: V(:,:)
   type(t_mesh)::m
   double precision, external :: ddot
   double precision ::normloc
   integer :: i,j
   double precision,allocatable :: Vdump(:,:)
   type(t_GramSchmidt) :: GS
   allocate(Vdump(m%N,nvec))
   call srand(seed)
   do i=1,nvec
      do j=1,m%N
         Vdump(j,i)=rand()
      end do
   end do
   do i=1,nvec
      normloc=ddot(m%N,Vdump(:,i),1,Vdump(:,i),1)
      normloc=1.0/sqrt(normloc)
      call dscal(m%N,normloc,Vdump(:,i),1)
   end do
   GS%nindep=1
   call GramSchmidt(V,Vdump,nvec,m,GS)
   deallocate(Vdump)
 end subroutine init_basis_set
 ! -----
 ! Ref.: D. G. Clayton "Gram-Schmidt Orthogonalization", J. Roy. Stat. Soc. C 20, 335 (1971)
 subroutine GramSchmidt(Vout, Vin, nvec, m, GS)
   implicit none
   integer :: nvec
   double precision,allocatable :: Vin(:,:),Vout(:,:)
   type(t_mesh)::m
```

```
type(t_GramSchmidt) :: GS
  integer :: i,k,i0
  double precision, parameter :: ETA=1.0e-6
  double precision,allocatable :: a(:)
  double precision :: normloc
  double precision, external :: ddot
  allocate(a(nvec))
  print *,'GS> -----'
 print *,"GS> Gram-Schmidt algorithm"
 print *,'GS> -----'
 print *,'GS> ',nvec,' vectors to orthogonalize'
 print *,'GS> ',GS%nindep,' vectors are already orthogonalized'
 Vout(:,1:GS%nindep)=Vin(:,1:GS%nindep)
 GS%ndep=0
  i0=GS%nindep
 do i=i0+1,nvec
    Vout(:,GS%nindep+1)=Vin(:,i)
    do k=1,GS%nindep
        ! We compute the projection of Vini(:,i) on V(:,1-nindep)
       a(k)=ddot(m%N, Vout(:,k),1, Vin(:,i),1)
       !print *,'GS > ',i,k,a(k)
        ! then we remove V(:,k) from V(:,nindep+1)
       call daxpy(m%N,-a(k),Vout(:,k),1,Vout(:,GS%nindep+1),1)
     end do
     ! now wre compute the norm of V(:,nindep+1)
    normloc=sqrt(ddot(m%N, Vout(:,GS%nindep+1),1,Vout(:,GS%nindep+1),1))
     !print *,'GS > norm(',i,')=',norm
     if (normloc.le.ETA) then
       GS%ndep=GS%ndep+1 ! V(:,nindep+1) is not linearly inependent
     else
       normloc=1.0/normloc
       call dscal(m%N,normloc,Vout(:,GS%nindep+1),1)
       !
                      do k=2,icur
        !
                         print *,'<U',k-1,'|U',i,'>=',ddot(N,V(:,k-1),1,V(:,icur),1)
                      end do
       GS%nindep=GS%nindep+1
     end if
  print *,'GS> ',GS%ndep,' vectors linearly dependant'
 print *,'GS> ',GS%nindep,' vectors linearly independent'
 print *,'GS> Size of the basis from ',nvec,' to ',GS%nindep
  !call check_ortho(Vout,nvec,m)
  !stop
 nvec=GS%nindep
 deallocate(a)
end subroutine GramSchmidt
l -----
subroutine check_ortho(P,nvec,m)
  implicit none
  integer :: nvec
 double precision :: P(:,:)
```

```
type(t_mesh)::m
 double precision, parameter :: ETA=1.0e-6
  integer :: i,j,nfail
 double precision :: pscal
 double precision, external :: ddot
 nfail=-nvec
 print *,"--- check_ortho() ---"
 do i=1,nvec
    do j=1,nvec
       pscal=ddot(m%N,P(:,i),1,P(:,j),1)
       write(*,'(F10.2)',advance='no') pscal
       if (pscal.gt.ETA) nfail=nfail+1
    write(*,*)
 end do
  if (nfail.gt.0) then
    print *,nfail,' fail(s)'
    stop
  end if
end subroutine check_ortho
! -----
subroutine init_mesh(m,param)
 implicit none
 type(t_mesh)::m
 type(t_param)::param
 double precision:: Lwidth
 Lwidth=param%box_width
 m%Nx=param%Nx
  m\%Nx=5
 m\%Ny=m\%Nx
 m\%Nz=m\%Nx
 m%N=m%Nx*m%Ny*m%Nz
 m%dx=Lwidth/(m%Nx+1)
 m%dy=Lwidth/(m%Ny+1)
 m%dz=Lwidth/(m%Nz+1)
 m\%dv=m\%dx*m\%dy*m\%dz
  !m\%dv=1.0
 m%center(1)=Lwidth/2
 m%center(2)=Lwidth/2
 m%center(3)=Lwidth/2
 allocate(m%n_neighbors(m%N))
  allocate(m%list_neighbors(m%N,6)) !
 m%list_neighbors(:,:)=0
 m%n_neighbors(:)=0
  call compute_list_neighbors(m)
end subroutine init_mesh
```

```
subroutine free_mesh(m)
 implicit none
 type(t_mesh) :: m
 deallocate(m%n_neighbors)
 deallocate(m%list_neighbors) !
end subroutine free_mesh
! -----
subroutine compute_list_neighbors(m)
  implicit none
 type(t_mesh) :: m
  integer::i,j,k,nn
  !integer,allocatable::n_neighbors(:),list_neighbors(:,:)
 do k=1,m\%Nz
    do i=1,m%Nx
       do j=1,m\%Ny
          nn=j+(i-1)*m\%Ny+(k-1)*m\%Ny*m\%Nx
           if (k>1) then
             m%n_neighbors(nn)=m%n_neighbors(nn)+1
             m%list_neighbors(nn,m%n_neighbors(nn))=nn-m%Nx*m%Ny
           end if
           if (k<m%Nz) then
             m%n_neighbors(nn)=m%n_neighbors(nn)+1
             m%list_neighbors(nn,m%n_neighbors(nn))=nn+m%Nx*m%Ny
           end if
           if (i>1) then
             m%n_neighbors(nn)=m%n_neighbors(nn)+1
             m%list_neighbors(nn,m%n_neighbors(nn))=nn-m%Ny
          end if
           if (i<m%Nx) then
             m%n_neighbors(nn)=m%n_neighbors(nn)+1
             m%list_neighbors(nn,m%n_neighbors(nn))=nn+m%Ny
           end if
           if (j>1) then
             m%n_neighbors(nn)=m%n_neighbors(nn)+1
             m%list_neighbors(nn,m%n_neighbors(nn))=nn-1
           end if
           if (j<m%Ny) then
             m%n_neighbors(nn)=m%n_neighbors(nn)+1
             m%list_neighbors(nn,m%n_neighbors(nn))=nn+1
          end if
       end do
     end do
  end do
end subroutine compute_list_neighbors
```

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end program Hbinitio

References

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