# Electronic Structure Calculation using Plane Wave Basis Set

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# **Contents**

1	Introduction	1
2	Outline	1
3	PWGrid_xx.jl 3.1 PWGrid_01.jl	<b>1</b>
4	Visualizing real-space grid points	3
5	Solving Poisson equation	3
6	Solving Schrodinger equation	5
7	Formulae	5

## 1 Introduction

This document is part of package ffr-ElectronicStructure.jl using plane wave basis set.

### 2 Outline

First, I will describe the top level description about what the a typical electronic structure calculation based on density functional theory is carried out. After that, I will break it apart into smaller pieces which are hopefully easier to implement and be understood.

In a typical DFT calculation we are trying to solve the so-called Kohn-Sham equations:

$$H_{KS}\psi_{KS} = e_{KS}\psi_{KS} \tag{1}$$

3 PWGrid\_xx.jl

xx stands for the version (or variant) of PWGrid.

#### **3.1** PWGrid\_01.jl

In this file, a type PWGrid is defined:

```
type PWGrid
Ns::Array{Int64}
LatVecs::Array{Float64,2}
RecVecs::Array{Float64,2}
Npoints::Int
Q::Float64
r::Array{Float64,2}
```

```
G::Array{Float64,2}
G2::Array{Float64}
end
```

The fields of this type are:

- Ns is an integer array which defines number of sampling points in each lattice vectors.
- Lat Vecs is  $3 \times 3$  matrix which defines lattice vectors of unit cell in real space.
- RecVecs is  $3 \times 3$  matrix which defines lattice vectors of unit cell in reciprocal space. It is calculated according to (3).
- Npoints Total number of sampling points
- Ω Unit cell volume in real space
- r Real space grid points
- G G-vectors
- G2 Magnitude of **G**-vectors

Constructor for PWGrid is defined as follow.

```
function PWGrid( Ns::Array{Int,1}, LatVecs::Array{Float64,2} )
  Npoints = prod(Ns)
  RecVecs = 2*pi*inv(LatVecs')
  \Q = det(LatVecs)
  R,G,G2 = init_grids( Ns, LatVecs, RecVecs )
  return PWGrid( Ns, LatVecs, RecVecs, Npoints, Q, R, G, G2 )
end
```

The function init\_grid() is defined as follow. It takes Ns, LatVecs, and RecVecs as the arguments.

```
function init_grids( Ns, LatVecs, RecVecs )
```

First, grid points in real space are initialized:

```
Npoints = prod(Ns)
r = Array (Float 64, 3, Npoints)
ip = 0
for k in 0:Ns[3]-1
for j in 0:Ns[2]-1
for i in 0:Ns[1]-1
  ip = ip + 1
 r[1, ip] = LatVecs[1, 1]*i/Ns[1] + LatVecs[2, 1]*j/Ns[2]
           + LatVecs[3,1]*k/Ns[3]
  r[2, ip] = LatVecs[1, 2]*i/Ns[1] + LatVecs[2, 2]*j/Ns[2]
            + LatVecs[3,2]*k/Ns[3]
  r[3,ip] = LatVecs[1,3]*i/Ns[1] + LatVecs[2,3]*j/Ns[2]
            + LatVecs[3,3]*k/Ns[3]
end
end
end
```

In the next step, grid points in reciprocal space, or G-vectors and also their squared values are initialized

```
G = Array (Float 64, 3, Npoints)
G2 = Array(Float64, Npoints)
ip
      = 0
for k in 0:Ns[3]-1
for j in 0:Ns[2]-1
for i in 0:Ns[1]-1
  gi = mm_to_nn(i, Ns[1])
  gj = mm_to_nn(j, Ns[2])
  gk = mm_to_nn(k, Ns[3])
  ip = ip + 1
  G[1,ip] = RecVecs[1,1]*qi + RecVecs[2,1]*qj + RecVecs[3,1]*qk
  G[2,ip] = RecVecs[1,2]*gi + RecVecs[2,2]*gj + RecVecs[3,2]*gk
  G[3,ip] = RecVecs[1,3]*gi + RecVecs[2,3]*gj + RecVecs[3,3]*gk
  G2[ip] = G[1,ip]^2 + G[2,ip]^2 + G[3,ip]^2
end
end
end
```

The function mm\_to\_nn defines mapping from real space to Fourier space:

```
function mm_to_nn(mm::Int,S::Int)
  if mm > S/2
    return mm - S
  else
    return mm
  end
  end
end
```

Finally, the variables r, G, and G2 are returned.

```
return r,G,G2
```

We give an example of creating a PWGrid object:

```
Ns = [40, 40, 40]
LatVecs = 10*diagm(ones(3))
pw = PWGrid( Ns, LatVecs )
```

# 4 Visualizing real-space grid points

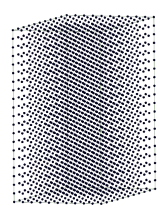
In the directory pwgrid\_01, we visualize grid points in real space using Xcrysden program. Originally Xcrysden, is meant to visualize crystalline structure, however, we also can use it to visualize grid points, taking periodic boundary conditions into consideration. This is useful to check whether grid points are generated correctly or not.

# 5 Solving Poisson equation

Poisson equation is relatively easy to solve in periodic boundary condition using Fourier method (FFT). This is different from other discretization method, such as finite difference or Lagrange function.

An example a program to solve Poisson equation is given in directory poisson\_01. In this program, a charge density is constructed from difference between two Gaussian charge density. Total charge (integrated charge density) is restricted to zero. From this charge density, we calculate the electrostatic (Hartree) potential by solving Poisson equation.

Function to generate vector dr:



```
function gen_dr( r, center )
  Npoints = size(r)[2]
  dr = Array(Float64, Npoints)
  for ip=1:Npoints
    dx2 = ( r[1,ip] - center[1] )^2
    dy2 = ( r[2,ip] - center[2] )^2
    dz2 = ( r[3,ip] - center[3] )^2
    dz2 = ( r[3,ip] - center[3] ) ^2
    dr[ip] = sqrt( dx2 + dy2 + dz2 )
  end
  return dr
end
```

Function to generate charge density:

```
function gen_rho( dr, \sigma1, \sigma2 )
   Npoints = size(dr)[1]
   rho = Array( Float64, Npoints )
   c1 = 2*\sigma1^2
   c2 = 2*\sigma2^2
   cc1 = sqrt(2*\sigmai^2\sigma)^3
   cc2 = sqrt(2*\sigmai^2\sigma2)^3
   for ip=1:Npoints
      g1 = exp(-dr[ip]^2/c1)/cc1
      g2 = exp(-dr[ip]^2/c2)/cc2
      rho[ip] = g2 - g1
   end
   return rho
end
```

Function to solve Poisson equation:

```
function solve_poisson( pw_grid::PWGrid, rho )

Q = pw_grid.Q

G2 = pw_grid.G2

Ns = pw_grid.Ns

Npoints = pw_grid.Npoints

ctmp = 4.0*pi*R_to_G( Ns, rho )

for ip = 2:Npoints

ctmp[ip] = ctmp[ip] / G2[ip]

end
```

```
ctmp[1] = 0.0
phi = real( G_to_R( Ns, ctmp ) )
return phi
end
```

# 6 Solving Schrodinger equation

Using energy minimization:

Introduction to minimization

simple 2D minimization, using steepest-descent and conjugate gradient method

Using iterative diagonalization: Davidson and LOBPCG

background information about iterative diagonalization Eigenvalue problems

### 7 Formulae

Plane wave basis  $b_{\alpha}(\mathbf{r})$ :

$$b_{\alpha}(\mathbf{r}) = \frac{1}{\sqrt{\Omega}} e^{\mathbf{G}_{\alpha} \cdot \mathbf{r}} \tag{2}$$

Lattice vectors of unit cell in reciprocal space:

$$\mathbf{b} = 2\pi \left( a^T \right)^{-1} \tag{3}$$

**G**-vectors:

$$\mathbf{G} = i\mathbf{b}_1 + j\mathbf{b}_2 + k\mathbf{b}_3 \tag{4}$$

Structure factor:

$$S_I(\mathbf{G}) = \sum_{\mathbf{G}} e^{-\mathbf{G} \cdot \mathbf{X}_I} \tag{5}$$