### 2023 Virtual School on Many-Body Calculations using EPW and BerkeleyGW

Resistivity, drift and Hall mobility with EPW

Hands-on Session (Mon.4)

Hands-on based on Quantum ESPRESSO (v7.2) and EPW v5.7  $\,$ 

# Introduction

In this tutorial, we will show how to compute the intrinsic electron and hole low-field drift and Hall mobility of the polar cubic semiconductor BN using the linearised iterative Boltzmann transport equation (IBTE) and the self-energy relaxation time approximation (SERTA), with or without external magnetic field. We will also see how to compute the electric resistivity of metals. You are advised to prepare the following script file, e.g. run.sh:

--#!/bin/bash #SBATCH -J job.ph # Job name #SBATCH -N 1 # Total # of nodes #SBATCH --ntasks-per-node 8 #SBATCH -t 01:00:00 # Run time (hh:mm:ss) #SBATCH -A EPSchool2022 #SBATCH -P small #SBATCH -p small #SBATCH --reservation=EPW-SCHOOL-Mon # Launch MPI code... export PATHQE=/work2/06868/giustino/SCHOOL/q-e-qe-7.2/ ibrun \$PATHQE/bin/pw.x < scf.in | tee scf.out ibrun \$PATHQE/bin/ph.x < ph.in | tee ph.out

For the description for all input flags please follow the link:

https://epwdoc.gitlab.io/source/doc/Inputs.html

# Exercise 1

### 1.1 Theory

In this example we are going to calculate the drift and Hall hole carrier mobility of c-BN. The drift mobility is obtained with:

$$\mu_{\alpha\beta}^{\rm d} = \frac{-1}{V^{\rm uc}n^{\rm c}} \sum_{n} \int \frac{\mathrm{d}^{3}k}{\Omega^{\rm BZ}} v_{n\mathbf{k}\alpha} \partial_{E_{\beta}} f_{n\mathbf{k}}$$
(1)

where the out of equilibrium occupations are obtained by solving the BTE:

$$\partial_{E_{\beta}} f_{n\mathbf{k}} = e v_{n\mathbf{k}\beta} \frac{\partial f_{n\mathbf{k}}^{0}}{\partial \varepsilon_{n\mathbf{k}}} \tau_{n\mathbf{k}} + \frac{2\pi\tau_{n\mathbf{k}}}{\hbar} \sum_{m\nu} \int \frac{\mathrm{d}^{3}q}{\Omega_{\mathrm{BZ}}} |g_{mn\nu}(\mathbf{k},\mathbf{q})|^{2} \\ \times \left[ (n_{\mathbf{q}\nu} + 1 - f_{n\mathbf{k}}^{0}) \delta(\varepsilon_{n\mathbf{k}} - \varepsilon_{m\mathbf{k}+\mathbf{q}} + \hbar\omega_{\mathbf{q}\nu}) + (n_{\mathbf{q}\nu} + f_{n\mathbf{k}}^{0}) \delta(\varepsilon_{n\mathbf{k}} - \varepsilon_{m\mathbf{k}+\mathbf{q}} - \hbar\omega_{\mathbf{q}\nu}) \right] \partial_{E_{\beta}} f_{m\mathbf{k}+\mathbf{q}}.$$
(2)

run.sh

The scattering rate in Eq. (2) is defined as:

$$\tau_{n\mathbf{k}}^{-1} \equiv \frac{2\pi}{\hbar} \sum_{m\nu} \int \frac{d^3q}{\Omega_{\rm BZ}} |g_{mn\nu}(\mathbf{k},\mathbf{q})|^2 [(n_{\mathbf{q}\nu} + 1 - f_{m\mathbf{k}+\mathbf{q}}^0) \\ \times \delta(\varepsilon_{n\mathbf{k}} - \varepsilon_{m\mathbf{k}+\mathbf{q}} - \hbar\omega_{\mathbf{q}\nu}) + (n_{\mathbf{q}\nu} + f_{m\mathbf{k}+\mathbf{q}}^0) \delta(\varepsilon_{n\mathbf{k}} - \varepsilon_{m\mathbf{k}+\mathbf{q}} + \hbar\omega_{\mathbf{q}\nu})].$$
(3)

A common approximation to Eq. (2) is called the self-energy relaxation time approximation (SERTA) and consists in neglecting the second term in the right-hand of the equation which gives:

$$\mu_{\alpha\beta}^{\text{SERTA}} = \frac{-e}{V^{\text{uc}}n^{\text{c}}} \sum_{n} \int \frac{\mathrm{d}^{3}k}{\Omega^{\text{BZ}}} \frac{\partial f_{n\mathbf{k}}^{0}}{\partial \varepsilon_{n\mathbf{k}}} v_{n\mathbf{k}\alpha} v_{n\mathbf{k}\beta} \tau_{n\mathbf{k}}.$$
(4)

The the low-field phonon-limited carrier mobility in the presence of a small finite magnetic field  ${\bf B}$  is given by:

$$\mu_{\alpha\beta}(B_{\gamma}) = \frac{-1}{V^{\mathrm{uc}}n^{\mathrm{c}}} \sum_{n} \int \frac{\mathrm{d}^{3}k}{\Omega^{\mathrm{BZ}}} v_{n\mathbf{k}\alpha} [\partial_{E_{\beta}} f_{n\mathbf{k}}(B_{\gamma}) - \partial_{E_{\beta}} f_{n\mathbf{k}}], \tag{5}$$

again solving the BTE with finite (small) magnetic field:

$$\left[1 - \frac{e}{\hbar} \tau_{n\mathbf{k}} (\mathbf{v}_{n\mathbf{k}} \times \mathbf{B}) \cdot \nabla_{\mathbf{k}}\right] \partial_{E_{\beta}} f_{n\mathbf{k}} (B_{\gamma}) = e v_{n\mathbf{k}\beta} \frac{\partial f_{n\mathbf{k}}^{0}}{\partial \varepsilon_{n\mathbf{k}}} \tau_{n\mathbf{k}} + \frac{2\pi \tau_{n\mathbf{k}}}{\hbar} \sum_{m\nu} \int \frac{\mathrm{d}^{3}q}{\Omega^{\mathrm{BZ}}} |g_{mn\nu}(\mathbf{k},\mathbf{q})|^{2} \\
\times \left[(n_{\mathbf{q}\nu} + 1 - f_{n\mathbf{k}}^{0}) \delta(\varepsilon_{n\mathbf{k}} - \varepsilon_{m\mathbf{k}+\mathbf{q}} + \hbar\omega_{\mathbf{q}\nu}) + (n_{\mathbf{q}\nu} + f_{n\mathbf{k}}^{0}) \delta(\varepsilon_{n\mathbf{k}} - \varepsilon_{m\mathbf{k}+\mathbf{q}} - \hbar\omega_{\mathbf{q}\nu})\right] \partial_{E_{\beta}} f_{m\mathbf{k}+\mathbf{q}} (B_{\gamma}). \quad (6)$$

The Hall factor and Hall mobility are then obtained as:

$$r_{\alpha\beta}(\hat{\mathbf{B}}) \equiv \lim_{\mathbf{B}\to 0} \sum_{\delta\epsilon} \frac{[\mu_{\alpha\delta}^{\mathrm{d}}]^{-1} \, \mu_{\delta\epsilon}(\mathbf{B}) \, [\mu_{\epsilon\beta}^{\mathrm{d}}]^{-1}}{|\mathbf{B}|} \tag{7}$$

$$\mu_{\alpha\beta}^{\text{Hall}}(\hat{\mathbf{B}}) = \sum_{\gamma} \mu_{\alpha\gamma}^{\text{d}} r_{\gamma\beta}(\hat{\mathbf{B}}),$$
(8)

where  $\hat{\mathbf{B}}$  is the direction of the magnetic field. More information can be found in the review Rep. Prog. Phys. **83**, 036501 (2020).

lonized impurity scattering is an important mobility limiting mechanism alongside thermal lattice vibrations in doped materials or materials with native ionized point defects. For 3D bulk crystals containing randomly distributed ionized impurities, we can exploit the Kohn-Luttinger ensemble average to obtain the partial transition rates of carriers by the ionized impurities.

$$\tau_{n\mathbf{k}\to m,\mathbf{k}+\mathbf{q}}^{-1,\mathrm{ii}} = n_{\mathrm{ii}} \frac{2\pi}{\hbar} \left[ \frac{e^2}{4\pi\varepsilon_0} \frac{4\pi Z}{\Omega_{\mathrm{u.c}}} \right]^2 \sum_{\mathbf{G}\neq -\mathbf{q}} \frac{|\langle u_{m\mathbf{k}+\mathbf{q}}|e^{i\mathbf{G}\cdot\mathbf{r}}|u_{n\mathbf{k}}\rangle_{\mathrm{u.c}}|^2}{|(\mathbf{q}+\mathbf{G})\cdot\varepsilon^0\cdot(\mathbf{q}+\mathbf{G})|^2} \delta(\epsilon_{n\mathbf{k}}-\epsilon_{m\mathbf{k}+\mathbf{q}}), \tag{9}$$

Here,  $n_{ii}$  is the concentration of ionized impurities in the system, Z is the charge of the ionized impurities, and  $\varepsilon^0$  is the low-frequency dielectric tensor which includes contributions from both electronic and ionic polarizability. More information can be found in the article Phys. Rev. B **107**, 125207 (2023).

### 1.2 Preliminary calculations with Quantum Espresso

First download the exercise files:

```
$ cp /work2/06868/giustino/SCHOOL/tutorials/Mon.4.Ponce_Leveillee.tar .
$ tar -xvf Mon.4.Ponce_Leveillee.tar
$ cd exercise1/
```

▶ Make a self-consistent calculation for c-BN.

```
--
&control
    calculation
                    = 'scf'
   prefix
                   = 'bn'
    restart_mode = 'from_scratch'
   pseudo_dir
                    = './'
                    = './'
    outdir
 1
&system
                    = 2
   ibrav
    celldm(1)
                    = 6.833
                    = 2
    nat
                    = 2
    ntyp
    ecutwfc
                    = 40
 1
&electrons
    diagonalization = 'david'
                 = 0.7
    mixing_beta
    conv_thr
                    = 1.0d-13
1
ATOMIC_SPECIES
 B 10.811 B-PBE.upf
N 14.0067 N-PBE.upf
ATOMIC_POSITIONS {crystal}
B 0.00 0.00 0.00
N -0.25 0.75 -0.25
K_POINTS automatic
888000
```

**Note**: In practice the **k**-point grid needs to be fairly large in order to get converged dielectric function and Born effective charges during the following phonon calculation.

```
$ ibrun $PATHQE/bin/pw.x -in scf.in | tee scf.out
```

► Compute the vibrational properties of c-BN on a coarse 4x4x4 q-point grid.

```
--
&inputph
    tr2_ph=1.0d-17,
    prefix='bn',
    amass(1)=10.811,
    amass(2)=14.0067,
    outdir='./',
    fildyn='bn.dyn.xml',
    fildvscf='dvscf'
    ldisp=.true.,
    epsil=.true.,
    nq1 = 4,
    nq2 = 4,
    nq3 = 4
/
```

**Note**: We have the input variable epsil=.true. which computes the macroscopic dielectric constant in non-metallic systems. If you add .xml after the name of the dynamical matrix file, it will produce the data in XML format (preferred).

ph.in

scf.in

**Note 2**: The input variable responsible to produce the electron-phonon matrix element is **fildvscf**. Always make sure that this variable is present.

**Note 3**: Notice the very tight tr2\_ph threshold parameter on the self-consistent first-order perturbed wavefunction. This is crucial to obtain good vibrational properties.

\$ ibrun \$PATHQE/bin/ph.x -in ph.in | tee ph.out

The calculation should take about 5 min on 4 cores. During the run, notice the IBZ q-point grid:

```
Dynamical matrices for (4, 4, 4) uniform grid of q-points
( 8 q-points):
 Ν
          xq(1)
                       xq(2)
                                    xq(3)
     0.00000000 0.00000000
                              0.00000000
 1
 2 -0.25000000 0.25000000 -0.25000000
 3
     0.50000000 -0.50000000
                             0.50000000
 4
     0.000000000
                0.50000000
                              0.00000000
 5
     0.75000000 -0.25000000
                               0.75000000
 6
     0.500000000
                 0.000000000
                               0.50000000
 7
     0.000000000
                -1.000000000
                              0.00000000
 8 -0.50000000 -1.00000000
                              0.00000000
```

#### as well as the dielectric function and Born effective charge tensor:

	Dielec	tric cor	istai	nt in	cartesian	axi	S					
	•	-0.0000	0000	52       -0.00000000         50       4.597197252         50       0.00000000			52	0.00000000			)	
E*y	atom ( ( (	1 1.892 -0.000 0.000	B 277 000	Mean	Drce / dE) Z*: -0.00000 1.89277 -0.00000 Z*:	1.	89277 -0.0 -0.0	0000	)))	with	asr	applied:
	(	-1.892 0.000	277 )00		0.00000 -1.89277 0.00000		0.0	0000	))			

The experimental dielectric constant in c-BN is about 4.46. More accurate values can be obtained with larger k-point grids. c-BN is a polar material and has a Born effective charge of 1.89 which is very close to theoretical value of 1.91.

Finally, we need to post-process some of the data to make it ready for EPW. To do so, we can use a python script (usually provided in QE/EPW/bin/pp.py but copied here for convenience).

- ▶ Run the python post-processing to create the save folder
- \$ python3 \$PATHQE/EPW/bin/pp.py

The script will ask you to enter the prefix used for the calculation. In this case enter "bn". The script will create a new folder called "save" that contains the dvscf potential files, pattern files, and dynamical matrices on the IBZ.

► Compute the low-frequency dielectric constant including ionic polarization for c-BN.

We need one final ingredient before moving on to running EPW if we intend to examine ionized impurity scattering, and that is to determine the zone-center dielectric polarizability of c-BN including ionic polarization. We have the electronic contribution from ph.x, but we will use dynmat.x to determine  $\varepsilon^0$ . Below is input for dynmat.x, dynmat.in.

dynmat.in

```
&input
fildyn='bn.dyn1.xml',
amass(1)=10.811,
amass(2)=14.0067,
asr='simple',
lperm=.true. ! calculate ionic polarizability
filout='bn.dynmat.out',
fileig='bn.eig'
/
```

To do this, run:

#### \$ ibrun \$PATHQE/bin/dynmat.x -in dynmat.in | tee dynmat.out

We can find the following lines in the dynmat.out file:

```
Electronic dielectric permittivity tensor (F/m units)

4.597197 0.000000 0.000000

-0.000000 4.597197 -0.000000

0.000000 0.000000 4.597197

... with zone-center polar mode contributions

6.629671 0.000000 -0.000000

0.000000 6.629671 -0.000000

-0.000000 -0.000000 6.629671
```

We will need the value of  $\approx$  6.63 for  $\epsilon_0$  to include ionized impurity scattering later on.

#### 1.3 Interpolation of the electron-phonon matrix element in real-space with EPW

▶ Do a non self-consistent calculation on a 4×4×4 uniform and  $\Gamma$ -centered k-point grid with crystal coordinates in the interval [0,1]

Such a grid can be for example generated with the wannier90 utility with kmesh.pl 4 4 4. The nscf.in file is as follow:

```
&control
                  = 'nscf'
   calculation
                  = 'bn'
   prefix
                  = 'from_scratch'
   restart_mode
  pseudo_dir
                  = './'
                  = './'
   outdir
&system
                  = 2
   ibrav
  celldm(1)
                  = 6.833
                  = 2
   nat
                  = 2
   ntyp
   ecutwfc
                  = 40
                  = 20
   nbnd
```

\_\_\_

nscf.in

```
&electrons
   diagonalization = 'david'
   mixing_beta = 0.7
                  = 1.0d-13
   conv_thr
ATOMIC_SPECIES
 B 10.811 B-PBE.upf
 N 14.0067 N-PBE.upf
ATOMIC_POSITIONS {crystal}
B 0.00 0.00 0.00
N -0.25 0.75 -0.25
K_POINTS crystal
64
 0.0000000 0.0000000 0.0000000 1.562500e-02
 0.00000000 0.00000000 0.25000000 1.562500e-02
 0.00000000 0.0000000 0.50000000 1.562500e-02
. . .
```

#### \$ ibrun \$PATHQE/bin/pw.x -in nscf.in | tee nscf.out

The reason for the non-self consistent calculation is that EPW needs the wavefunctions on the full BZ on a grid between 0 and 1.

**Note**: Since we are also interested in electron mobility, we will need the conduction bands. Notice that we added the input nbnd = 20 in nscf.in

▶ Perform an EPW calculation to Fourier-transform the electron-phonon matrix element from a coarse  $4\times4\times4$  k and q-point grids to real space and then interpolate the electronic band structure and phononic dispersion along the  $L - \Gamma - X - K - \Gamma$  high symmetry line by reading the file LGXKG.txt.

```
epw1.in
&inputepw
              = 'bn'
 prefix
 outdir
              = './'
 elph
              = .true.
 epbwrite
             = .true.
 epbread
              = .false.
 epwwrite
              = .true.
 epwread
              = .false.
 etf_mem
             = 1
                        ! polar material
 lpolar
             = .true.
              = 'dipole'
 vme
 nbndsub
             = 3
 bands_skipped = 'exclude_bands = 1, 5-20'
 wannierize = .true.
              = 50000
 num iter
 iprint
              = 2
 dis_win_max = 12.0
 dis_win_min = -1.0
 proj(1)
             = 'N:p'
 wdata(1) = 'bands_plot = .true.'
 wdata(2) = 'begin kpoint_path'
 wdata(3) = ' L 0.500 0.500 0.500 G 0.000 0.000 '
 wdata(4) = ' G 0.000 0.000 0.000 X 0.500 0.000 0.500 '
wdata(5) = ' X 0.500 0.000 0.500 K 0.375 0.375 0.750 '
 wdata(6) = ' K 0.375 0.375 0.750 G 0.000 0.000 '
 wdata(7) = 'end kpoint_path'
 wdata(8) = 'bands_plot_format = gnuplot'
 wdata(9) = 'guiding_centres = .true.'
 wdata(10) = 'dis_num_iter
                                 = 5000'
```

```
wdata(11) = 'num_print_cycles = 10'
wdata(12) = 'dis_mix_ratio
                              = 1.0'
wdata(13) = 'conv_tol = 1E-12'
wdata(14) = 'conv_window = 4'
wdata(15) = 'use_ws_distance = T'
            = 100
fsthick
           = 0.001
degaussw
dvscf_dir = './save'
band_plot = .true.
filkf
            = './LGXKG.txt'
            = './LGXKG.txt'
filqf
             = 4
 nk1
 nk2
             = 4
 nk3
             = 4
 nq1
             = 4
             = 4
 nq2
             = 4
 nq3
1
```

\$ ibrun \$PATHQE/bin/epw.x -npool 8 -input epw1.in | tee epw1.out

**Note**: The number of pool -npool has to be the same as the total number of core -np since k-point parallelization is (almost) the only parallelization level allowed. **G**-vector parallelization will be introduced in EPW v6.0.

The calculation should take less than 2 min. Note that the code should have detected the presence of the quadrupole.fmt file and correctly read the quadrupole tensor. Look in the output for the line Quadrupole tensor is correctly read:. In this hands-on we will not cover how to obtain the quadrupole tensor and they are simply given here. There are two ways to obtain them:

- Using perturbation theory. This is implemented in a recent version of the Abinit software.
- Fitting the perturbed density or the electron-phonon matrix elements in the long wavelength limit obtained by direct DFPT calculations.

More information can be found in Phys. Rev. Research 3, 043022 (2021)

At the end of the calculation, because of the keyword band\_plot = .true., the code should produce the band.eig and phband.freq files that contain the electronic band structure and phononic dispersion along a path given in the filkf and filqf files.

If you want to have files in an easy gnuplot format, you can use the plotband.x tool by doing

\$ \$PATHQE/bin/plotband.x

and follow the instructions. You should check that both plots look reasonable.

▶ Do a restart calculation (restarting from the bn.epmatwp1 file) and compute the hole mobility of c-BN.

```
$ ibrun $PATHQE/bin/epw.x -npool 8 -input epw2.in | tee epw2.out
```

The input file is as follow:

```
&inputepw
prefix = 'bn'
```

= './' outdir = .true. elph = .false. epwwrite = .true. epwread etf\_mem = 3 ! generate k-points within fsthick lpolar = .true. vme = 'dipole' mp\_mesh\_k = .true. = 3 nbndsub bands\_skipped = 'exclude\_bands = 1, 5-20' scattering = .true. scattering\_serta = .true. int\_mob = .false. = .true. = -1E13 carrier ncarrier iterative\_bte = .true. epmatkqread = .false. mob\_maxiter = 300 broyden\_beta= 1.0 = 0.0d0 bfieldx bfieldy = 0.0d0 bfieldz = 1.0d-10 ! Apply a magnetic field along Cart. z = 1 nstemp = 300 temps restart = .true.  $restart_step = 1000$ wannierize = .false. num\_iter = 50000 iprint = 2 iprint dis\_win\_max = 12.0 dis\_win\_min = -1.0 proj(1) = 'N:p' elecselfen = .false. phonselfen = .false. = .false. a2f = 0.4 ! 0.3 eV fsthick degaussw = 0.0 efermi\_read = .true  $fermi_energy = 11.246840$ dvscf\_dir = './save' nkf1 = 30 nkf2 = 30 nkf3 = 30 = 30 nqf1 = 30 nqf2 = 30 nqf3 nk1 = 4 = 4 nk2 nk3 = 4 = 4 nq1 nq2 = 4 = 4 nq3 /

Notes:

- The value of fermi\_energy was obtained from the output of the previous calculation epw1.in
- epwread allows for the restart from the bn.epmatwp1 file

- int\_mob allows to perform both electron and hole calculations at the same time but is not recommanded.
- carrier and noarrier define the carrier concentration. If carrier = .true. then the intrinsic mobility with noarrier concentration (in cm<sup>-3</sup>) is computed. If noarrier is positive it will compute the electron mobility and if it is negative it will compute the hole mobility. The resulting mobility should be independent of the choice of carrier concentration in reasonable ranges  $10^{10}$   $10^{16}$  cm<sup>-3</sup>.
- iterative\_bte asks for the iterative solution of the BTE in addition to SERTA.
- nstemp and temps define the lattice temperature at which the mobility is evaluated.
- restart and restart\_step will create restart point every (in this case) 1000 q-points. You can try breaking the run after a restart point and restart to test this feature.
- bfieldz adds a (small) finite magnetic field along the Cartesian z direction (in unit of Tesla). This will automatically trigger the calculation of the Hall factor.
- mob\_maxiter is the maximum number of iterations for the BTE solution.
- degaussw = 0.0 means that adaptive smearing is used. Positive values give Gaussian smearing.

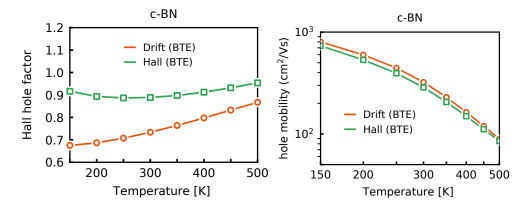
The run should take about 4 min. The fine  $\mathbf{k}$  and  $\mathbf{q}$  point grids need to be much denser for real calculations. However, we can already get relatively decent results.

▶ Re-run the code with multiple temperatures (using nstemp = 4 and temps = 100, 200, 400, 500). You should remove the restart.fmt file before doing so.

Try filling the table below for the hole mobility:

T (K)	hole $\varepsilon_F$ (eV)	drift SERTA $\mu$ (cm $^2$ /Vs)	drift BTE $\mu$ (cm $^2$ /Vs)	Hall BTE $\mu$ (cm $^2$ /Vs)
100				
200				
300				
400				
500				

At convergence you should get <sup>1</sup>:



where the room temperature values with SOC should be around 319  $cm^2/Vs$  for the drift BTE and 281  $cm^2/Vs$  for the Hall mobility with a Hall factor of 0.88.

▶ Try to increase the fine grids and add a few more temperatures and see if you can get a result closer to convergence.

► Try adding SOC

▶ Try removing or renaming the file quadrupole.fmt to do the interpolation with dipole only and see the impact on the results.

<sup>1</sup>The figure is from Phys. Rev. Research **3**, 043022 (2021)

### 1.4 Compute the spectral decomposition

 $\blacktriangleright$  Do a restart calculation (restarting from the bn.epmatwp1 file) and compute the hole spectral decomposition of c-BN.

> You should remove the restart.fmt file.

\$ ibrun \$PATHQE/bin/epw.x -npool 8 -input epw3.in | tee epw3.out

The input file is as follow (we show only the difference wrt epw2.in):

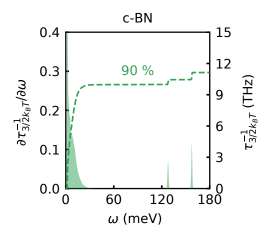
```
&inputepw
iverbosity = 3
mob_maxfreq = 160
mob_nfreq
            = 640 ! To have 0.25 meV intervals
            = 60
nkf1
nkf2
             = 60
             = 60
nkf3
            = 60
nqf1
nqf2
            = 60
nqf3
             = 60
```

At the end of the calculation, the code should have produced a file named inv\_tau\_freq.fmt. You should open the file and look for the maximum number of k-points within the fsthick, here you should have 272. You should also look for the number of bands, here you should find 3. Edit the gaussian-h.py python script to correspond to the calculation you have been doing. Then run it.

#### \$ python3 gaussian-h.py

The script should produce a file named inv\_tau\_freq.fmt-gaussian1.0 which you can plot with your favorite software. Note that the results are not converged but it should be clear that acoustic scattering is dominating in c-BN.

At convergence you should get <sup>2</sup>:



<sup>2</sup>The figure is from Phys. Rev. Research **3**, 043022 (2021)

epw3.in

### 1.5 Include carrier-ionized impurity scattering

► You will now run a calculations that includes the influence of ionized impurity scattering on the hole mobility of c-BN.

▶ You should once again remove the restart.fmt file:

```
$ rm restart.fmt
```

Then, run EPW:

\$ ibrun \$PATHQE/bin/epw.x -npool 8 -input epw4.in | tee epw4.out

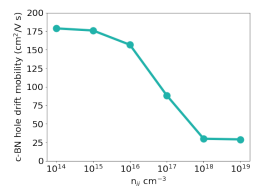
The input file for epw4.in is as follow (we show only the difference wrt epw2.in):

	epw4.in
&inputepw	
ncarrier = -1E17 ! absolute value MUST match ii_n, negative sign for holes.	
ii_g = .true.	
ii_charge = 1.0d0	
ii_n = 1.0d17 ! must match ncarrier, keep positive for both holes and electrons	
ii_scattering = .true.	
ii_only = .false.	
ii_eps0 = 6.62967d0	

Notes:

- $ii_g = .true$ . tells EPW to calculate ionized impurity matrix elements
- ii\_charge is the charge of the ionized impurities in units of the electron charge.
- ii\_n is the density of ionized impurities in units of cm<sup>-3</sup>. For bulk, typical doping values range from  $10^{15}$  to  $10^{18}$  cm<sup>-3</sup>, higher would result in degenerate doping.
- ii\_scattering = .true. tells EPW to compute the ionized impurity scattering rates.
- ii\_only: if ii\_only were set to .true., this would tell EPW to only include ionized impurity scattering and omit phonon scattering. Since we want phonon and ionized impurity scattering, we set ii\_only = .false..
- ii\_eps0 is the low-frequency dielectric constant, determined by dynmat.x.

After running epw.x with input epw4.in, we should find a hole mobility at 300 K of 88.25 cm<sup>2</sup>/V s, reduced from a phonon-only limited mobility of 178.87 cm<sup>2</sup>/V s previously calculated. The presence of ionized impurities strongly influences the drift mobilities. As an exercise, you can modify ii\_n (and ncarrier accordingly), between 1.0d15 and 1.0d19 to see how the hole mobility at 300 K changes as the concentration of ionized impurities increases. Here is the plot of the expected results for this example.



NOTE: lonized impurity scattering and Hall mobilities have not been tested together. The use of the ionized impurity functionality in EPW should be currently limited to drift mobilities only.

# **Exercise 2**

In this example we are going to calculate the electric resistivity of fcc Pb using the Ziman formula and the Boltzmann transport equation (BTE). First go in the first exercise:

\$ cd exercise2

► Make a self-consistent calculation for Pb and a phonon calculation on a homogeneous 3×3×3 q-point grid.

Note: The ecutwfc need to be much larger for real calculations.

```
$ ibrun $PATHQE/bin/pw.x < scf.in | tee scf.out
$ ibrun $PATHQE/bin/ph.x < ph.in | tee ph.out</pre>
```

```
scf.in
&control
   calculation = 'scf'
prefix = 'pb'
restart_mode = 'from_scratch'
   pseudo_dir = './'
outdir = ' /'
                    = './'
    outdir
    verbosity
tprnfor
                    = 'high'
                    = .true.
    tstress
                     = .true.
 1
&system
    ibrav
                    = 2
    celldm(1)
                    = 9.27
                     = 1
    nat
    ntyp
                    = 1
    ecutwfc
                    = 30
    occupations = 'smearing'
                  = 'mp'
    smearing
    degauss
                     = 0.025
 1
&electrons
    diagonalization = 'david'
    mixing_beta = 0.7
conv_thr = 1.0d-12
    conv_thr
ATOMIC_SPECIES
Pb 207.2 pb_s.UPF
ATOMIC_POSITIONS crystal
Pb 0.0 0.0 0.0
K POINTS automatic
12 12 12 0 0 0
                                                                                                             ph.in
&inputph
```

%inputph
recover = .false.
tr2\_ph = 1.0d-17,
prefix = 'pb',
fildyn = 'pb.dyn.xml',
fildvscf = 'dvscf'
ldisp = .true.,
nq1 = 3,
nq2 = 3,
nq3 = 3
/

The important keyword is fildvscf as it will tell the code to write to file the change of potential due to ionic displacement  $\partial_{q\nu} V^{scf}$ . The ldisp input allows to calculate phonons for a grid of q-points specified by nq1, nq2, and nq3.

Note 1: For real calculations the coarse q-point grid should be converged on and is typically 6x6x6 or 8x8x8.

**Note 2**: The tr2\_ph variable is the threshold on the perturbed wavefunction obtained by solving the Sternheimer equation and should be very small.

**Note 3**: It is recommended to add XXX.xml at the end of the fildyn as it will force the code to write the output in XML format.

This should take about 5 min to be completed. In the output file, locate the list of 4 irreducible q points in the Brillouin Zone (IBZ):

```
Dynamical matrices for ( 3, 3, 3) uniform grid of q-points
( 4 q-points):
N xq(1) xq(2) xq(3)
1 0.000000000 0.000000000 0.000000000
2 -0.33333333 0.33333333 -0.333333333
3 0.000000000 0.6666666667 0.000000000
4 0.66666666667 -0.00000000 0.6666666667
```

For each **q**-point, a pb.dynX.xml file containing the dynamical matrix has been produced. The pb.dvscf files are located inside the \_ph0 folder.

▶ Gather the .dyn, .dvscf and patterns<sup>3</sup> files into a new save directory. This can easily be done using the pp.py python script.

```
$ python3 $PATHQE/EPW/bin/pp.py
```

The script will ask you to provide the prefix of your calculation (here "pb").

**b** Do a non self-consistent calculation on a homogeneous 3x3x3 uniform and  $\Gamma$ -centered grid between [0,1[ in crystal coordinates.

```
___
                                                                                                  nscf.in
&control
   calculation
                   = 'nscf'
   prefix
                   = 'pb'
                   = 'from_scratch'
   restart mode
                   = './'
   pseudo_dir
                   = './'
   outdir
                   = 'high'
    verbositv
 /
&system
   ibrav
                   = 2
   celldm(1)
                   = 9.27
   nat
                   = 1
                   = 1
   ntyp
   ecutwfc
                   = 30
   occupations
                   = 'smearing'
                   = 'mp'
   smearing
                   = 0.025
   degauss
   nbnd
                   = 10
&electrons
   diagonalization = 'david'
   mixing_beta = 0.7
                   = 1.0d-12
    conv_thr
ATOMIC_SPECIES
Pb 207.2
          pb_s.UPF
ATOMIC_POSITIONS crystal
Pb 0.00000000
                0.000000000
                               0.00000000
```

 $<sup>^{3}\</sup>mathsf{The}\ \mathtt{patterns}\ \mathtt{file}\ \mathtt{contains}\ \mathtt{the}\ \mathtt{basis}\ \mathtt{in}\ \mathtt{which}\ \mathtt{the}\ \mathtt{.dvscf}\ \mathtt{are}\ \mathtt{defined}.$ 

\$ ibrun \$PATHQE/bin/pw.x -input nscf.in | tee nscf.out

▶ Perform an EPW calculation to interpolate the electron-phonon matrix element from a coarse 3x3x3 to a dense 18x18x18 k-point and q-point grids.

```
___
                                                                                                 epw1.in
&inputepw
             = 'pb'
 prefix
             = './'
 outdir
 dvscf_dir = './save'
 elph
             = .true.
            = .true.
 epbwrite
             = .false.
 epbread
            = .true.
 epwwrite
 epwread
             = .false.
             = 'dipole'
 vme
 nbndsub
            = 4
 bands_skipped = 'exclude_bands = 1-5'
 wannierize = .true.
 num_iter = 300
 dis_win_max = 21
 dis_froz_min= -3
 dis_froz_max= 13.5
 proj(1) = 'Pb:sp3'
 wdata(1) = 'bands_plot = .true.'
 wdata(2) = 'begin kpoint_path'
 wdata(3) = 'G 0.00 0.00 0.00 X 0.00 0.50 0.50'
 wdata(4) = 'X 0.00 0.50 0.50 W 0.25 0.50 0.75'
 wdata(5) = 'W 0.25 0.50 0.75 L 0.50 0.50 '
 wdata(6) = 'L 0.50 0.50 0.50 K 0.375 0.375 0.75'
 wdata(7) = 'K 0.375 0.375 0.75 G 0.00 0.00 '
 wdata(8) = 'G 0.00 0.00 0.00 L 0.50 0.50 '
 wdata(9) = 'end kpoint_path'
 wdata(10) = 'bands_plot_format = gnuplot'
 elecselfen = .false.
 phonselfen = .true.
 a2f
             = .true.
 delta_approx = .true.
             = 4.0d0 ! Number of carriers for the Ziman resistivity formula
 nc
 fsthick
             = 6
                    ! eV
             = 1 ! K
 temps
             = 0.1 ! eV
 degaussw
              = 0.05 ! meV
 degaussq
 assume_metal = .true.
 ngaussw
              = -99 ! we want F-D distribution for metals
            = 18
 nkf1
 nkf2
             = 18
 nkf3
             = 18
 nqf1
             = 18
             = 18
 nqf2
 nqf3
             = 18
             = 3
 nk1
 nk2
             = 3
```

nk3 = 3 nq1 = 3 nq2 = 3 nq3 = 3

There are two ways to compute the resistivity of Pb:

- using the phonon self-energy and the Eliashberg transport spectral function in conjunction with the Ziman formula
- using the Boltzmann transport equation

We will do both and start with the first one for which we need the isotropic transport spectral function:

$$\alpha_{\rm tr}^2 F(\omega) = \frac{1}{2} \sum_{\nu} \int_{\rm BZ} \frac{d\mathbf{q}}{\Omega_{\rm BZ}} \omega_{\mathbf{q}\nu} \lambda_{\rm tr, \mathbf{q}\nu} \delta(\omega - \omega_{\mathbf{q}\nu}), \tag{10}$$

where the mode-resolved transport coupling strength is defined by:

$$\lambda_{\mathrm{tr},\mathbf{q}\nu} = \frac{1}{N(\varepsilon_F)\omega_{\mathbf{q}\nu}} \sum_{nm} \int_{\mathrm{BZ}} \frac{d\mathbf{k}}{\Omega_{\mathrm{BZ}}} |g_{mn,\nu}(\mathbf{k},\mathbf{q})|^2 \delta(\varepsilon_{n\mathbf{k}} - \varepsilon_{\mathrm{F}}) \delta(\varepsilon_{m\mathbf{k}+\mathbf{q}} - \varepsilon_{\mathrm{F}}) \left(1 - \frac{v_{n\mathbf{k}} \cdot v_{m\mathbf{k}+\mathbf{q}}}{|v_{n\mathbf{k}}|^2}\right).$$
(11)

The calculation of the spectral function in EPW is given by the two keywords phonselfen = .true. and a2f = .true.

Note 1: The dvscf\_dir = './save' specify the place where we have placed the .dyn, .dvscf and patterns using the python script.

Note 2: Here we are using the delta\_approx = .true. to approximate the double  $\delta$  in Eq. (2). In this case the broadening of Dirac deltas is approximated by a Gaussian of widths given by degaussw = 0.1 and the Dirac delta in Eq. (1) by degaussq = 0.05.

**Note 3**: The variable ngaussw is used for calculation of the Fermi level and DOS and is a Fermi-Dirac distribution function (input -99) of electronic temperature given by temps = 1 K such that all the files names will end in XXX.1.000K.

```
$ ibrun $PATHQE/bin/epw.x -npool 8 -input epw1.in | tee epw1.out
```

The calculation should take about 6 min to be completed. While the calculation is running, notice in the epw1.out the different steps a full EPW run goes into. First the Wannierization, then the unfolding into the full  $3\times3\times3$  BZ, then the Fourier transform to real space and then finally the interpolation into the fine  $18\times18\times18$  k and q grids.

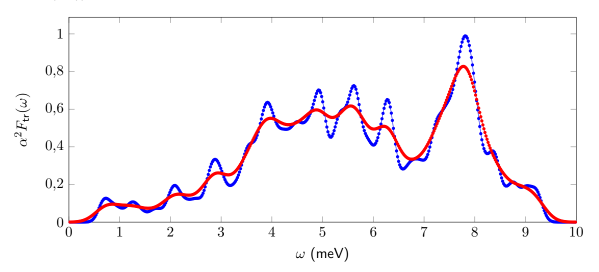
At the end of the calculation, you should get:

```
Eliashberg Spectral Function in the Migdal Approximation
_____
lambda :
        1.9271341
lambda_tr : 1.4564165
Estimated Allen-Dynes Tc
logavg =
        0.0002066 \ l_a 2f = 1.9281703
m11 =
     0.10 \ {\rm Tc} =
                   4.573698490498 K
     0.12 \ {\rm Tc} =
                    4.363659787196 K
mu =
mu =
     0.14 \ {\rm Tc} =
                    4.152675719793 K
     0.16 \ Tc =
                    3.941000842563 K
mu =
     0.18 \ {\rm Tc} =
                    3.728922987482 K
m11 =
mu =
     0.20 \ Tc =
                    3.516766489845 K
```

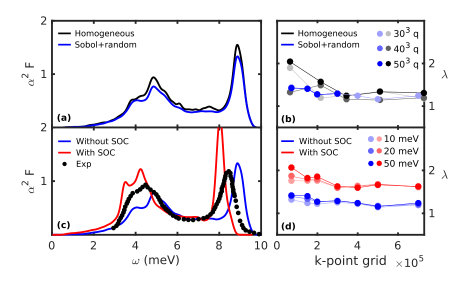
Note that the converged value for  $\lambda$  and  $\lambda_{tr}$  should be around 1.1. In addition the files pb.a2f.01.1.000 and pb.a2f\_tr.01.1.000 which contain the Eliashberg spectral function and transport spectral function for different broadening values should have been produced.

Tip: Look at the end of the pb.a2f\_tr.01.1.000 file to know which column corresponds to which broadening.

You should get something similar to this (here shown for two broadening values 0.15 meV (blue) and 0.3 meV (red)):



Again this is unconverged. At convergence you should get something closer to <sup>4</sup>:



► Compute the resistivity of Pb using the Ziman's formula for metals:

$$\rho(T) = \frac{4\pi m_e}{ne^2 k_B T} \int_0^\infty d\omega \,\hbar\omega \,\alpha_{\rm tr}^2 F(\omega) \,n(\omega, T) \big[ 1 + n(\omega, T) \big],\tag{12}$$

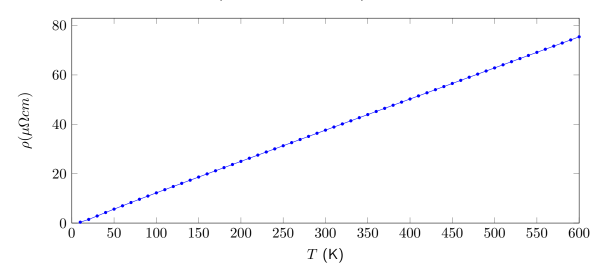
where n is the number of electrons per unit volume and  $n(\omega, T)$  is the Bose-Einstein distribution. Usually this means the number of electrons that contribute to the mobility which is 4.0 (can be fractional) for the case of Pb and given with the input variable nc = 4.0d0.

<sup>4</sup>The figure is from Comput. Phys. Commun. **209**, 116 (2016).

The resistivity was actually computed during the previous run. A file named pb.res.01.1.000 should have been created. The file contains the resistivity (in  $\mu$ Ohm cm) for various temperatures and smearing values (in meV).

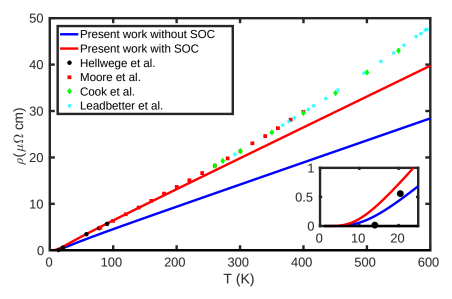
# Temperature [K]				Resistivity [micro Ohm cm] for different Phonon smearing (meV)							
#		0.0500000	0.1000000	0.1500000	0.2000000	0.2500000	0.3000000	0.3500000	0.4000000	0.4500000	0.5000000
	10	0.3512365	0.3525075	0.3547220	0.3580869	0.3631532	0.3708099	0.3812646	0.3936785	0.4068816	0.4199550
	20	1.4725265	1.4750212	1.4793713	1.4859905	1.4959803	1.5111188	1.5318215	1.5564122	1.5825522	1.6084046
	30	2.8584829	2.8621542	2.8685611	2.8783246	2.8930970	2.9155452	2.9462931	2.9828262	3.0216362	3.0599667
	40	4.2577891	4.2626474	4.2711290	4.2840617	4.3036484	4.3334453	4.3742842	4.4228124	4.4743510	4.5252226
	50	5.6291937	5.6352487	5.6458209	5.6619449	5.6863747	5.7235552	5.7745264	5.8350971	5.8994175	5.9628891

You should get the following graph (for 0.15 meV smearing):



Note that in this case, as it is an integrated quantity, it is not so dependent on smearing. Compare your result with other smearing.

At convergence you should get  $^5\colon$ 



▶ Now compute the conductivity of Pb using the BTE:

$$\sigma_{\alpha\beta} = \frac{-e}{V_{\rm uc}} \sum_{n} \int \frac{\mathrm{d}^{3}k}{\Omega_{\rm BZ}} v_{n\mathbf{k}}^{\alpha} \partial_{E_{\beta}} f_{n\mathbf{k}}$$
(13)

<sup>5</sup>The figure is from Comput. Phys. Commun. **209**, 116 (2016).

\$ ibrun \$PATHQE/bin/epw.x -npool 8 -input epw2.in | tee epw2.out

```
epw2.in
&inputepw
 -
prefix
              = 'pb'
              = './'
  outdir
  dvscf_dir = './save'
  elph
              = .true.
  epwwrite
              = .false. ! Restarting
              = .true. ! Restarting by reading the pb.epmatwp file
  epwread
              = 'dipole'
  vme
             = 4
  nbndsub
  bands_skipped = 'exclude_bands = 1-5'
  wannierize = .false. ! Restarting.
 num_iter = 300
  dis_win_max = 21
  dis_froz_min= -3
 dis_froz_max= 13.5
            = 'Pb:sp3'
  proj(1)
  wdata(1) = 'bands_plot = .true.'
  wdata(2) = 'begin kpoint_path'
  wdata(3) = 'G 0.00 0.00 0.00 X 0.00 0.50 0.50'
  wdata(4) = 'X 0.00 0.50 0.50 W 0.25 0.50 0.75'
  wdata(5) = 'W 0.25 0.50 0.75 L 0.50 0.50 0.50'
  wdata(6) = 'L 0.50 0.50 0.50 K 0.375 0.375 0.75'
  wdata(7) = 'K 0.375 0.375 0.75 G 0.00 0.00 0.00'
  wdata(8) = 'G 0.00 0.00 0.00 L 0.50 0.50 '
  wdata(9) = 'end kpoint_path'
  wdata(10)= 'bands_plot_format = gnuplot'
  elecselfen = .false.
  phonselfen = .false.
  a2f
              = .false.
              = 0.4 ! eV - we only need states close to Fermi level
  fsthick
            degaussw
  assume_metal = .true.
  ngaussw
              = -99 ! we want F-D distribution for metals
 int_mob
              = .true.
  iterative_bte = .true. ! SERTA and iterative BTE
 scattering = .true. ! compute scattering rates
carrier = .false. ! This is a metal, we do not specify carrier concentration
 mp_mesh_k
               = .true. ! Use crystal symmetries
 epmatkqread = .false. ! Can be used to just perform BTE iterations
mob_maxiter = 200 ! Max nb of BTE iterations
broyden_beta = 0.7 ! Broyden mixing during iterations
           = .true. ! Activate possible restart
 restart
  restart_step = 50
                           ! Write restart points every 50 q-points
  selecqread = .false.
               = 9
                           ! compute conductivity at 9 temperatures
  nstemp
               = 100 500
  temps
              = 30
  nkf1
  nkf2
              = 30
  nkf3
              = 30
 nqf1
              = 30
  nqf2
              = 30
              = 30
  nqf3
              = 3
  nk1
 nk2
              = 3
 nk3
              = 3
              = 3
 nq1
              = 3
 nq2
              = 3
 nq3
```

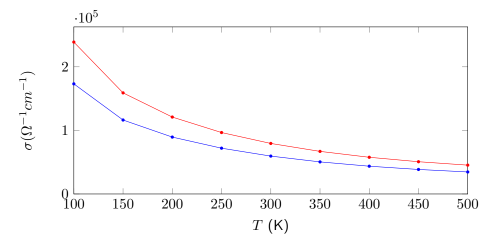
**Note 1**: In this calculation, we are restarting from the electron-phonon matrix elements written in real space in the pb.epmatwp file.

**Note 2**: In the case of BTE, temps corresponds to the real lattice temperature and degaussw is used to approximate the Dirac deltas. In this case it has the value 0.0 eV which means that an adaptative smearing is used. The value of the smearing is therefore band and k-point dependent and depends on the fine grid size: the denser the grids, the smaller the smearing.

**Note 3**: Because of the restart = .true. input parameter, if you want to do a clean restart, you need to remove the restart file by doing rm restart.fmt.

► Check the output file to find the minimum and maximum values of smearing reported. You can try changing the fine grids to see how it affects the smearing.

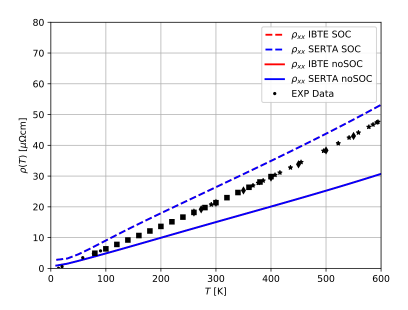
► Extract the SERTA and BTE conductivity as a function of temperature from the output file. You should get something like this (blue SERTA, red BTE):



Note: The resistivity is the inverse of the conductivity, therefore you can also obtain it.

At convergence you should get (BTE and SERTA are almost the same in this case, figure courtesy of Félix Goudreault):

| /



► You can try to include SOC using noncolin =.true. and lspinorb =.true. in scf.in and re-do everything to see the impact. Note that the calculations will be longer and that you need to double the number of Wannier functions and bands.