# Tutorial on phonon calculations with Quantum-Espresso

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## **Outline**

#### Introduction

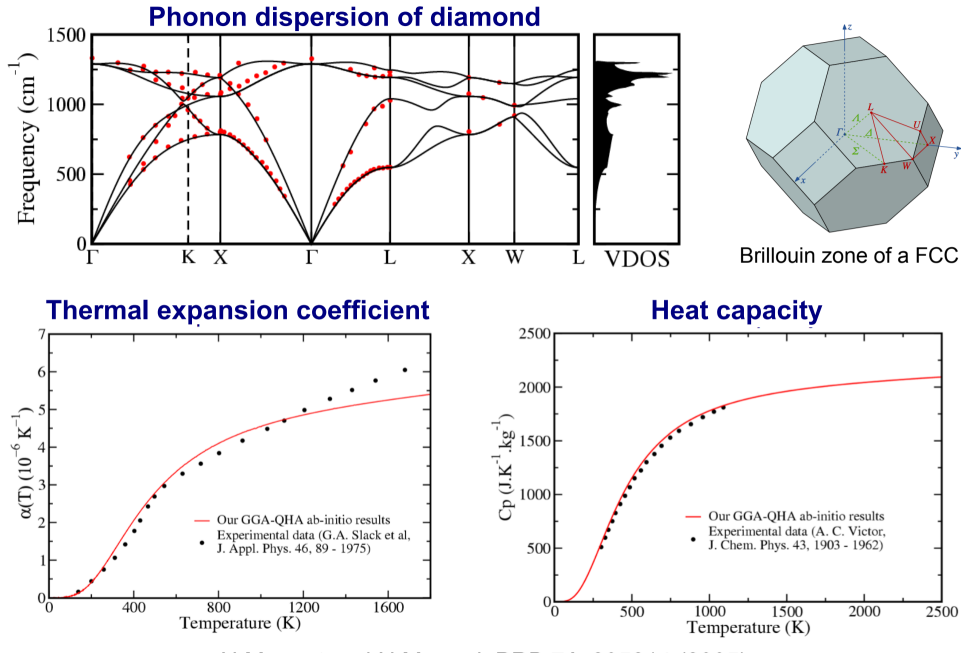
Exercise 1: Phonon calculations at Γ

Exercise 2: Phonons in polar materials

Exercise 3: Phonon dispersion and density of states

**Exercise 4: Tests** 

## Phonons and related quantities



N.Mounet and N.Marzari, PRB 71, 205214 (2005)

## **Phonons**

Normal mode frequencies,  $\omega$ , and eigenvectors,  $\mathbf{u}$ , are determined by the secular equation:

$$\sum_{s'\beta} D_{s\alpha s'\beta}(\boldsymbol{q}) \boldsymbol{u}_{s'\beta}(\boldsymbol{q}) = \omega_{\boldsymbol{q}}^2 \boldsymbol{u}_{s\alpha}(\boldsymbol{q})$$

where

$$D_{s \alpha s' \beta}(q) = \frac{1}{\sqrt{M_s M_{s'}}} \sum_{\nu} \left[ \frac{\partial^2 E_{tot}}{u_{\mu s \alpha} u_{\nu s' \beta}} \right] e^{i q (R_{\nu} - R_{\mu})}$$
Interatomic force constants (IFC)

is the dynamical matrix of the solid.

This matrix can be calculated from the linear response  $\partial n(\mathbf{r})/\partial \mathbf{u}_{s\alpha}(\mathbf{q})$  and diagonalized to get phonon modes at  $\mathbf{q}$ .

## Part 1: Calculation of the phonon frequencies of silicon at Γ

To perform the calculation use the script run\_ph\_G\_Si.sh in the directory exercise1:

```
prompt> ./run_ph_G_Si.sh &
```

The script allows you to perform:

- 1) A scf calculation for silicon at the equilibrium structure using pw.x
- 2) A phonon calculation at  $\Gamma$  using ph.x

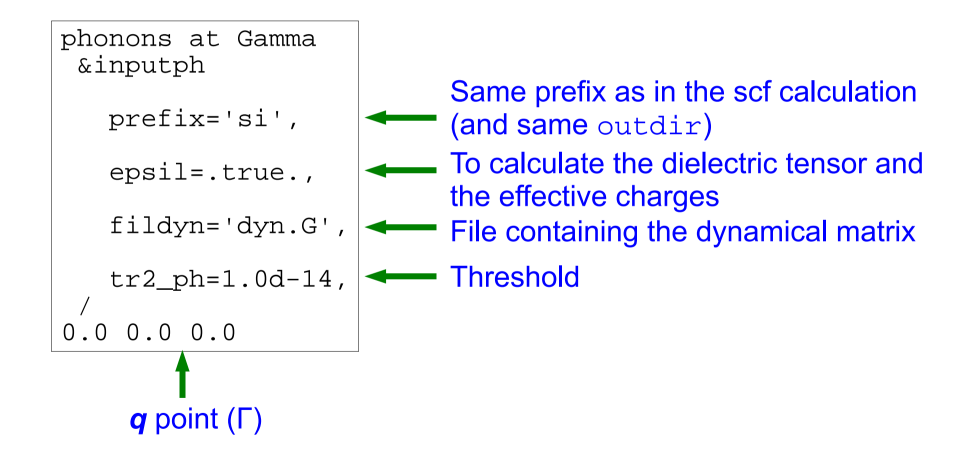
1) A scf calculation for silicon at the equilibrium structure using pw.x

```
&control
    calculation='scf'
    restart_mode='from scratch',
    prefix='si',
   pseudo_dir = '../',
 &system
    ibrav= 2,
    celldm(1) = 10.187,
    nat = 2,
    ntyp=1,
    ecutwfc =16
 &electrons
    conv thr = 1.0d-10
ATOMIC SPECIES
 Si 0.0 Si.pz-rrkj.UPF
ATOMIC POSITIONS
 Si 0.00 0.00 0.00
 Si 0.25 0.25 0.25
K POINTS AUTOMATIC
4 4 4 1 1 1
```

Smaller than for simple total energy

2) Phonon calculation at  $\Gamma$  using ph.x

Input file si.phG.in of ph.x:



#### Output file si.phG.out:

```
Dielectric constant in cartesian axis
             13.99782
                          0.00000
                                       0.00000
             0.00000
                         13.99782
                                        0.00000
             0.0000
                          0.00000
                                       13.99782
     Effective charges (d Force / dE) in cartesian axis
      atom
                   Si
            -0.07434
                          0.00000
                                       0.00000
  Ex
             0.00000
                          -0.07434
                                       0.00000
  E_{\mathbf{V}}
             0.00000
                          0.00000
                                       -0.07434
  E_{Z}
                   Si
      atom
            -0.07434
                          0.00000
                                       0.00000
  Ex
             0.00000
                          -0.07434
                                       0.00000
  Εy
                          0.00000
  Ez
             0.00000
                                       -0.07434)
 Diagonalizing the dynamical matrix
         0.00000000
                     0.00000000
                                 0.00000000
 a = (
omega(1) =
                 0.081752 [THz] =
                                     2.726975 [cm-1]
                 0.081752
                         [THz] =
                                     2.726975
 omega(
                                             [cm-1]
 omega(
                 0.081752
                         [THz] =
                                     2.726975
                                            [cm-1]
 omega(4) =
                15.549026
                         [THz] =
                                   518.663131
 omega(
                15.549026
                         [THz] =
                                   518.663131 [cm-1]
                15,549026
                                   518.663131
 omega (
```

15.549026

0.000000 - 0.547867

0.000000 - 0.444530

0.000000 - 0.047270

0.547867

0.444530

0.047270

15.549026

15.549026 [THz] =

## Dynamical matrix file dyn.G:

omega(4)

omega(5) =

omega(6)

0.000000

0.000000

0.000000

-0.445151

0.445151

-0.535639

0.535639

0.122195

-0.122195

```
Diagonalizing the dynamical matrix
   a = (
           0.00000000
                       0.00000000
                                   0.000000000)
2.726975 [cm-1]
                   0.081752 [THz] =
   omega(1) =
                             0.000000 - 0.483206
                                               0.000000
  0.400413
           0.000000 - 0.325855
                                                         Acoustic
           0.000000 - 0.325855
  0.400413
                             0.000000 - 0.483206
                                               0.000000
                                                          modes
                   0.081752 [THz] =
                                        2.726975 [cm-1]
   omega(2) =
                                               0.000000
           0.000000 0.294894
                             0.000000
                                      0.280354
  0.578307
           0.000000 0.294894
  0.578307
                             0.000000
                                      0.280354
                                               0.000000
   omega(3) =
                   0.081752 [THz] =
                                        2.726975 [cm-1]
  0.072322
           0.000000 - 0.553946
                             0.000000
                                      0.433490
                                               0.000000
  0.072322
           0.000000 - 0.553946
                             0.000000
                                               0.000000
                                      0.433490
```

518.663131 [cm-1]

518.663131 [cm-1]

518.663131 [cm-1]

0.041012

0.124435

0.694863

0.000000 - 0.041012

0.000000 - 0.124435

0.000000 - 0.694863

0.000000

0.000000

0.000000

0.000000

0.000000

0.000000)

[THz] =

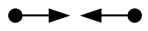
[THz] =

0.000000

0.000000

0.000000

**Optical** modes



Problems with the frequency of the acoustic phonon modes at  $\Gamma$  and with the effective charges.

Because of numerical inaccuracies the interatomic force constants and the effective charges do not strictly satisfy the following rules due to the translational invariance of the system:

Translational rules 
$$\sum_{\mathbf{L},j} C_{\alpha i,\beta j}(\mathbf{R_L}) = 0 \quad \forall \alpha, \beta \text{ and } i$$

$$\sum_{j} Z *_{j,\alpha \beta} = 0 \quad \forall \alpha, \beta$$

Reasons for numerical inaccuracies:

- Insufficiently accurate scf thresholds (in pw.x and/or in ph.x).
   DO NOT USE large thresholds in pw.x to save time!!
- XC energy is computed in real space. More problematic for GGA than in LDA. For US pseudopotentials it could require large ecutrho.
- k point sampling not accurate enough (in particular Z\* and ε require a very dense k-point sampling).

The *acoustic sum rules* can however be imposed after the phonon calculation.

To do this we use the code *dynmat.x* which imposes the acoustic sum rules on the elements of the dynamical matrix and dyagonalizes it. The input of *dynmat.x* in this case is:

```
%input
fildyn='dyn.G',
asr='simple',
/
File containing the dynamical matrix
A way to impose the acoustic sum rules
(if you put 'no' instead of 'simple'
the code does not impose the asr)
```

To submit the calculation:

```
prompt> dynmat.x < si.dynmat.in > si.dynmat.out
```

Dynmat.x produces the file *dynmat.out* which contains the new frequencies:

```
diagonalizing the dynamical matrix ...
        0.0000
                 0.0000
                            0.0000
a =
   omega(1) =
                 0.000000 [THz] =
                                    0.000000 [cm-1]
  0.000000 \quad 0.000000 \quad -0.707107 \quad 0.000000
                                          0.000000
                                                   0.000000
  0.00000
           0.000000 - 0.707107
                               0.00000
                                          0.000000
                                                   0.000000
   omega(2) = 0.000000 [THz] =
                                    0.000000 [cm-1]
 -0.707107 0.000000 0.000000 0.000000
                                         0.000000
                                                   0.00000
 -0.707107 0.000000 0.000000
                               0.00000
                                         0.000000
                                                   0.000000
                 0.000000 [THz] =
                                     0.000000 [cm-1]
   omega(3) =
  0.000000
           0.00000
                     0.000000 \quad 0.000000 \quad -0.707107
                                                   0.00000
                     0.00000
  0.00000
           0.00000
                               0.000000 - 0.707107
                                                   0.000000
  omega( 4) = 15.548811 [THz]
                                   518.655959 [cm-1]
  0.000000
           0.000000 0.707107 0.000000
                                                   0.000000
                                         0.000000
           0.000000 - 0.707107
                               0.00000
  0.00000
                                         0.000000
                                                   0.000000
  omega(5) = 15.548811 [THz]
                                   518.655959 [cm-1]
  0.000000
           0.00000
                     0.000000 \quad 0.000000 \quad -0.707107
                                                   0.000000
           0.000000
  0.00000
                      0.00000
                               0.00000
                                         0.707107
                                                   0.00000
   omega(6) = 15.548811 [THz]
                                   518.655959 [cm-1]
-0.707107 0.000000
                      0.00000
                               0.00000
                                         0.000000
                                                   0.000000
  0.707107
           0.000000
                      0.000000
                               0.000000
                                          0.000000
                                                   0.000000
```

Dynmat.x creates also the file *dynmat.axsf* which contains the phonon eigendisplacements. You can use xcrysden to visualize them:

```
prompt> xcrysden -axsf dynmat.axsf &
```

Polar materials in the  $\mathbf{q}=0$  limit: a macroscopic electric field appear as a consequence of long-rangeness of Coulomb interactions. Incompatible with Periodic Boundary Conditions! A non-analytic term must be added to force constants at  $\mathbf{q}=\mathbf{0}$ :

$${}^{na}\widetilde{C}_{st}^{\alpha\beta} = \frac{4\pi}{\Omega} \frac{(\mathbf{q} \cdot \mathbf{Z}^{\star}_{s})_{\alpha} (\mathbf{q} \cdot \mathbf{Z}^{\star}_{t})_{\beta}}{\mathbf{q} \cdot \boldsymbol{\epsilon}^{\infty} \cdot \mathbf{q}}$$

Effective charges  $\mathbb{Z}^*$  are related to polarization P induced by a lattice distortion:

$$Z_{s}^{\star \alpha \beta} = \Omega \frac{\partial \mathsf{P}_{\alpha}}{\partial u_{s}^{\beta}(\mathbf{q} = 0)}.$$

Dielectric tensor  $\epsilon_{\infty}^{\alpha\beta}$  are related to polarization induced by an electric field E:

$$\epsilon_{\infty}^{\alpha\beta} = \delta_{\alpha\beta} + 4\pi \left. \frac{\partial \mathsf{P}_{\alpha}}{\partial \mathsf{E}_{\beta}} \right|_{\mathsf{u}_{s}(\mathsf{q}=0)=0}.$$

All of the above can be calculated from (mixed) second derivatives of the energy.

9.33612

#### Part 2: Phonons in polar materials

Use the script run\_ph\_G\_AlAs.sh to calculate the phonons at Γ:

Dielectric constant in cartesian axis

0.00000

0.00000)

```
prompt> ./run_ph_G_AlAs.sh &
```

```
9.33612
                                            0.00000)
                0.00000
                              0.00000
                0.00000
                                            9.33612
        Effective charges (d Force / dE) in cartesian axis
         atom
                     Αl
                2.12370
                              0.00000
                                            0.00000)
    Ex
                0.00000
                              2.12370
                                            0.00000)
    EУ
                0.00000
                              0.00000
                                            2.12370
    Ez
                     As
         atom
                                            0.00000)
               -2.17270
                              0.00000
    Ex
                0.00000
                             -2.17270
                                            0.00000)
    Еy
                0.00000
                              0.00000
                                           -2.17270
    Ez
   Diagonalizing the dynamical matrix
           0.00000000
                        0.00000000
                                    0.000000000)
5.171595
   omega(1) =
                    0.155039 [THz] =
                                                 [cm-1]
                                         5.171595
   omega(2) =
                   0.155039
                            [THz] =
                                                 [cm-1]
   omega(3) =
                   0.155039
                            [THz] =
                                         5.171595 [cm-1]
   omega(4) =
                   11.220317
                            [THz] =
                                       374.271992
                                                 [cm-1]
                   11.220317
                                       374.271992
   omega(
                            [THz] =
                                                 [cm-1]
                   11.220317 [THz] =
   omega(6) =
```

No LO-TO splitting

To add the non-analytic term to the force constants and recalculate the frequencies we use dynmat.x.

In this case the input for dynmat.x is:

Re-calculate the phonons at  $\Gamma$ :

```
prompt> dynmat.x < alas.dynmat.in > alas.dynmat.out
```

#### Check the output file:

```
#
  mode
        [cm-1]
                    [THz]
                               TR
          0.00
                  0.0000
                           0.0000
   1
          0.00
                  0.0000
                           0.0000
          0.00
                  0.0000
                           0.0000
        374.24 11.2195
                           5.3672
                                      LO-TO
   5
        374.24 11.2195 5.3672
                                     splitting
        410.67 12.3115
                           5.3672
```

Infrared Intensities:

$$I_{IR}(\nu) = \sum_{\alpha} \left| \sum_{s\beta} Z_{s}^{\star \alpha \beta} U_{s}^{\beta}(\nu) \right|^{2}$$

can be calculated directly from effective charges and phonon displacement patterns.

#### Phonon calculation for $q \neq 0$

The procedure to calculate phonons for  $q \neq 0$  is the same as the one for a calculation at  $\Gamma$ .

Modify the script run\_ph\_G\_Si.sh and calculate the phonons at X = (0,0,1):

- 1) prompt> cp ../exercise1/run\_ph\_G\_Si.sh run\_ph\_X\_Si.sh
- 2) Use your favorite editor to remove epsil=.true. (that works only at  $\Gamma$ ), change the q point and the name of the dynamical matrix.
- 3) Execute the new script:

```
prompt> ./run_ph_X_Si.sh &
```

Notice that the output file for the phonon calculation is slightly different from the one at q=0. It is because for  $q\neq 0$  the code has to calculate also the wavefunctions  $\psi_{k+q}$  (in addition to the  $\psi_k$ 's in the scf calculation). Notice also that the symmetry of the *small group of* q is used (not the crystal symmetry).

## First step: Phonon calculation on a grid of q points

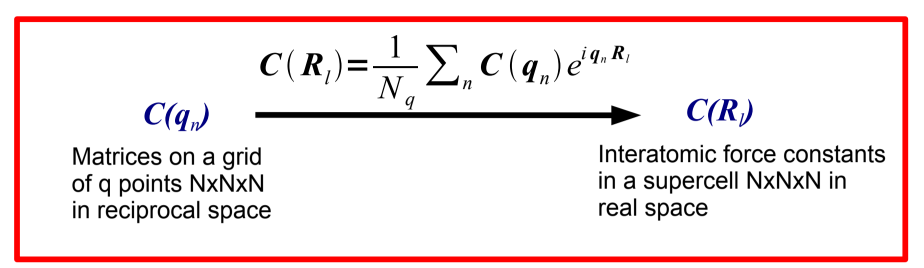
Modify the script run\_ph\_X\_Si.sh as described below.

- 1) prompt> cp run\_ph\_X\_Si.sh run\_ph\_Si.sh
- 2) The script should allow you to perform both scf and phonon calculations. The input file for the phonon calculations should be like this:

```
phonons on a grid & inputph prefix='si', epsil=.true., ldisp=.true. Option for the calculation on a grid fildyn='dyn', nq1=4, nq2=4, nq3=4, tr2_ph=1.0d-14, \mathbf{q}_{ijk} = \frac{i-1}{n_1}\mathbf{G}_1 + \frac{j-1}{n_2}\mathbf{G}_2 + \frac{k-1}{n_3}\mathbf{G}_3
```

The phonon code generates the files of the dynamical matrices on the specified grid of points. The files are dyn1, dyn2, ... dyn8. The file dyn0 contains the list of the inequivalent q points (8, in this case).

#### Second step: calculation of the interatomic force constants



We use the code q2r.x. The input file for q2r.x is:

```
Einput
fildyn='dyn',
zasr='simple',
flfrc='si.444.fc'

Output file of the interatomic force
constants
```

To perform the calculation:

```
prompt> q2r.x < q2r.in > q2r.out
```

Third step: phonons at a generic q' point using the IFC

$$C(R_l) = \sum_{l} C(R_l) e^{-iq'R_l}$$
Interatomic force constants in a supercell NxNxN in real space

Fourier interpolation

$$C(q') = \sum_{l} C(R_l) e^{-iq'R_l}$$
Dynamical matrix at a generic q' point

We use the code matdyn.x. The input file for matdyn.x is:

```
&input
    asr='simple',
    amass(1)=28.0855,
    flfrc='si.444.fc',
    flfrq='si.freq'

0.300000

0.200000

0.050000

List of q points
```

To perform the calculation:

```
prompt> matdyn.x < matdyn_ex.in > matdyn_ex.out
```

## Si phonon dispersion

Using the script run\_dispersion\_Si.sh you can calculate and generate a plot of the phonon dispersion:

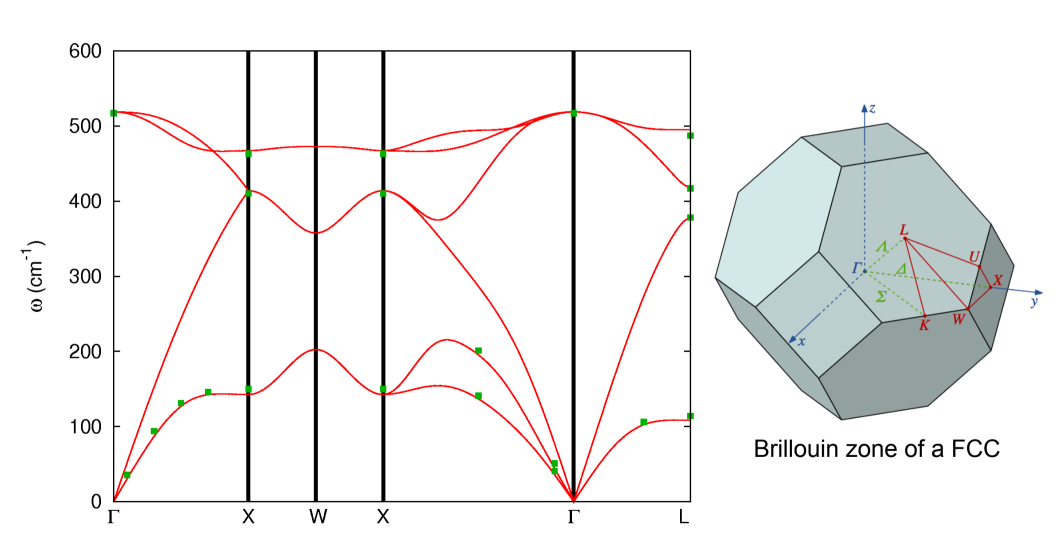
```
prompt> ./run_dispersion_Si.sh &
```

The script includes a *matdyn.x* calculation on a list of q points (along particular lines in the Brillouin zone) and a *plotband.x* post-processing calculation that generates a user-friendly file of the frequencies and a postscript figure.

Using a gnuplot script (*plot\_dispersion.gnu*) you can plot the theoretical results together with some experimental data (exp.dat):

```
prompt> gnuplot plot_dispersion.gnu
```

Si phonon dispersion (file gnuplot\_dispersion.eps)



#### Si vibrational density of states

The code matdyn.x can also be used to calculate the phonon density of states. In this case the input file should be like this:

```
&input
    asr='simple',
    amass(1)=28.0855,
    flfrc='si.444.fc',
    flfrq='si.dos.freq'
    dos=.true.,
    fldos='si.dos'
    deltaE=1.d0,
    nk1=4, nk2=4, nk3=4,

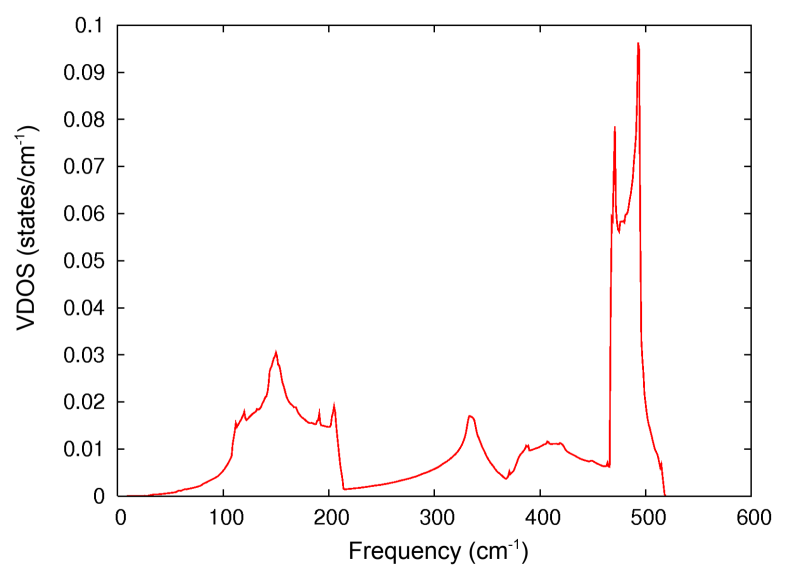
    Grid of q points for the DOS
    calculation
```

Modify the matdyn.x input file and perform the calculation. You can use the gnuplot script plot dos.gnu to generate a picture.

```
prompt> gnuplot plot_dos.gnu
```

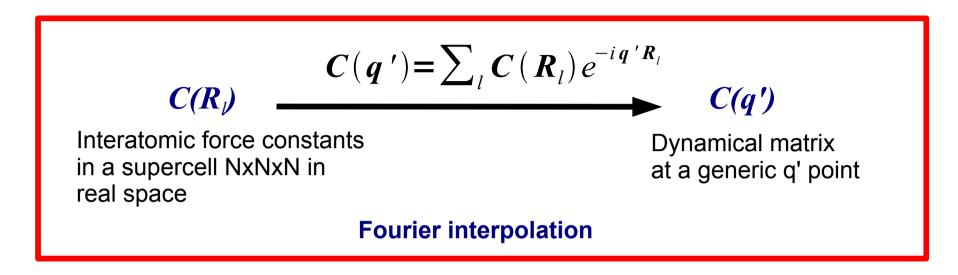
Calculate the phonon density of states using different grid of q points. Which grid gives a converged result?

## Si vibrational density of states



Notice that the VDOS is normalized to  $3\ensuremath{N_{\text{at}}}$  , where  $\ensuremath{N_{\text{at}}}$  is the number of

# **Exercise 4: checking the Fourier interpolation**



We need to check if the Fourier interpolation works properly. To do that we can calculate:

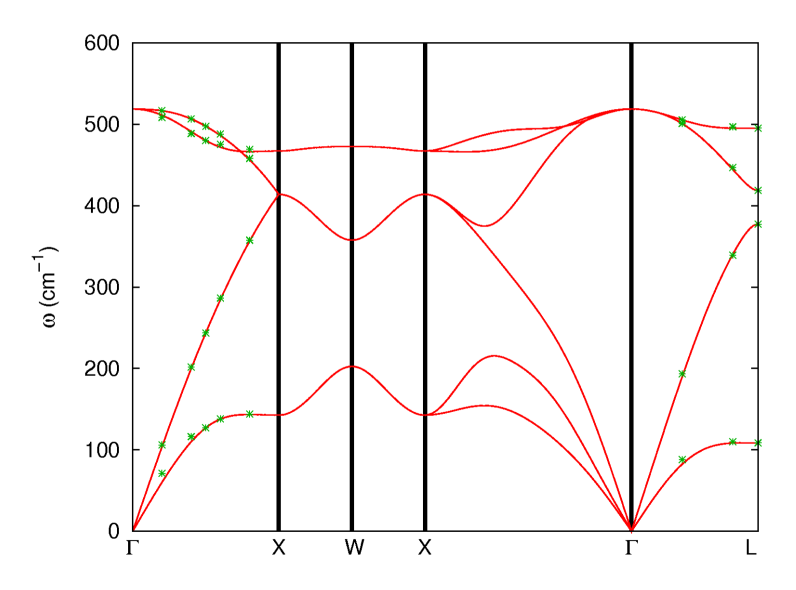
- 1) the phonon frequencies at a certain q point (not belonging to the grid on which we calculated the dynamical matrices) using matdyn.x and our file of the interatomic force constants (from exercise3).
- 2) the phonon frequencies at the same point using ph.x (i.e. performing the *ab initio* calculation).

and compare the results.

You can run a ph.x calculation modifying the script of exercise3 and perform a matdyn.x calculation as in exercise3.

## **Exercise 4: checking the Fourier interpolation**

You can use the script run\_check\_Si.sh and use plot\_check.gnu Where the comparison is done automatically for a few q points.



# **Exercise 4: checking the Fourier interpolation**

The Fourier interpolation works if the interatomic force constants are known on a sufficiently large supercell (i.e, large enough grid of q points in the phonon calculation).

There are cases in which the interatomic force constants are long range and the Fourier interpolation does not work properly:

- When there are Kohn anomalies in metals. In this case the dynamical matrices are not a smooth function of q and the interatomic force constants are long range.
- 2) In polar insulators where the atomic displacements generate long range electrostatic interactions and the dynamical matrix is non analytic for q → 0. As we have seen, this case, however, can be dealt with by calculating the Born effective charges and the dielectric tensor of the material.

Extra slides...

## Phonons and thermo-mechanical properties

Phonons are determined by this secular equation:

$$\sum_{t,\beta} \left( \widetilde{C}_{st}^{\alpha\beta}(\mathbf{q}) - M_s \omega^2(\mathbf{q}) \delta_{st} \delta_{\alpha\beta} \right) U_t^{\beta}(\mathbf{q}) = 0$$

Free energy in quasi-harmonic approximation:

$$\begin{split} F(\{a_i\},T) &= E(\{a_i\}) + F_{vibr}(\{a_i\},T) \\ &= E(\{a_i\}) + \sum_{q,j} \frac{\hbar \, \omega_{q,j}(\{a_i\})}{2} + k_{\scriptscriptstyle B} T \sum_{q,j} \ln \left(1 - \exp\left(-\frac{\hbar \, \omega_{q,j}(\{a_i\})}{k_{\scriptscriptstyle B} T}\right)\right) \\ & \qquad \qquad \qquad \\ \hline \alpha_i &= \frac{1}{a_i} \frac{\partial a_i}{\partial T} \end{split} \quad \text{Thermal expansion coefficient} \end{split}$$

And also: heat capacity, temperature dependence of the elastic constants....

# **Density functional perturbation theory (1)**

Let us assume that the external potential depends on some parameter  $\lambda$ 

$$V_{\lambda}(\mathbf{r}) \simeq V(\mathbf{r}) + \lambda \frac{\partial V(\mathbf{r})}{\partial \lambda} + \frac{1}{2} \lambda^2 \frac{\partial^2 V(\mathbf{r})}{\partial \lambda^2} + \dots$$

(all derivatives calculated at  $\lambda = 0$ ) and expand the charge density

$$n_{\lambda}(\mathbf{r}) \simeq n(\mathbf{r}) + \lambda \frac{\partial n(\mathbf{r})}{\partial \lambda} + \frac{1}{2} \lambda^2 \frac{\partial^2 n(\mathbf{r})}{\partial \lambda^2} + \dots$$

and the energy functional into powers of  $\lambda$ :

$$E_{\lambda} \simeq E + \lambda \frac{\partial E}{\partial \lambda} + \frac{1}{2} \lambda^2 \frac{\partial^2 E}{\partial \lambda^2} + \dots$$

The first-order derivative  $\partial E/\partial \lambda$  does not depend on any derivative of  $n(\mathbf{r})$  (Hellmann-Feynman theorem):

$$\frac{\partial E}{\partial \lambda} = \int n(\mathbf{r}) \frac{\partial V(\mathbf{r})}{\partial \lambda} d\mathbf{r}$$

# **Density functional perturbation theory (2)**

The second-order derivative  $\partial^2 E/\partial \lambda^2$  depends on the first-order derivative of the charge density,  $\partial n(\mathbf{r})/\partial \lambda$ :

$$\frac{\partial^2 E}{\partial \lambda^2} = \int \frac{\partial V(\mathbf{r})}{\partial \lambda} \frac{\partial n(\mathbf{r})}{\partial \lambda} d\mathbf{r} + \int n(\mathbf{r}) \frac{\partial^2 V(\mathbf{r})}{\partial \lambda^2} d\mathbf{r}$$

The result can be generalized to mixed derivatives:

$$\frac{\partial^2 E}{\partial \lambda \partial \mu} = \int \frac{\partial V(\mathbf{r})}{\partial \lambda} \frac{\partial n(\mathbf{r})}{\partial \mu} d\mathbf{r} + \int n(\mathbf{r}) \frac{\partial^2 V(\mathbf{r})}{\partial \lambda \partial \mu} d\mathbf{r}$$

(the order of derivatives can be exchanged)

In general, the (2n+1)-th derivative of energy depends only on derivatives up to order n of the charge density ((2n+1) theorem).

# **Density functional perturbation theory (3)**

The first-order derivative of the charge density can be obtained as in traditional linearresponse theory, by applying perturbation theory to the KS Hamiltonian:

$$\frac{\partial \psi_{v}(\mathbf{r})}{\partial \lambda} = \sum_{c} \psi_{c}(\mathbf{r}) \frac{1}{\epsilon_{v} - \epsilon_{c}} \langle \psi_{c} | \frac{\partial V_{KS}}{\partial \lambda} | \psi_{v} \rangle = \frac{1}{\epsilon_{v} - H_{KS}} P_{c} \frac{\partial V_{KS}(\mathbf{r})}{\partial \lambda} \psi_{v}(\mathbf{r})$$

(v denotes occupied KS states, c empty states,  $P_c$  is the projector over empty states). The self-consistent potential response  $\partial V_{KS}/\partial \lambda$ :

$$\frac{\partial V_{KS}(\mathbf{r})}{\partial \lambda} = \frac{\partial V(\mathbf{r})}{\partial \lambda} + \int \frac{1}{|\mathbf{r} - \mathbf{r}'|} \frac{\partial n(\mathbf{r}')}{\partial \lambda} d\mathbf{r}' + \int \frac{\delta V_{xc}(\mathbf{r})}{\delta n(\mathbf{r}')} \frac{\partial n(\mathbf{r}')}{\partial \lambda} d\mathbf{r}'$$

depends on the first-order variation  $\partial n(\mathbf{r})/\partial \lambda$  of the charge density:

$$\frac{\partial n(\mathbf{r})}{\partial \lambda} = 2 \operatorname{Re} \sum_{v} \psi_{v}^{*}(\mathbf{r}) \frac{\partial \psi_{v}(\mathbf{r})}{\partial \lambda}$$

The quantities  $\partial \psi_v(\mathbf{r})/\partial \lambda$  and all other needed quantities are thus determined by the solution of a self-consistent set of *linear equations*.

## Frozen phonons vs DFPT

Frozen phonons is an older and alternative way to calculate phonons. The monochromatic perturbation is frozen in with a finite amplitude in the system, which is described by a supercell having **q** as reciprocal lattice vector.

Fourier transform of force constants at **q** are calculated from *finite differences of forces* induced on all the atoms of the supercell by the monochromatic perturbation.

#### Advantages:

straightforward to implement

#### Disadvantages:

• limited to small supercells, i.e.  $\mathbf{q} = \mathbf{G}/n$ , where  $\mathbf{G}$  is a reciprocal lattice vector of the original cell, n=2,3,4,..., but in any case a small number.

Note that this is *not* the algorithm used by Quantum-Espresso! What if we want the entire dispersions for all **q**-vectors in the Brillouin Zone?

## Calculation of IR and Raman intensities

Infrared Intensities:

$$I_{IR}(\nu) = \sum_{\alpha} \left| \sum_{s\beta} Z_{s}^{\star \alpha \beta} U_{s}^{\beta}(\nu) \right|^{2} \qquad Z_{s}^{\star \alpha \beta} = \Omega \frac{\partial P_{\alpha}}{\partial u_{s}^{\beta}(\mathbf{q} = 0)}$$

can be calculated directly from effective charges and phonon displacement patterns.

Non-resonant Raman intensities:

$$I_{\text{Stokes}}(\nu) \propto \frac{(\omega_i - \omega_\nu)^4}{\omega_\nu} r_{\alpha\beta}(\nu), \qquad r_{\alpha\beta}(\nu) = \left| \frac{\partial \chi_{\alpha\beta}}{\partial U(\nu)} \right|^2$$

where  $\chi$  is the electric polarizability of the system.

Raman coefficients are third-order derivatives of the energy that can be calculated in various ways. The most convenient way is to use second-order response to an electric field: M.Lazzeri and F.Mauri, Phys. Rev. Lett. 90, 036401 (2003).