

LetzElPhC Documentation

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Chapter 1

About the Code

LetzElPhC is a C code designed to compute electron-phonon coupling matrix elements from the outputs of standard Density Functional Theory (DFT) and Density Functional Perturbation Theory (DFPT) calculations. Currently, it only supports the Quantum Espresso code, with long plans to support the Abinit code. The main objective of this project is to facilitate electron-phonon related calculations within the YAMBO code (version 5.2 and above), and it works only with norm-conserving pseudo-potentials. The code is released under the MIT license and hosted on GitHub [\[link\]](#).

1.1 Main Features

- Utilizes full crystal symmetries, ensuring compatibility with the YAMBO code, without encountering phase issues.
- Implements multiple levels of parallelization, including OpenMP, plane-wave, k-point, and q-point parallelization.
- Utilizes fully parallel IO via parallel NetCDF-4/HDF5 libraries.
- Highly portable. The code can be compiled on various CPU architectures and operating systems with minimal to no changes in the source code.

Chapter 2

Installing the Code

2.1 Mandatory Requirements

- GNU Make
- C99 compiler with complex number support, such as GCC, Clang, ICC, AMD C-Compiler, MinGW (for Windows), PGI, or Arm C compilers.
- MPI implementation supporting at least MPI-standard 2.1 standard, such as Open-MPI, MPICH and its variants, Intel MPI compiler, or Microsoft MPI (for Windows).
- FFTW-3 or Intel-MKL.
- HDF5 and NETCDF-4 libraries with Parallel IO support (compiled with MPI).
- A BLAS library, such as OpenBLAS, BLIS, Intel-MKL, or Atlas.

2.2 Installation Process

LetzElPhC employs a standard make build system. Sample make files are available in the [sample_config](#) directory. Copy one to the [src](#) directory and rename it as [make.inc](#). Navigate to the [src](#) directory and edit [make.inc](#) according to your requirements. Then, in the same directory, execute the following commands:

```
1 $ make
2 ##### To compile the code in parallel, use the -j option
3 $ make -j n
4 ##### where n is the number of processes.
```

Upon successful compilation, you should find the *lelphc* executable located in the *src* directory. If you encounter difficulties in locating the required libraries, go to the YAMBO code installation directory and open the *report* file in the *config* directory, which lists all necessary libraries and include paths.

Below are the list of variables in the *make.inc* file with explanations.

```

1 CC                := mpicc
2 ##### MPI C compiler mpicc/mpicc (for intel),
3 CFLAGS            := -O3
4 ##### -O3 is to activate compiler optimizations
5 LD_FLAGS          :=
6 ##### use this to pass any flags to linker
7
8 ##### ***** OPENMP BUILD ***
9 ##### If you wish to build the code with openmp support
10 ##### uncomment the below line
11 # OPENMP_FLAGS     := -DELPH_OMP_PARALLEL_BUILD
12 ##### Additionally, you need to add openmp compiler flag to
13 ##### CFLAGS and LD_FLAGS.
14 ##### Also uncomment the below two lines
15 # CFLAGS           += -fopenmp ## use -qopenmp for intel
16 # LD_FLAGS         += -fopenmp ## use -qopenmp for intel
17
18 ##### FFTW3 include and libs (see FFT flag in yambo config/report)
19 FFTW_INC           := -I/opt/homebrew/include
20 FFTW3_LIB          := -L/opt/homebrew/lib -lfftw3_threads -
    lfftw3f -lfftw3f_omp -lfftw3_omp -lfftw3
21 ##### Note if using FFTW
22 ##### Yambo uses double precision FFTW regardless of the precision
    with which Yambo is built. In contrast, you need to link single
    (double) precision FFTW for single (double) precision LetzElPhC.
    please refer to https://www.fftw.org/fftw3\_doc/Precision.html .
    Also you refer to https://www.fftw.org/fftw3\_doc/
    Multi_002dthreaded-FFTW.html if compiling with openmp support.
23
24 ##### If you want to compile the code in double precession,
    uncomment the below
25 #CFLAGS            += -DCOMPILER_ELPH_DOUBLE
26
27 ##### Blas and lapack libs (see BLAS and LAPACK flag in yambo config
    /report)
28 BLAS_LIB           := -L/opt/homebrew/opt/openblas/lib -lopenblas
29 ##### you need to add both blas and lapack libs for ex : -lblas -
    llapack
30
31 ##### netcdf libs and include
32 ##### (see NETCDF flag in yambo config/report)
33 NETCDF_INC         := -I/Users/murali/software/core/include
34 NETCDF_LIB         := -L/Users/murali/software/core/lib -lnetcdf
35
36 ##### hdf5 lib (see HDF5 flag in yambo config/report)
37 HDF5_LIB           := -L/opt/homebrew/lib -lhdf5
38
39 ##### incase if you want to add additional include dir and libs

```

```
40 INC_DIRS      :=
41 LIBS          :=
42
43
44 ##### Notes Extra CFLAGS
45 ### add -DCOMPILER_DOUBLE if you want to compile the code in
    double precision
46 ### if you are using yambo <= 5.1.2, you need to add "-
    DYAMBO_LT_5_1" to cflags
47 ### for openmp use -DELPH_OMP_PARALLEL_BUILD in CFLAGS and set -
    fopenmp in LD_FLAGS and CFLAGS
```

Chapter 3

Running the Code

3.1 Running DFT and DFPT (Step 0)

Before using LetzElPhC, ensure you have obtained the following quantities:

- Kohn-Sham wavefunctions (obtained from a non-self-consistent calculation after obtaining the ground state density of the system).
- Phonon eigenvectors and perturbed Hartree and Exchange potentials due to phonon modes (obtained from a DFPT run after finding the ground state).

With the Quantum Espresso code, follow these steps (an example is provided in [examples/qe/silicon](#)):

- Perform a self-consistent field (SCF) calculation to obtain the ground state.
- Perform a DFPT calculation using the **ph.x** executable to obtain dynamical matrices and changes in potentials on a uniform q-point grid.
- Perform a non-self-consistent field (NSCF) calculation to obtain the wavefunctions on a uniform k-point grid. The k-grid and q-grid of phonons must be commensurate.

Note: Set the *dvscf* flag in the **ph.x** input to save the change in potentials. If you forget to set this variable, you have to rerun the entire calculation. Additionally, make sure that the q-grid is commensurate with the k-grid used in the NSCF calculation (Although, the choice of kgrid to converge the SCF calculation is irrelevant). Once these steps are completed successfully, go to the NSCF folder and enter the *prefix.save* directory, where the wavefunctions are stored. Then, execute *p2y* followed by *yambo* to generate the **SAVE** folder:

```

1 $ p2y
2 #### Generates the YAMBO SAVE directory
3 $ yambo
4 #### Further processing creates additional files

```

Upon successful completion of these steps, we are ready to use LetzElPhC.

3.2 Running the Preprocessor (Step 1)

Once the **SAVE** directory is obtained, we need to create the **ph_save** folder. Navigate to the phonon calculation directory and run the preprocessor with the **-pp** flag:

```

1 $ cd /path/to/phonon calculation directory
2 #### Run the preprocessor
3 $ lelphc -pp --code=qe -F PH.X_input_file
4 #### Where PH.X_input_file is the input file of ph.x code used for
   computing phonons

```

Upon successful execution, the **ph_save** directory will be created, containing all necessary files. If you wish to create soft symlinks instead of copying these files, set the following environment variable (works only on linux):

```

1 $ export ELPH_COPY_CMD="ln -sr"

```

and then run the preprocessor. Additionally, the name of the **ph_save** directory can be changed using another environment variable:

```

1 $ export ELPH_SAVE_DIR=ph_save_name_you_want

```

Remarks:

- The new format XML dynamical matrix files are currently not supported.
- The preprocessor does not directly work with the image parallelization. In case if you use image parallelization,, copy all `prefix.q_*` in each `_ph*` folder (other than `_ph0`) to the `_ph0` directory and run the preprocessor.

3.3 Performing the ELPH Calculation (Final Step)

Once both the **SAVE** and **ph_save** folders are created, the ELPH calculation can be executed. Run the following command with the LetzElPhC input file in any directory where you wish to perform the calculation:


```

1 $ mpirun -n 4 lelphc -F LetzElPhC_input_file
2 ## Here, we are using 4 MPI processes.

```

A detailed description of the input file is provided below:

```

1  nkpool          = 1
2  # k point parallelization (number of kpools)
3
4  nqpool          = 1
5  # q point parallelization (number of qpools)
6
7  ## note (nkpool*nqpool) must divide total number of cpus.
8
9  ## For example, if you run the code on 12 processes,
10 ## and set nkpool = 3 and nqpool = 2
11 ## then, we have 2 sets of cpus working subset of qpoints
12 ## with each group having 3 sub groups which that work on
13 ## subset of kpoints. So in total, we have 6 subgroups, each
14 ## having 2 cpus that distribute plane waves
15
16 ## {1,2,3,4,5,6,7,8,9,10,11,12} (total cpus)
17 ## -----|-----
18 ## |      divided in to 2 qpools      |
19 ## (qpool 1) {1,2,3,4,5,6}      (qpool 2) {7,8,9,10,11,12}
20 ## -----|-----|-----|-----
21 ## |      |      |      |      |      |
22 ## kp1    kp2    kp3  kp1    kp2    kp3
23 ## where kp1 are kpools each containg 2
24 ## cpus work on subset of plane waves
25
26 start_bnd        = 1
27 # starting band to consider in elph calculation
28
29 end_bnd          = 40
30 # last band to consider in elph calculation
31
32 save_dir          = SAVE
33 # SAVE dir you created with yambo
34
35 ph_save_dir       = ph_save
36 # ph_save directory that was created with preprocessor
37
38 kernel           = dfpt
39 ## type of kernel, only dfpt supported right Now
40
41 convention        = standard
42 # standard/yambo, If standard (default)
43 # <k+q|dV|k> is computed. if yambo, <k|dV|k-q> is outputed
44
45 ### ##, !, ; are considered as comments

```

Bibliography