



# PWneb User's Guide (v. 7.2)

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## 1 Introduction

This guide covers the usage of `PWneb`, version 7.2: an open-source package for the calculation of energy barriers and reaction pathway using the Nudged Elastic Band (NEB) method.

This guide assumes that you know the physics that `PWneb` describes and the methods it implements. It also assumes that you have already installed, or know how to install, `QUANTUM ESPRESSO`. If not, please read the general User's Guide for `QUANTUM ESPRESSO`, found in subdirectory `Doc/` of the main `QUANTUM ESPRESSO` directory, or consult the web site: <http://www.quantum-espresso.org>.

`PWneb` is part of the `QUANTUM ESPRESSO` distribution and uses the `PWscf` package as electronic-structure computing tools ("engine"). It is however written in a modular way and could be adapted to use other codes as "engine". Since v.4.3 the NEB calculation is performed by a separate executable `neb.x` and no longer by `pw.x`. Also note that NEB with Car-Parrinello molecular dynamics is no longer implemented since v.4.3.

## 2 People and terms of use

The current maintainers of `PWneb` are Layla Martin-Samos, Paolo Giannozzi, Stefano de Gironcoli. The original QUANTUM ESPRESSO implementation of NEB was written by Carlo Sbraccia.

`PWneb` is free software, released under the GNU General Public License.

See <http://www.gnu.org/licenses/old-licenses/gpl-2.0.txt>, or the file `License` in the distribution).

We shall greatly appreciate if scientific work done using the QUANTUM ESPRESSO distribution will contain an acknowledgment to the following references:

P. Giannozzi, S. Baroni, N. Bonini, M. Calandra, R. Car, C. Cavazzoni, D. Ceresoli, G. L. Chiarotti, M. Cococcioni, I. Dabo, A. Dal Corso, S. Fabris, G. Fratesi, S. de Gironcoli, R. Gebauer, U. Gerstmann, C. Gougoussis, A. Kokalj, M. Lazzeri, L. Martin-Samos, N. Marzari, F. Mauri, R. Mazzarello, S. Paolini, A. Pasquarello, L. Paulatto, C. Sbraccia, S. Scandolo, G. Sclauzero, A. P. Seitsonen, A. Smogunov, P. Umari, R. M. Wentzcovitch, *J.Phys.: Condens.Matter* 21, 395502 (2009)

and

P. Giannozzi, O. Andreussi, T. Brumme, O. Bunau, M. Buongiorno Nardelli, M. Calandra, R. Car, C. Cavazzoni, D. Ceresoli, M. Cococcioni, N. Colonna, I. Carnimeo, A. Dal Corso, S. de Gironcoli, P. Delugas, R. A. DiStasio Jr, A. Ferretti, A. Floris, G. Fratesi, G. Fugallo, R. Gebauer, U. Gerstmann, F. Giustino, T. Gorni, J Jia, M. Kawamura, H.-Y. Ko, A. Kokalj, E. Küçükbenli, M. Lazzeri, M. Marsili, N. Marzari, F. Mauri, N. L. Nguyen, H.-V. Nguyen, A. Otero-de-la-Roza, L. Paulatto, S. Poncé, D. Rocca, R. Sabatini, B. Santra, M. Schlipf, A. P. Seitsonen, A. Smogunov, I. Timrov, T. Thonhauser, P. Umari, N. Vast, X. Wu, S. Baroni, *J.Phys.: Condens.Matter* 29, 465901 (2017)

Users of the GPU-enabled version should also cite the following paper:

P. Giannozzi, O. Baseggio, P. Bonfà, D. Brunato, R. Car, I. Carnimeo, C. Cavazzoni, S. de Gironcoli, P. Delugas, F. Ferrari Ruffino, A. Ferretti, N. Marzari, I. Timrov, A. Urru, S. Baroni, *J. Chem. Phys.* 152, 154105 (2020)

Note the form QUANTUM ESPRESSO for textual citations of the code. Please also see package-specific documentation for further recommended citations. Pseudopotentials should be cited as (for instance)

[ ] We used the pseudopotentials `C.pbe-rrjkus.UPF` and `O.pbe-vbc.UPF` from <http://www.quantum-espresso.org>.

## 3 Compilation

`PWneb` is a package of QUANTUM ESPRESSO and requires package `PWscf` for compilation. For instruction on how to download and compile QUANTUM ESPRESSO, please refer to the general Users' Guide, available in file `Doc/user_guide.pdf` under the main QUANTUM ESPRESSO directory, or in web site <http://www.quantum-espresso.org>.

Once QUANTUM ESPRESSO is correctly configured, PWneb can be automatically downloaded, unpacked and compiled by just typing `make neb`, from the main QUANTUM ESPRESSO directory. `make neb` will produce the following codes in NEB/src:

- `neb.x`: calculates reaction barriers and pathways using NEB.
- `path_interpolation.x`: generates a reaction path (a set of points in the configuration space of the atomic system, called “images”), by interpolating the supplied path. The new path can have a different number of images than the old one and the initial and final images of the new path can differ from the original ones. The utility `path_interpolation.sh` in the `tools/` directory shows how to use the code.

Symlinks to executable programs will be placed in the `bin/` subdirectory of the main QUANTUM ESPRESSO directory.

### 3.1 Running examples

As a final check that compilation was successful, you may want to run the examples (presently only one). See the general documentation for instructions on how to run the examples.

Go to `NEB/examples/example01` and execute:

```
./run_example
```

This will create a subdirectory `results/` containing the input and output files generated by the calculation.

The `reference/` subdirectory contains verified output files, that you can check your results against. The exact numbers depend upon the hardware, software stack, execution mode: tiny differences are not a problem, but a plain diff of your results against the reference data doesn’t work, or at least, it requires human inspection.

## 4 Parallelism

The PWneb code is interfaced to PWscf, which is used as computational engine for total energies and forces. It can therefore take advantage from the two parallelization paradigms currently implemented in QUANTUM ESPRESSO, namely Message Passing Interface (MPI) and OpenMP threads, and exploit all PWscf-specific parallelization options. For a detailed information about parallelization in QUANTUM ESPRESSO, please refer to the general documentation.

In addition, PWneb makes several independent evaluations of energy and forces at each step of the path optimization: one per “image”, that is, a point in the path, corresponding to a set of atomic positions. It is thus possible and often convenient to distribute images among processors, using the “image” parallelization, as described in the general documentation. The number of image groups is specified using the option `-ni N` (or, equivalently, `-nimage N`) after the executable name (e.g., `neb.x`) in the command line. The default is a single image group (no image parallelization)

Images are loosely coupled calculations: processors belonging to different image groups communicate only once in a while, whereas processors within the same image group are tightly coupled and communications are more significant (please refer to the user guide of PWscf).

## 5 Using PWneb

NEB calculations with `neb.x` can be started in two different ways:

1. by reading a single input file, specified with the command line option `-i` (or `-in`, or `-inp`);
2. by specifying the number  $N$  of images with the command line option `-input_images N`, and providing the input data for PWneb in a file named `neb.dat` and for the PWscf engine in the files `pw_X.in` ( $X = 1, \dots, N$ , see also below).

In the first case, the input file contains keywords (described here below) that enable the code to distinguish between parts of the input containing NEB-specific parameters and parts containing instructions for the computational engine (only PW is currently supported).

**N.B.:** the `neb.x` code does not read from standard input, so that input redirection (e.g., `neb.x < neb.in ...`) cannot be used.

The general structure of the file to be parsed should be as follows:

```
BEGIN
BEGIN_PATH_INPUT
~... neb specific namelists and cards
END_PATH_INPUT
BEGIN_ENGINE_INPUT
~...pw specific namelists and cards
BEGIN_POSITIONS
FIRST_IMAGE
~...pw ATOMIC_POSITIONS card
INTERMEDIATE_IMAGE
~...pw ATOMIC_POSITIONS card
LAST_IMAGE
~...pw ATOMIC_POSITIONS card
END_POSITIONS
~... other pw specific cards
END_ENGINE_INPUT
END
```

After the parsing is completed, several files are generated by PWneb, more specifically: `neb.dat`, with NEB-related input data, and a set of input files in the PWscf format, `pw_1.in`, ..., `pw_N.in`, one for each set of atomic position (image) found in the original input file. For the second case, the `neb.dat` file and all `pw_X.in` should be already present in the directory where the code is started. A detailed description of all NEB-specific input variables is contained in the input description files `Doc/INPUT_NEB.*`, while for the PWscf engine all the options of a `scf` calculation apply (see `PW/Doc/INPUT_PW.*` and `example01` in the `NEB/examples` directory).

A NEB calculation will produce a number of output files containing additional information on the minimum-energy path. The following files are created in the directory where the code is started:

`prefix.dat` is a three-column file containing the position of each image on the reaction coordinate (arb. units), its energy in eV relative to the energy of the first image and the residual error for the image in  $\text{eV}/a_0$ .

`prefix.int` contains an interpolation of the path energy profile that pass exactly through each image; it is computed using both the image energies and their derivatives

`prefix.path` information used by QUANTUM ESPRESSO to restart a path calculation, its format depends on the input details and is undocumented

`prefix.axsf` atomic positions of all path images in the XCrySDen animation format: to visualize it, use `xcrysdn --axsf prefix.axsf`

`prefix.xyz` atomic positions of all path images in the generic xyz format, used by many quantum-chemistry softwares

`prefix.crd` path information in the input format used by `pw.x`, suitable for a manual restart of the calculation

where `prefix` is the `PWscf` variable specified in the input. The more verbose output from the `PWscf` engine is not printed on the standard output, but is redirected into a file stored in the image-specific temporary directories (e.g. `outdir/prefix_1/PW.out` for the first image, etc.).

NEB calculations are a bit tricky in general and require extreme care to be setup correctly. Sometimes it can easily take hundreds of iterations for them to converge, depending on the number of atoms and of images. Here you can find some advice (courtesy of Lorenzo Paulatto):

1. Don't use Climbing Image (CI) from the beginning. It makes convergence slower, especially if the special image changes during the convergence process (this may happen if `CI_scheme='auto'` and if it does it may mess up everything). Converge your calculation, then restart from the last configuration with CI option enabled (note that this will *increase* the barrier).
2. Carefully choose the initial path. If you ask the code to use more images than those you have supplied on input, the code will make a linear interpolation of the atomic positions between consecutive input images. You can visualize the `.axsf` file with XCrySDen as an animation: take some time to check if any atoms overlap or get very close in some of the new images (in that case you will have to supply intermediate images). Remember that QUANTUM ESPRESSO assumes continuity between two consecutive input images to initialize the path. In other words, periodic images are not used by default, so that an unwanted path could result if some atom crosses the border of the unit cell and it is refolded in the unit cell in the input image. The problem can be solved by activating the `minimum_image` option, which chooses an appropriate periodic replica of any atom that moves by more than half the size of the unit cell between two consecutive input images. If this does not work either, you may have to manually translate an atom by one or more unit cell base vectors in order to have a meaningful initial path.
3. Try to start the NEB process with most atomic positions fixed, in order to converge the more "problematic" ones, before leaving all atoms move.
4. Especially for larger systems, you can start NEB with lower accuracy (less k-points, lower cutoff) and then increase it when it has converged to refine your calculation.
5. Use the Broyden algorithm instead of the default one: it is a bit more fragile, but it removes the problem of "oscillations" in the calculated activation energies. If these oscillations persist, and you cannot afford more images, focus to a smaller problem, decompose it into pieces.

6. A gross estimate of the required number of iterations is (number of images) \* (number of atoms) \* 3. Atoms that do not move should not be counted. It may take half that many iterations, or twice as many, but more or less that's the order of magnitude, unless one starts from a very good or very bad initial guess.

The code `path_int.x` is a tool to generate a new path (what is actually generated is the restart file) starting from an old one through interpolation (cubic splines). The new path can be discretized with a different number of images (this is its main purpose), images are equispaced and the interpolation can be also performed on a subsection of the old path. The input file needed by `path_int.x` can be easily set up with the help of the self-explanatory `path_interpolation.sh` shell script in the `NEB/tools` folder.

## 6 Performances

`PWneb` requires roughly the time and memory needed for a single SCF calculation, times `num_of_images`, times the number of NEB iterations needed to reach convergence. We refer the reader to the `PW user_guide` for more information.

## 7 Troubleshooting

Almost all problems in `PWneb` arise from incorrect input data and result in error stops. Error messages should be self-explanatory, but unfortunately this is not always true. If the code issues a warning messages and continues, pay attention to it but do not assume that something is necessarily wrong in your calculation: most warning messages signal harmless problems.