CAP optimization manual

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1 Introduction

This algorithm aims to determine the optimal Complex Absorbing Potential for a specific system, so that it produces a transmission (and sometimes reflection) as close as possible from the exact system:

$$T_{CAP}(E,k) = T_{obj}(E,k)$$

It is done by minimizing the square of the difference between the exact and the CAP-computed transmission (and sometimes reflection), with various k processing methods that can be chosen by the user. The python module *scipy.optimize* is used for that purpose.

2 Complex absorbing potential approach

In the Complex Absorbing Potential (CAP) approximation, the environment is represented by a finite part close to the system.

Under the NEGF (Non-Equilibrium Green Function) formalism, the Hamiltonian of the system is written as following:

$$H' = \begin{bmatrix} H_L + W_L & V_L^{\dagger} & 0 \\ V_L & H_D & V_R^{\dagger} \\ 0 & V_R & H_R + W_R \end{bmatrix}$$

Where $W_{L/R}$ is the CAP matrix. We write

$$W_{\mu,v} = \langle \mu | W | v \rangle = v_{\mu\nu} - i w_{\mu\nu} , \ v_{\mu\nu} \in R, w_{\mu\nu} \in R^+$$

The imaginary part of the self-energy can be rewritten as

$$\Gamma_{L/R} = -2\Im[W_{L/R}]$$

We compute the Green function of the system

$$G(E,k) = [(E+i\eta)I - H'(k)]^{-1}, \ \eta \ll 1$$

The transmission and reflection coefficients are defined as

$$\begin{split} T(E,k) = & \text{Tr}[\mathbf{\Gamma}_{L}(E,k)\mathbf{G}(E,k)\mathbf{\Gamma}_{R}(E,k)\mathbf{G}^{\dagger}(E,k)] \\ R(E,k) = & T_{\text{bulk}}(E,k) - \left\{ i \text{Tr}[(\mathbf{G}(E,k) - \mathbf{G}(E,k)^{\dagger})\mathbf{\Gamma}_{\text{elec}}(E,k)] \right. \\ & - & \text{Tr}[\mathbf{G}(E,k)\mathbf{\Gamma}_{L}(E,k)\mathbf{G}(E,k)^{\dagger}\mathbf{\Gamma}_{L}(E,k)] \right\} \end{split}$$

 Γ must be positive-definite (i.e. with strictly positive eigenvalues) to ensure that the transmission (and reflection) is positive.

3 Content

- job_opt : job file to run the algorithm on the cluster
- cap_opt.py : python algorithm
- input.csv : user inputs for the algorithm
- Tobj: folder containing the exact transmission as a .npy file
- Robj: folder containing the exact reflection as a .npy file
- geom: folder containing the geometry or hamiltonian of the device, left and right electrode, as .xyz or .TSHS files
- res_opt : folder containing the result of the optimization, as a .npy file

4 Input

- \bullet Emin/Emax : minimum/maximum energy defining the optimization range.
- \bullet ne: number of energy points.
- \bullet kmin/kmax: minimum/maximum transverse periodic boundary condition number defining the optimization range.
- nk: number of k-points.
- eta: infinitesimal imaginary part of energy, for convergence of Green function calculation.
- axis: transport direction (0, 1 or 2).
- dev_file/elec_left_file/elec_right_file: name of device, left electrode and right electrode file inside the geom folder. The device must contain the left and right electrode, with same orbital order than in the electrode files, and in this order: left device right.

- if .xyz format, it is assumed to be a graphene-like system, therefore
 the hamiltonian is defined with the hopping elements that are written
 at the beginning of cap_opt.py file.
- if .TSHS format, the hamiltonian is directly used as is all 3 files must be in the same format.
- nsc: number of super cells, list of 3 ints like "[1,3,1]". The supercell number in the transport direction must be 1. In the periodic boundary condition direction it is typically 3.
- Tobj_file/Robj_file: name of exact transmission/reflection inside the Tobj/Robj folder. These are .npy files with shape (ne,nk). If Tobj_file/Robj_file is set to None, the transmission/reflection will be recomputed. Robj_file is only used if the keyword "refl" is written in the 'mode' input (see below).
- kproc : either "none", "avg" or "inde".
 - if "none" each k-point is considered in the least square objective function : $\min_{CAP} \sum_{E} \sum_{k} (T_{CAP}(E,k) T_{obj}(E,k))^2$
 - if "avg" the average of the transmission over k is considered in the least square objective function : $\min_{CAP} 1/n_k^2 \sum_E [\sum_k (T_{CAP}(E,k) T_{obj}(E,k))]^2$
 - if "inde" a different optimized CAP is computed for each k-point : $\forall k, \min_{CAP(k)} \sum_{E} (T_{CAP(k)}(E,k) T_{obj}(E,k))^2$
- *tol* : termination condition of the optimization algorithm, the gradient of the objective function must be inferior to this value.
- save_file: name of the .npy file where the optimized CAP is saved, inside the res_opt folder.
- *mode*: list or string of keywords, the possible keywords are "diag, real, hop, X0_zero, refl, sym, transym".
 - if "diag" the on-site potential of each orbital are optimised, the imaginary part is negative to satisfy the condition of definite-positiveness of Γ. $w_{\mu\mu}$ is a variable, with bound $w_{\mu\mu} > 0$. By default, $v_{\mu\mu} = 0$, the CAP is pure imaginary.
 - if "hop" the hopping elements between orbitals are added to the optimization, W is now a symmetric matrix. This requires a new condition for Γ to be definite-positive. A simple and linear condition is enforcing the matrix to be diagonally dominant: $\forall \mu, w_{\mu\mu} > \sum_{\nu \neq \mu} w_{\mu\nu}$, $\forall \mu\nu, w_{\mu\nu} > 0$. Thus, linear constraints are added to the optimization problem, resulting in much higher computation time. NOT COMPATIBLE WITH "real" AND "transym". WHEN "hop" IS ACTIVATED, "sym" MUST BE ACTIVATED ASWELL.

- if "real" the elements of the CAP (diag or/and hop) have also a real part, independent from the imaginary part. $v_{\mu\nu}$ is non-zero, subject to no bounds or constraints. This results in twice the number of variables to optimize so a multiplication of the computation time.
- if "X0-zero" the first guess of the optimization process is a zero matrix. If "X0-zero" is not in mode, then a standard diagonal CAP is used as a first approximation: $W_{\mu\mu} = -i\frac{\hbar^2}{2m}\left(\frac{2\pi}{r_1}\right)^2\frac{4}{c^2}\left[\frac{r_1^2}{(r_1-r_{\mu\mu})^2} + \frac{r_1^2}{(r_1+r_{\mu\mu})^2} 2\right]$, where r_1 is the length of the electrode along the transport direction, $r_{\mu\mu}$ is the distance of orbital μ to the device, projected onto the transport direction, and c=2.62.
- if "refl" the reflection is optimized as well as the transmission.
- if "sym" the right and left electrodes are symmetric. The the same CAP are added to both electrodes: $W_L = W_R$. Use the '.mirror()' method in sisl to build the electrodes. This divides by two the number of variables and improves greatly the efficiency.
- if "transym" both electrodes are symmetric along the periodic boundary condition direction: $W_{\mu\nu} = W_{\mu+no/2,\nu+no/2}$. This divides by two the number of variables and improves greatly the efficiency.

5 Run

The algorithm can be run either by running directly "python3 cap_opt.py" in a command shell, or by running "bsub $\angle job_opt$ " on the cluster. After it is finished, all the information about the run can be found in registre.csv and the result in the res_opt folder.