Exploring the limits of the self consistent Born approximation for inelastic electronic transport

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The non equilibrium Green function formalism is today the standard computational method for describing elastic transport in molecular devices. This can be extended to include inelastic scattering by the so called self-consistent Born approximation (SCBA), where the interaction of the electrons with the vibrations of the molecule is assumed to be weak and it is treated perturbatively. The validity of such an assumption and therefore of the SCBA is difficult to establish with certainty. In this work we explore the limitations of the SCBA by using a simple tight-binding model with the electron-phonon coupling strength $\alpha$ chosen as a free parameter. As model devices we consider Au mono-atomic chains and a $\text{H}_2$ molecule sandwiched between Pt electrodes. In both cases our self-consistent calculations demonstrate a breakdown of the SCBA for large $\alpha$ and we identify a weak and strong coupling regime. For weak coupling our SCBA results compare closely with those obtained with exact scattering theory. However in the strong coupling regime large deviations are found. In particular we demonstrate that there is a critical coupling strength, characteristic of the materials system, beyond which multiple self-consistent solutions can be found depending on the initial conditions in the simulation. We attribute these features to the breakdown of the perturbative expansion leading to the SCBA.

I. INTRODUCTION

Central to the field of molecular electronics are phenomena involving the interaction between the electron current and the internal degrees of freedom of the molecule investigated. In scanning tunnel microscopy (STM) the molecular vibrational modes (phonons) have been exploited to desorb or to move a molecule on a surface, paving the way for phonon assisted surface chemistry. At the same time, STM inelastic tunnelling spectroscopy uses the fingerprints of vibrations in the $I$-$V$ curve to probe the orientation and/or to identity molecules on surfaces. Switching devices exploiting phonons have also been reported.

Broadly speaking, in molecular devices phonons are important for two reasons. Firstly, they play a role in transport by opening new conductance channels through which the itinerant electrons can propagate, and by suppressing the transmission of purely elastic channels. More dramatically, for large electron-phonon coupling the charge carriers become quasi-particles consisting of coupled electrons and phonons. Secondly, from a technological point of view, phonons limit the efficiency of molecular devices because of energy dissipation. This causes heating, power loss and instability.

Transport experiments at the nano-scale are difficult to interpret since the atomically precise device geometry is rarely known. Therefore one usually relies on atomistic simulation techniques in order to understand the results. For elastic transport, when electron-phonon interaction is not considered and the electron-electron interaction is treated at the mean field level, methods of note for predicting the current flowing through devices include the non-equilibrium Green function formalism (NEGF) and scattering theory (ST). Some of these methods have been adapted to include electron-phonon interaction, notably an extension of scattering theory (EST) and the self consistent Born approximation (SCBA) within the NEGF formalism. In addition time-dependent methods for describing correlated electron-ion dynamics have been recently proposed.

The focus of this paper is the SCBA. This is attractive from a practical point of view since it has moderate computational requirements and it has been used extensively for calculating transport properties of a number of different material systems. However, it is a perturbative approach appropriate only for weak electron-phonon (e-p) coupling. As the e-p coupling strength increases the SCBA will eventually breakdown, however it is unclear whether such breakdown is either sharp or smooth with the e-p coupling strength. Our work explores this question in detail.

The paper is organized as follows. We begin by presenting the NEGF formalism for a two probe device, and by recalling the foundations of the SCBA. We then consider a 1D tight-binding model where the e-p interaction in the scattering region is described by the Su-Schrieffer-Heeger (SSH) Hamiltonian. The parameters for the model Hamiltonian are chosen for mimicking two systems which have been studied experimentally: $\text{H}_2$ molecules sandwiched between Pt electrodes ($\text{H}_2$-$\text{Pt}$) and Au monatomic chains comprising R atoms (RC's). The parameters for $\text{H}_2$-$\text{Pt}$ are the same as those used by Jean and Sanvito, who previously employed exact scattering theory (EST) to describe phononic effects. In contrast to the SCBA, EST is valid for both strong and weak coupling and therefore it is a good benchmark for the SCBA. Accordingly we compare the SCBA results directly with EST over a range of different e-p couplings to establish the limit of validity for the SCBA and to investigate its breakdown.
II. METHODOLOGY

A. Non Equilibrium Green Function Formalism

A two probe device consists of two crystalline electrodes attached on either side of a scattering region, which is in general a collection of atoms breaking the electrodes translational symmetry. The leads are also charge reservoirs, so that the device may be viewed as two charge reservoirs bridged by the central region. Thermodynamically we characterise the left-hand side (L) and right-hand side (R) lead by defining their chemical potentials $\mu_L$ and $\mu_R$. If $\mu_L = \mu_R$, equilibrium is established and no current flows. When $\mu_L \neq \mu_R$ the system is dragged out of equilibrium, and net charge will move from the reservoir with the higher chemical potential across the central region to the reservoir of lower chemical potential in an attempt to re-establish equilibrium. If a battery is attached to the two reservoirs keeping $\mu_L - \mu_R = eV$ ($V$ is the bias and $e$ the electron charge) the system cannot return to equilibrium and will eventually reach a steady state with a constant current flow.

At the Hamiltonian level the problem can be formulated by using a basis set comprising a linear combination of atomic orbitals (LCAO). It is convenient to write the Hamiltonian of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO). It is convenient to write the Hamiltonian level of the semi-infinite periodic leads in terms of atomic orbitals (LCAO).

\[ \begin{pmatrix}
  H_{11} & H_{12} & \cdots & H_{1N} \\
  H_{21} & H_{22} & \cdots & H_{2N} \\
  \vdots & \vdots & \ddots & \vdots \\
  H_{N1} & H_{N2} & \cdots & H_{NN}
\end{pmatrix} \]

describe respectively the interactions between PLs and within a PL. The scattering region in general is described by $M$ basis functions. The $M \times M$ matrix, $H_{LM}$, describes its internal interaction, while the matrices $H_{LM}$ ($N \times M$) and $H_{RM}$ ($M \times N$) contains the interaction between the PLs of the leads adjacent to the scattering region and the scattering region itself. The entire system is thus described by the infinite tri-diagonal Hamiltonian $H$

\[
H = \begin{pmatrix}
  H_{0} & H_{1} & \cdots & H_{1} \\
  H_{0} & H_{0} & \cdots & H_{1} \\
  \vdots & \vdots & \ddots & \vdots \\
  H_{0} & H_{0} & \cdots & H_{0}
\end{pmatrix}
\]

Time reversal symmetry sets $H_{-1} = H_{1}^\dagger$, $H_{ML} = H_{LM}^\dagger$, and $H_{MR} = H_{RM}^\dagger$. The retarded Green function, $G^R$ associated to the entire system (leads plus scattering region) is defined as

\[
[g^R(\omega) = I, \quad (1)]
\]

where $\omega' = \lim_{\delta \rightarrow 0^+} (\omega + i\delta)$, $\omega$ is the energy and $I$ is the infinite dimensional identity matrix. For transport calculations however one does not need the Green’s function of the entire system but only that relative to the scattering region, $G_M$, in presence of the leads. This can be written as

\[
G_M(\omega) = [\omega' I_M - H_M - \Sigma_L(\omega) - \Sigma_R(\omega)]^{-1}, \quad (2)
\]

where the presence of the leads have been accounted via the introduction of the self-energies for the left- and right-hand side lead $\Sigma_L(\omega)$ and $\Sigma_R(\omega)$. $I_M$ is the $M \times M$ identity matrix. The self-energies are $M \times M$ matrices defined as

\[
\Sigma_L = H_{ML} g_L H_{LM}, \quad \Sigma_R = H_{MR} g_R H_{RM}, \quad (3)
\]

where $g_L(\omega)$ and $g_R(\omega)$ are the retarded surface Green functions of the leads, namely the retarded Green functions of the isolated semi-infinite leads evaluated at the PLs adjacent to the scattering region. These are calculated by considering the retarded Green function of the corresponding infinite system (periodic) and by applying appropriate boundary conditions. External bias voltage is introduced under the assumption that the leads are good metals maintaining local charge neutrality. The effect of a bias is therefore only that of shifting rigidly in energy the leads electronic structure, so that

\[
\Sigma_{L/R}(\omega, V) = \Sigma_{L/R}(\omega \pm eV/2, 0). \quad (4)
\]

We now proceed to evaluate the non-equilibrium charge density in the scattering region and the two-probe current by using the NEGF scheme. The lesser ($<$) and greater ($>$) Green functions $G^<_M(\omega)$ are defined as

\[
G^<_M(\omega) = G_M(\omega)\Sigma^<_M(\omega)G^>_M(\omega), \quad (5)
\]

with self-energies

\[
\begin{align*}
\Sigma^<_M(\omega) &= \sum_{\alpha=L,R} \Sigma^\alpha_M(\omega), \\
\Sigma^\alpha_M(\omega) &= i n^\alpha_R(\omega) \Gamma^\alpha_M(\omega), \\
\Gamma^\alpha_M(\omega) &= i [\bar{n}^\alpha_L(\omega) - 1] \Gamma^\alpha_M(\omega).
\end{align*}
\]

FIG. 1: Schematic representation of a system composed of two semi infinite leads and a scattering region (rectangular dashed box). The matrices $H_0$ and $H_1$ describe the lead principal layers, $H_M$ describes the scattering region and $H_{LM}$, $H_{RM}$ the interaction between the scattering region and the last principal layers of the leads.
Here \( n_F^0(\omega) = n_F(\omega - \mu_o) \) is the Fermi function evaluated at \( \omega - \mu_o \) and temperature \( T \), \( \mu_o = E_F \pm eV/2 \) with \( E_F \) the leads Fermi energy, and we have introduced the coupling matrix for the \( \alpha \) lead (\( \alpha = L/R \))

\[
\Gamma_\alpha(\omega, V) = i[\Sigma_\alpha(\omega, V) - \Sigma_\alpha^\dagger(\omega, V)].
\] (9)

The non-equilibrium charge density matrix for the scattering region is

\[
\rho = \frac{1}{2\pi i} \int_{-\infty}^{\infty} d\omega' G_M^\dagger(\omega') .
\] (10)

If \( H_M \) has a functional dependence on \( \rho \) the equations [2] and [10] can be solved self-consistently. The net current flowing through the device is then

\[
J_{\text{unp}}(V) = \frac{2e}{h} \int_{-\infty}^{\infty} d\omega \text{Tr}[\Gamma_L G_M^\dagger \Gamma_R G_M](n_F^0 - n_F^R),
\] (11)

where the subscript “unp” stands for “unperturbed”, meaning that no e-p interaction is included. The term \( T(\omega, V) = \text{Tr}[\Gamma_L G_M^\dagger \Gamma_R G_M] \) is the standard Landauer Büttiker transmission coefficient, although in this case it is explicitly bias dependent. The conductance \( G \) in the linear response limit is

\[
G = \frac{2e^2}{h} T(E_F, 0),
\] (12)

while more generally at a given bias \( V \) one has

\[
G(V) = \frac{dJ_{\text{unp}}}{dV} \bigg|_V .
\] (13)

B. Self Consistent Born Approximation

We now discuss the main concepts associated with introducing e-p scattering into the NEGF transport scheme. In general, inelastic scattering produces loss of phase coherence, similarly to what happens when an electron is absorbed by a reservoir. In fact one may think of inelastic processes as resulting from the coupling of the scattering region to a “fictitious” charge reservoir that does not exchange a net current. Thus e-p interaction can be introduced via a self-energy \( \Sigma_{ph}(\omega) \) and the retarded Green function becomes

\[
G_M(\omega) = [\omega I_M - H_M - \Sigma_L(\omega) - \Sigma_R(\omega) - \Sigma_{ph}(\omega)]^{-1}.
\] (14)

The exact form for \( \Sigma_{ph}(\omega) \) is unknown, however convenient approximations can be derived from the perturbative expansion over the e-p coupling strength [16,24,25,28,34]. In this work we consider the SCBA where only the Hartree and Fock diagrams of the perturbative expansion are retained (see figure [2]). This is equivalent to evaluating the first order diagrams at the interacting electronic Green function. Thus the phonon self-energy reads

\[
\Sigma_{ph}(\omega) = \Sigma^F(\omega) + \Sigma^H,
\] (15)

where the retarded Hartree (H) and Fock (F) contributions to the self-energies are respectively [28,35]

\[
\Sigma^H = i \sum_{\lambda} \frac{4}{\Omega} \int_{-\infty}^{\infty} \frac{d\omega'}{2\pi} M^H \text{Tr}[G_M^<(\omega') M^L],
\] (16)

\[
\Sigma^F(\omega) = \sum_{\lambda} \left[ \frac{1}{2} \Sigma_\lambda^<(\omega) - \Sigma_\lambda^>(\omega) \right]
- \frac{i}{2} \Omega \{ \Sigma_\lambda^<(\omega') - \Sigma_\lambda^>(\omega') \}(\omega).
\] (17)

In equation [17] \( H_\omega \) is the Hilbert transform

\[
H_\omega \{ f(x) \}(y) = \frac{1}{\pi} \mathcal{P} \int_{-\infty}^{\infty} \frac{f(x)}{x-y},
\] (18)

and \( \mathcal{P} \) stands for the principal part of the integral. The phonon energy and e-p coupling matrix for a particular mode \( \lambda \) are respectively \( \Omega_\lambda \) and \( M^\lambda \). Finally the e-p lesser and greater self-energy are given by

\[
\Sigma^F_\lambda(\omega) = \sum_\lambda \Sigma_\lambda^F(\omega),
\] (19)

\[
\Sigma^S_\lambda(\omega) = M^\lambda \left[ (N_\lambda + 1) G_M^S(\omega \mp \Omega_\lambda) + N_\lambda G_M^S(\omega \mp \Omega_\lambda) \right] M^\lambda,
\] (20)
which is simply a sum of the lesser self-energies over all the possible modes $\lambda$. The occupancy of each phonon mode is $N^\lambda$.

We assume that the phonons are in thermal equilibrium with a bath so that for a given temperature $N^\lambda$ is simply the Bose-Einstein distribution $N^\lambda = (e^{\Omega^\lambda/\kB T} - 1)^{-1}$, with $\kB$ the Boltzmann constant. From equation (20) we note that the lesser (greater) self-energy contains contributions from two scattering processes: electrons with energy $\omega - \Omega^\lambda$ may absorb (emit) a phonon and/or electrons with energy $\omega + \Omega^\lambda$ may emit (absorb) a phonon of energy $\Omega^\lambda$. When $T \sim 0$, electrons may only emit phonons since $N^\lambda \sim 0$, i.e. no phonons are present in the scattering region provided that the phonon lifetime is much smaller than that of the electrons. The total lesser self-energy of equation (20) must be adjusted to include the phonon lesser self-energy,

$$
\Sigma^\beta(\omega) = \sum_{\alpha=\text{L,R}} \Sigma^{\alpha\beta}_{\alpha\beta}(\omega) + \Sigma^\beta_{\text{ph}}(\omega)
$$

(21)

and $G^\beta(\omega)$ from equation (14) are now evaluated using the perturbed Green function and lesser self-energy from equations (14) and (21). The general expression for the interacting (including e-p coupling) current through lead $\beta$ may be written as the sum of an elastic and an inelastic contribution

$$
J^\beta(V) = J^\beta_{\text{el}}(V) + J^\beta_{\text{inel}}(V),
$$

(22)

where

$$
J^\beta_{\text{el}} = \frac{2e}{\hbar} \int_{-\infty}^{\infty} d\omega \text{Tr}[\Gamma G^\beta \Gamma G^\beta_{\text{M}}(n_F - n_{F}^\beta)]
$$

(23)

and

$$
J^\beta_{\text{inel}} = \frac{2e}{\hbar} \int_{-\infty}^{\infty} d\omega \text{Tr} \left[ \Sigma^\beta G^\beta \Sigma^\beta_{\text{ph}} G^\beta_{\text{M}} - \Sigma^\beta G^\beta \Sigma^\beta_{\text{ph}} G^\beta_{\text{M}} \right].
$$



### III. NUMERICAL METHOD

#### A. Model Hamiltonian and Coupling Matrices $M^\lambda$

The systems under investigation are 1D linear atomic chains described by a $s$-orbital nearest-neighbour tight-binding model. The scattering region comprises $R$ atoms plus one PL (one atom) from each lead, so that it contains $M = R + 2$ orbitals. Henceforth we refer to this system as RC. Furthermore, we assume that the two leads are identical. The matrices $H_0$, $H_1$ for a 1D tight-binding model reduce to c-numbers: $\epsilon_L = H_0$ and $\gamma_L = H_1$, where $\epsilon_L, \gamma_L$ are the lead onsite energy and hopping parameter respectively. The leads’ Hamiltonians thus read

$$
H_{\text{L}} = \epsilon_{\text{L}} \sum_{i=-\infty}^{-1} c_i^\dagger c_i + \gamma_{\text{L}} \sum_{i=-\infty}^{-2} [c_i^\dagger c_{i+1} + c_{i+1}^\dagger c_i],
$$

(24)

$$
H_{\text{R}} = \epsilon_{\text{L}} \sum_{i=R+2}^{\infty} c_i^\dagger c_i + \gamma_{\text{L}} \sum_{i=R+2}^{\infty} [c_i^\dagger c_{i+1} + c_{i+1}^\dagger c_i].
$$

(25)

Finally e-p interaction is included into the scattering region in the form of an SSH Hamiltonian\cite{SSH}, comprising three terms

$$
H_M = H_e + H_{\text{ep}} + H_{\text{ph}},
$$

(31)

with

$$
H_e = \sum_{i=0}^{R+1} \epsilon_i c_i^\dagger c_i + \sum_{i \neq j} \gamma_{ij}^0 (c_i^\dagger c_j + c_j^\dagger c_i),
$$

(32)

$$
H_{\text{ph}} = \lambda_{\text{max}} \sum_{\lambda=1}^{R+1} (b_{\lambda}^\dagger b_{\lambda} + 1/2) \hbar \omega_{\lambda},
$$

(33)

$$
H_{\text{ep}} = \lambda_{\text{max}} \sum_{\lambda=1}^{R+1} \sum_{i \neq j} M_{ij} (b_{\lambda}^\dagger b_{\lambda} + b_{\lambda} b_{\lambda}^\dagger) (c_i^\dagger c_j + c_j^\dagger c_i).
$$

(34)

$H_e$ is the electronic Hamiltonian of the scattering region with onsite energies $\epsilon_i$ and the unperturbed hopping parameters $\gamma_{ij}^0$ (we assume $\gamma_{ij}^0 = 0$ for $j \neq i \pm 1$). $H_{\text{ph}}$ is the non-interacting phonon Hamiltonian, written in terms of the phononic creation and annihilation operators $\{b_{\lambda}^\dagger, b_{\lambda}\}$ and the phonon energies $\Omega_{\lambda} = \hbar \omega_{\lambda}$,
with the index \( \lambda \) running over all the modes. The final term, \( H_{ep} \), is the Hamiltonian describing the e-p interaction within the scattering region. The details of such interaction are included in the e-p coupling matrices \( M^\lambda \).

In order to calculate the matrices \( M^\lambda \) and the longitudinal phonon frequencies we consider a simple nearest neighbours elastic model in which

\[
M^\lambda_{ij} = \alpha_{ij} \left\{ \frac{e^i - e^j}{\sqrt{m_i}} - \frac{e^j - e^i}{\sqrt{m_j}} \right\} \frac{\hbar}{2\omega}.
\]

In equation (35) the orthonormal vectors \( e^\lambda \) represent the ionic displacement associated to each mode \( \lambda \), \( m_i \) is the mass of the atom at site \( i \), and the constants \( \alpha_{ij} \) are the e-p coupling parameters. These latter are defined as the first order coefficient of the expansion of the tight-binding hopping parameter \( \gamma_{ij} \) about the atomic equilibrium positions

\[
\gamma_{ij} = \gamma^0_{ij} + \alpha_{ij}(u_i - u_j),
\]

where \( u_i \) is the displacement vector of the atom at site \( i \). From equation (36) it follows that \( \alpha_{ij} = -\alpha_{ji} \). In the nearest neighbour approximation the interaction is restricted to electrons moving between sites \( i \) and \( i \pm 1 \). Finally, we note that the eigenvectors \( e^\lambda \) are real. This implies that the matrices \( M^\lambda \) are real and symmetric with non-zero matrix elements for \( i = i \pm 1 \), so that \( M^\lambda \) has a tri-diagonal form for longitudinal phonons. We note that, although it is possible to calculate the coupling parameters \( \alpha_{ij} \) using first-principles electronic structure methods, here we set \( \alpha_{ij} = \alpha \) and \( \alpha \) is taken as a free parameter.

B. Numerical Integration and Self Consistency

The flowchart in figure 4 outlines the numerical procedure used to calculate the interacting current \( J^\beta(V) \) and the differential conductance \( G(V) \). Each simulation can be partitioned into three steps. The first two consist of two self-consistent loops which calculate the phonon self-energies \( \Sigma^H \) and \( \Sigma^F \) respectively. These are used in the third step to evaluate \( J^\beta(V) \) by using the equations (22) and (23).

Let us now discuss the three steps in some detail. We start by writing \( \Sigma^H \) as an explicit function of the density matrix \( \rho \)

\[
\Sigma^H(\rho) = -4 \sum \lambda \text{Tr} \left[ \frac{M^\lambda}{\Omega^\lambda} \right] M^\lambda, \tag{37}
\]

where we assume that all the elements of \( G^\lambda_M \) are integrable. \( \Sigma^H \) is thus nothing but a weighted sum of the matrices \( M^\lambda \). This can be written in the form

\[
\Sigma^H = \sum_{\lambda=1}^M r^\lambda R^\lambda, \tag{38}
\]

where the ratio matrices \( R^\lambda \) and their weighting coefficients \( r^\lambda \) are given by

\[
R^\lambda = \left| \gamma^0 \right|_{\text{min}} \frac{M^\lambda}{M^\lambda_{\text{max}}}, \tag{39}
\]

\[
r^\lambda = \frac{4 |M^\lambda|_{\text{max}}}{\Omega^\lambda |\gamma^0|_{\text{min}}} \sum_{i=1}^{M-1} R^\lambda_{i,i+1} (\rho_{i,i+1} + \rho_{i+1,i}), \tag{40}
\]

For a given mode \( \lambda \), the largest matrix element of the matrix \( M^\lambda \) is denoted as \( |M^\lambda|_{\text{max}} \). \( |\gamma^0|_{\text{min}} \) is the smallest among the hopping parameters of the unperturbed system (no e-p coupling), which, by construction, is equal to \( |R^\lambda|_{\text{max}} \). The matrices \( R^\lambda \) are independent of the e-p coupling \( \alpha \) and simply reflect the symmetry of the specific phonon mode considered. Thus \( r^\lambda \) measures the maximum fractional modification of the elements \( \gamma^0_{ij} \) in the electronic Hamiltonian as the result of e-p coupling.

The first self-consistent loop of figure 4 begins by choosing initial values for the weighting coefficients, \( \{r^\lambda\}_{\text{init}} = \{r^0_1, r^0_2, ..., r^0_R\} \), which are used to construct the density matrix at the first iteration, \( \rho^1 \). Then both
\( \{r_\lambda \} \) and \( \rho \) are varied until the convergence condition
\[
e^0 = \sum_\lambda \left( \frac{|r_\lambda^n - r_\lambda^{n-1}|}{|r_\lambda^n - r_\lambda^{n-1}|} \right) < t_H \tag{41} \]
is met for the chosen tolerance \( t_H \). The density matrix \( \rho \) is obtained by integrating \( G^< \) as in equation (10). In order to perform this integral we first take \( \Sigma^F = 0 \) under the assumption that this has little effect of the convergence of \( \rho \) (first self-consistent loop). Following previous work we write \( \rho = \rho_{eq} + \rho_V \), where
\[
\rho_{eq} = -\frac{1}{\pi} \int_{-\infty}^{\infty} d\omega \text{Im}[G_M] \frac{1}{n_F^{\dagger}} \tag{42} 
\]
and
\[
\rho_V = \frac{1}{2\pi} \int_{-\infty}^{\infty} d\omega G_M \Gamma_R G_M^{\dagger} (n_F^{\dagger} - n_F). \tag{43} 
\]
At equilibrium \( (\mu_L = \mu_R = \mu) \) one has \( \rho = \rho_{eq} \).

FIG. 5: Schematic representation of the integrals in equations (42) and (43) respectively. In a) the integration of \( \rho_V \) is bound between \( E_F + eV/2 \) and \( E_F - eV/2 \) by the Fermi functions. \( \Gamma_{P1} \) is the number of points of the real axis energy grid. For \( \rho_{eq} \) in b) the number of grid points along the circular path \( (\text{cp}_1) \) and the path in the complex plane parallel to the real axis \( (\text{cp}_2) \) must be chosen, as well as the poles in the Fermi functions.

As shown in figure 3, the integral of \( \rho_V \) (Eq. (12)) is along the real axis and it is bound between the chemical potentials \( \mu_L \) and \( \mu_R \). This is carried out over a numerical grid of sufficient fineness \( (dE)_H \). The calculation of \( \rho_{eq} \) involves an unbound integral. This is performed over a coarse grid in the complex plane using a contour integral method, since \( G_M \) is analytical and smooth in the imaginary energy plane. As shown in figure 5 a number of numerical parameters must be chosen. First the lower limit of integration \( E_R \) must lie below the lowest lying molecular states and below the lowest electrode bands. Secondly, the poles of the Fermi functions (Matsubara frequencies) which lie within the contour must be taken into account. The integration is then performed by using Gaussian quadrature.

The second self-consistent loop begins by calculating \( \{G^0_M, G^<0\} \) using equation (14), where the converged \( \Sigma^R \) from the first loop is used and \( \Sigma^F = 0 \). We then proceed to iterate \( \{\Sigma^F, \Sigma^R_{ph}\} \) and \( \{G_M, G^<\} \) until a second convergence condition is met
\[
e^2 = \frac{1}{N_p} \text{Max} \left\{ \left| G_M^{\omega_i} - G_M^{\omega_{i-1}}(\omega_i) \right| \right\} < t_F . \tag{44} 
\]

Note that the condition is over the largest of the matrix elements and runs over the \( N_p^F \) energy points \( \omega_i \) of the entire grid. Note also that the tolerance used, \( t_F \), is in general different to that used for the Hartree term. The Hilbert transform required for calculating the imaginary part of \( \Sigma^F \) (eq. (17)) is done by using a convolution method combined with a fast Fourier transform algorithm. In order to avoid end-point corrections we choose a grid of sufficient range while the grid fineness \( (dE)_F \) must be sufficiently fine to resolve phononic features which lie in the meV range.

Table 4 shows the numerical and system parameters used in our simulations. The parameters for the H$_2$-Pt junctions are identical to those used by Jean within the EST treatment of phonons. This set produces the same unperturbed \( G(0) \sim 97G_0 \). The spring constants and masses are chosen to give longitudinal phonon modes of energies 63 meV (CM mode) and 432 meV respectively while the ratio matrices for these two modes are
\[
R^1 = \begin{pmatrix}
0 & 3.2 & 0 & 0 \\
3.2 & 0 & 0 & 0 \\
0 & 0 & 0 & 3.2 \\
0 & 0 & -3.2 & 0
\end{pmatrix} \tag{45} 
\]
and
\[
R^2 = \begin{pmatrix}
0 & 1.6 & 0 & 0 \\
1.6 & 0 & -3.2 & 0 \\
0 & -3.2 & 0 & 1.6 \\
0 & 0 & 1.6 & 0
\end{pmatrix}. \tag{46} 
\]
The e-p coupling \( \alpha \) remains a free parameter.

The parameters for the Au RC’s match closely those used by Frederiksen of. The atoms in the leads are chosen as identical to the atoms in the scattering region, thus that a single onsite energy and hopping parameter characterise the electronic Hamiltonian. These parameters and the equilibrium potential \( \mu_{eq} \) are chosen so that the differential conductance for the unperturbed system is \( G_0 = 2e^2/h \) (perfect transmission).

IV. RESULTS: H$_2$-Pt JUNCTIONS

A. self-consistent simulations

The self-consistent \( G(V) \) in the range 0-200 meV for \( \alpha \) ranging between 0 and 3.1 eV/Å are presented in figure 6. For this system the characteristic signature of e-p interaction is a drop in the conductance at a threshold voltage \( V_{thr} \). This signals the onset of inelastic electron
processes involving the emission of phonons with energies $\Omega \approx \epsilon_{V_{\text{thr}}}$. We quantify this effect by defining the conductance drop (in units of $G_0$)

$$\Delta_{\text{thr}} = G(0) - G(V_{\text{thr}}).$$  \hspace{1cm} (47)

Numerical simulations are carried out in two different ways. First, we follow the exact numerical procedure of figure 4 ("full SCBA"), but we run simulations starting with different initial conditions, namely \( \{\lambda\}_{\text{init}}^{+} = +0.003125 \) and \( \{\lambda\}_{\text{init}}^{-} = -0.003125 \). In figure 4, \( \Delta_{\text{thr}} \) is plotted versus \( \alpha \) demonstrating good agreement between the full SCBA and EST for low \( \alpha \). However, the two methods disagree for \( \alpha \) beyond \( \alpha_{\text{crit}} \sim 1.8 \text{ eV/Å} \). \( \Delta_{\text{thr}} \) peaks sharply above \( \alpha_{\text{crit}} \), beyond which it becomes dependent on the initial condition \( \{\lambda\}_{\text{init}}^{+} \). This last situation is shown in figure 6c for \( \alpha = 2.0 \text{ eV/Å} \), where two different \( G(V) \) curves are predicted for different \( \{\lambda\}_{\text{init}}^{+} \) and a low-bias conductance of 0.0125 \( G_0 \) is observed in stark disagreement with the unperturbed value of \( \sim 0.97 G_0 \).

Since the Hartree self-consistent loop is performed before the Fock one, the dependence on the initial conditions suggests that \( \Sigma^H \) may be responsible for the behaviour observed for \( \alpha > \alpha_{\text{crit}} \). Stronger evidence to support this hypothesis is provided in figures 6b and 6d, where the results for our second set of simulations in which \( \Sigma^H \) is set to zero, are presented ("partial SCBA"). Figure 6d shows good agreement between the partial SCBA and the EST to the much higher coupling of \( \alpha \approx 4.0 \text{ eV/Å} \). The full SCBA disagrees with the EST at a considerably lower \( \alpha \approx 1.8 \text{ eV/Å} \). Above such a coupling strength \( G(V) \) depends on the initial conditions.

### B. Contribution from the individual modes

We now analyze the origin of the breakdown of the full SCBA. For \( V << V_{\text{thr}} \), the inelastic current \( J_{\text{inel}} \) is strongly suppressed by Pauli exclusion principle. At low temperature \( T = 4.0 \text{ K} \) we can approximate the Fermi distributions in \( J_\alpha \) by step functions to obtain

$$J_\alpha^\beta(V) \approx \frac{2}{\hbar} \int_{\mu_L}^{\mu_H} T(\omega) d\omega.$$

We can now easily probe the contribution of the Hartree term to the conductance by considering the test Green function

$$G_\alpha^H(\omega) = [\omega' I_M - H_M - \Sigma_L - \Sigma_R - r_\alpha R^\lambda]^{-1}.$$  \hspace{1cm} (48)
This is used to evaluate the transmission coefficient at $V = 0$ and it is useful to understand the influence of the individual modes $\lambda$ over the transmission. In figure 7 we show $T(\omega)$ as a function of $r_\lambda$ for the two longitudinal modes available in the H$_2$-Pt system (see figure 5). In the case of the rigid translational mode ($\lambda = 1$ and figure 4b) $T(\omega)$ is reduced in the region around $E_F$ as $|r_1|$ increases. However the general shape of $T(\omega)$ is little affected. This is somehow expected from the shape of the matrix $R^1$ (eq. 14)), which indicates that mode 1 causes simply a change in the hopping parameters $\gamma_{0}$ and $\gamma_{34}$ connecting the molecule to the leads. Importantly when $\gamma_{0}$ is increased, $\gamma_{34}$ is reduced by the same amount and vice-versa. Moreover there is a symmetry $r_1 \rightarrow -r_1$.

The results for the symmetric mode $\lambda = 2$ are presented in figure 7b. This time the peak in transmission is shifted in energy either to the left or to the right depending on the sign of $r_2$. When shifted to the left, the peak is broadened while a shift to the right narrows it. In either case the transmission around $E_F$ is reduced.

From the figures 4c, 8a, and 8b one can conclude that the deviation of the zero bias differential conductance from its unperturbed (no e-p interaction) value is a measure of the magnitude of the e-p perturbation. We define this deviation as

$$\Delta G = G_0 - G(0) \equiv G_0 - T(E_F),$$

where the last equality is valid for $T \rightarrow 0$. Figures 9a and b show the estimated deviation (in units of $G_0$) versus the weighting coefficients $r_\lambda$. The maximum deviation, $\Delta G = 1$, occurs when the chain actually breaks as for mode 1 and $r_1 = 1$. As expected, the curve in figure 9a for mode 1 is symmetric about 0 while figure 9b for mode 2 is not.

C. Discussion of the self-consistent results

We finally re-analyze our self-consistent results in the light of the discussion in the previous section. Figure 10a and 10b show the self-consistently calculated $r_\lambda$ as a function of $\alpha$ for $V = 0$, while figure 10c shows $\Delta G$ as defined in equation (18). The critical point $\alpha_{\text{crit}} \sim 1.8$ eV/Å marks a sharp transition in the behaviour of the weighting coefficients. This is evident in the abrupt change of magnitude and behaviour of $\Delta G$ (in figure 10b). In fact while for $\alpha < \alpha_{\text{crit}}$ the full SCBA agrees well with the EST, the two differ sharply as soon as $\alpha > \alpha_{\text{crit}}$. Going into more details we note that $r_1$ is identically zero ($\sim 10^{-14}$) for $\alpha < \alpha_{\text{crit}}$, while $r_2$ is small and negative. Importantly both $r_1$ and $r_2$ are independent of $\{r_\lambda\}_{\text{init}}$. By contrast, for $\alpha > \alpha_{\text{crit}}, r_2$ remains independent of $\{r_\lambda\}_{\text{init}}$ but this is not the case for $r_1$. In fact we obtain a strong dependence over the initial conditions with positive (negative) $r_1$ for positive (negative) $\{r_\lambda\}_{\text{init}}$. Importantly none of these features are found in the case when we neglect the Hartree self-energy.

Since $|r_1|$ is two orders of magnitude larger than $|r_2|$, it will largely determine $\Sigma^H$. As figure 10b shows, $r_1$
varies roughly linearly with $\alpha$ above $\alpha_{\text{crit}}$, therefore the self-consistent $\Delta G$ follows the estimated curve of figure [9] in this region of $\alpha$. The magnitude of $r_1$ for $\alpha > \alpha_{\text{crit}}$ suggests that the interaction with phonons becomes a strong perturbation of the electronic system. We define the region $\alpha < \alpha_{\text{crit}}$ as the weak coupling regime and the region $\alpha > \alpha_{\text{crit}}$ as the strong coupling regime.

Figure [10] adequately explains the causes of the massive reduction of $G(V)$ with respect to its unperturbed value observed in figure [6] at $V = 0$. Notably, as figure [6] shows, $\Delta_{\text{thr}}$, calculated with the full SCBA starts deviating from the EST result for $\alpha \approx 1.75$ eV/Å, i.e. at a value lower that $\alpha_{\text{crit}} = 1.865 \pm 0.005$ eV/Å calculated for $V = 0$. This seems to suggest that the critical value of $\alpha$ for the breakdown of the full SCBA somehow depends on the bias. Moreover at finite bias $\alpha_{\text{crit}}$ is characterized by a peak in $\Delta_{\text{thr}}(\alpha)$ for positive $\{r_\lambda\}_{\text{init}}$ and by a discontinuity for negative $\{r_\lambda\}_{\text{init}}$ (see figure [6]).

In order to explore the onset of the breakdown of the SCBA at finite bias in figure [11] we present $G(V)$ for $\alpha = 1.84$ eV/Å i.e. just below the zero-bias critical value. In addition, in figure [6] we plot the dominant coefficient $r_1$ for a small range of $\alpha$ about $\alpha_{\text{crit}}$ at three different bias, $V = 0$, $V = 0.1$ and $V = 0.2$ Volt. A clear result from figure [11] is that the presence of $\Sigma^H$ introduces a reduction of $G(V)$ with bias not present at $V = 0$. For $\alpha < \alpha_{\text{crit}}$ figure [11] shows that $|r_1|$ is a function of bias whose value increases as the bias increases. The difference between $r_1$ at $V = 0$ and at finite $V$ explains the deviation below $\alpha_{\text{crit}}$ and also the origin of the peak above it.

The discontinuity in $\Delta_{\text{thr}}$ for $\{r_\lambda\}_{\text{init}} > 0$ of figure [6] is explained by the discontinuities observed in $r_1$ (Fig. [11b]) and in the current $J^0(V)$ (Fig. [11]). In fact at $V = 0$, $r_1$ is single valued as long as $\alpha < \alpha_{\text{crit}}$. For $\alpha > \alpha_{\text{crit}}$ instead $r_1(\alpha)$ has a parabolic shape symmetric about $r_1 = 0$: the $\{r_\lambda\}_{\text{init}} < 0$ solution traces out the lower arm of the parabola and the $\{r_\lambda\}_{\text{init}} > 0$ solution follows the upper arm as $\alpha$ increases. For $V > 0$, it is seen that the two solutions for $r_1$ are identical, and asymptotically approach the lower arm of the $V = 0$ curve from below for $\alpha > \alpha_{\text{crit}}$. However, as $\alpha$ is further increased the $\{r_\lambda\}_{\text{init}} > 0$ solution jumps discontinuously above zero and then asymptotically approaches the upper arm, again from below.

The discontinuity in $r_1(\alpha, V)$ is determined by both the bias and the initial conditions. Generally, it is found that such discontinuity occurs for lower bias first; $r_1$ jumps discontinuously for $V = 0.1$ Volt before it does for $V = 0.2$ Volt. This explains the behaviour of the $\{r_\lambda\}_{\text{init}} = \{+0.003125\}$ solution for $J^0(V)$ in figure [11], which leads to a peak in its derivative (the conductance $G(V)$) and explains the discontinuity of $\Delta_{\text{thr}}$.

We note that after the discontinuity in $r_1$ for $V > 0$, the two solutions are no longer symmetric about $r_1 = 0$ and do not converge to the $V = 0$ solutions of either arm until $\alpha \approx 3.0$ eV/Å. This is highlighted in figure [11] where $|r_1|$ for the two solutions is plotted in a range of $\alpha$ just above $\alpha_{\text{crit}}$ at $V = 0.2$ Volt. Such differences explain why $\Delta_{\text{thr}}$ is not independent of the initial conditions beyond the discontinuity and also the different curves observed for $G(V)$ in figure [6].
We make a final comment about the discontinuities seen in figure 11. The value of α at which these occur is dependent on the initial conditions as mentioned already. Thus for a particular \( \{ r_\lambda \}_\text{init} \) it may be possible to reach the upper solution at \( \alpha_{\text{crit}} \) for all bias, so that \( r_1 \) has a parabolic shape for bias while being asymmetric about 0. We have not observed this and regard \( \alpha_{\text{crit}} \) as uniquely defined for \( V = 0 \) only.

V. RESULTS: Au Chains

By using the procedure outlined in section 3 of the full SCBA we used to calculate the transport of the Au RC’s. In general we observe a behaviour similar to that of the H$_2$-Pt system. For 3C, 4C, 5C, and 6C a weak coupling regime is identified where the shift \( \Delta G \) and the weighting coefficients are zero. A critical coupling strength \( \alpha_{\text{crit}} \) for \( V = 0 \) was discovered for each of the chains investigated with values \( \alpha_{\text{crit}} \approx 0.85, 0.9, 0.82, 0.83 \text{ eV/Å} \) respectively for 3C, 4C, 5C, and 6C. For weak coupling \( \Delta G \) matches closely the values calculated in previous works. As an example, in figure 12 we present \( G(V) \) and \( \Delta G(\alpha) \) for 4C. It is seen that the modes symmetric with respect to the centre of the scattering region (even numbered) induce drops in \( G(V) \) at threshold voltages corresponding to the energy of the modes. In general, only the symmetric modes are active in RC’s containing an even number of atoms and conversely the rigid and anti-symmetric modes are active.

For an odd RC.

The transition from weak to strong coupling can be appreciated for the 4C by looking at the inset of figure 12b, where for \( \alpha > \alpha_{\text{crit}} \) two different initial conditions lead to two different \( \Delta G \). The 6C shows the same behaviour of the 4C, while the 3C and 5C show a single curve for \( \Delta G \) due to the symmetry of the weighting coefficients in the strong regime. Beyond \( \alpha_{\text{crit}} \), for all the RC’s simulated, \( G(V) \) is reduced to zero as \( \alpha \) increased and the shift \( \Delta G \) rises to \( G_0 \).

Finally we want to investigate further the existence of multiple solutions depending on the initial conditions in the self-consistent loops. As an example of how convergence is achieved in figure 14 we show the coefficient \( r_4 \) as a function of the iteration number \( n \) for the 4C plotted for a single bias \( V = \) 0.02 Volt and coupling 0.902 eV/Å (\( \alpha > \alpha_{\text{crit}} \)). A number of simulations were run with different initial conditions. The figure indicates the existence of two stable minima: simulations which start at \( \{ r_\lambda \}_\text{init} > 0 \) (dashed lines) converge to the same positive final value, while simulations initialised with \( \{ r_\lambda \}_\text{init} < 0 \) converge to the same \( r_0 < 0 \) value. We note that the minima are not symmetric about \( r_0 = 0 \). The solution \( r_\lambda = 0 \) appears to be a minima in the weak coupling regime, but becomes unstable and evolves to a local maximum in the

![FIG. 12: \( G(V) \) for the 4C illustrating the onset of inelastic processes at threshold voltages. These are associated to the symmetric modes of energy \( \Omega_2 = 6.5 \text{ meV} \) and \( \Omega_4 = 12.2 \text{ meV} \). The rigid and anti-symmetric modes of energies respectively \( \Omega_1 = 3.1 \text{ meV} \) and \( \Omega_3 = 9.8 \text{ meV} \) have no effect for chains comprising an even number of atoms. No overall shift in \( G(V) \) is observed as \( \alpha \) lies below the critical coupling \( \alpha_{\text{crit}} \) which is clearly determined from the inset in panel b.](image)

![FIG. 13: Vibration modes of Au 4C. Modes 2 and 4 are symmetric modes (S) about the centre of the scattering region. Mode 1 is the rigid translational mode (R) while mode 3 is anti-symmetric (A). For all chains considered, the mass of the lead atoms are taken sufficiently larger than that of the chain atoms so that they do not vibrate.](image)

![FIG. 14: Test of the convergence of the weighting coefficient \( r_4 \) for the 4C and \( \alpha > \alpha_{\text{crit}} \) using a range \( \{ r_\lambda \}_\text{init} \), where \( n \) is the number of iterations. Results for the calculations with positive starting values are indicated with dashed lines while those with negative initial values are shown with solid lines. A single negative and positive solution for the converged \( r_4 \) is found.](image)
strong coupling regime.

In figure 15, \( r_0 \) is plotted versus the number of iterations \( n \) for \( \alpha > \alpha_{\text{crit}} \). This time we run different simulations in which the lower bound of the energy integration grid, \( EB \), is changed. In particular we explore situations where \( EB \) is not below the lower band-edge of the leads, that for our choice of parameters lies at -2 eV. Clearly when \( EB \geq -2 \) eV \( r_0 \) does not converge fully, i.e. it depends on \( EB \).

Finally we test the robustness of our integration method. Figure 16 shows the transmission coefficients \( T(\omega) \) for a 6C at a bias of \( V = 1 \) mV where only the sixth mode is considered. We run two simulations. In the first case, the full SCBA is used with the numerical parameters taken from table 1 and initial condition \( \{ r_0 \} = -0.00975 \). In the second case, the integration method outlined in section 11B is replaced by Simpson's rule along the real axis. The integration range used is \( [-3.093, 0.1] \) eV with a grid fineness \( \text{d}E \text{m} = 2.129 \times 10^{-5} \) eV. The Hartree self-consistent loop is started with \( \{ r_0 \} = 0 \) and finished when \( e_\ast < 10^{-12} \) (this tolerance was also used in the first case). All other parameters of the calculations are identical in the two cases. The figure shows that the two numerical methods produce the same \( T(\omega) \), specifically in the range of applied bias shown in the inset.

VI. CONCLUSIONS

By using a simple 1D tight binding model we have investigated the breakdown of the SCBA as a function of the e-p coupling strength \( \alpha \). We have identified two regimes. In the weak coupling regime there is a unique solution for both \( \Sigma^H \) and \( \Sigma^F \), independently of the initial conditions. In particular the Hartree self-energy is small and has little effect on the final conductance. In this weak coupling regime the characteristic conductance drops at voltages corresponding to the various phonon energies compare well with those calculated with the EST method.

As the coupling parameter \( \alpha \) is increased beyond some critical value \( \alpha_{\text{crit}} \), a sharp transition to the strong coupling regime occurs. In this limit the self-consistent \( \Sigma^H \) becomes unstable with respect to the initial conditions and exhibits multiple values for the same voltage. This results in a conductance that sharply deviates from that obtained with the exact EST. Such a sharp transition suggests a breakdown of the SCBA, indicating that the electron-phonon interaction cannot be treated perturbatively for \( \alpha > \alpha_{\text{crit}} \). Interestingly such a breakdown is suppressed when the Hartree self-energy is neglected completely from the calculation, as routinely done in practice. Our results however set a warning to quantitative calculations based on e-p parameters extracted from density functional theory. For these, no information is available on whether or not the obtained e-p parameters would be obtained with the exact EST. Such a sharp transition suggests a breakdown of the SCBA, indicating that the electron-phonon interaction cannot be treated perturbatively for \( \alpha > \alpha_{\text{crit}} \). Interestingly such a breakdown is suppressed when the Hartree self-energy is neglected completely from the calculation, as routinely done in practice. Our results however set a warning to quantitative calculations based on e-p parameters extracted from density functional theory. For these, no information is available on whether or not the obtained e-p parameters would be obtained with the exact EST.

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