Competition between the electronic and phonon–mediated scattering channels in the out–of–equilibrium carrier dynamics of semiconductors: an ab–initio approach

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Abstract. The carrier dynamics in bulk Silicon, a paradigmatic indirect gap semiconductor, is studied by using the Baym–Kadanoff equations. Both the electron–electron (e–e) and electron–phonon (e–p) self–energies are calculated fully ab–initio by using a semi–static out–of–equilibrium GW approximation in the e–e case and a Fan self–energy in the e–p case. By using the generalized Baym–Kadanoff ansatz the two–time evolution is replaced by the only dynamics on the macroscopic time axis. The enormous numerical difficulties connected with a real–time simulation of realistic systems is overcome by using a completed collision approximation that further simplifies the memory effects connected to the time evolution. The carrier dynamics is shown to reduce in such a way to have stringent connections to the well–known equilibrium electron–electron and electron–phonon self–energies. This link allows to use general arguments to motivate the relative balance between the e–e and e–p scattering channels on the basis of the carrier energies.

1. Introduction
Since the seminal works of two chemists, Norrish and Porter (recognized by the 1976 Nobel prize in Chemistry), laser technology has grown explosively until reaching in recent years laser durations of few attoseconds and power flux densities larger than $10^{18}$ W/cm$^2$ [1]. This led to the opening of the new era of femtosecond (fs) nanoscale physics, where combined non–linear and non–adiabatic phenomena [2] occur and where it is possible to monitor real–time the dynamics of photo–induced electronic excitations with unprecedented precision [3].

Ultrafast science at the nano-scale is clearing the path for nanostructure devices with efficient light emission spectra, high optical gain and controllable atomic deformations, thus paving the way to strategic applications in chemistry, biophysics and medicine. The design of nano-scale devices is, however, inevitably linked to a detailed knowledge of the structural and dynamical properties of these systems. Despite the massive number of available experimental results there are still scarce numerical and theoretical methods in use of the scientific community.

Nanostructures and biological systems are, indeed, formed by hundreds/thousands of atoms and their peculiar properties are related to their reduced dimensionality and extended surface. Any reliable theory is inevitably linked to a detailed knowledge of their structural and dynamical
properties. Due to the complexity of these systems, however, state-of-the-art methods are confined to rather simple models with empirical parameters. In this way the theory is deprived of its predictive aspiration, and of the possibility of inspiring new experiments and practical applications. This situation is unavoidably creating a gap between theory and experiment.

Density Functional Theory (DFT)[4] represents the most up-to-date, systematic and predictive approach to study the electronic, structural and optical properties of an impressive wide range of materials[5], including solids and nanostructures. DFT is, however, a ground-state theory that, as it will be shortly discussed in this paper, can only provide a suitable, parameter-free (and thus predictive) basis where the actual real-time simulations are carried on. All the class of methods developed starting from the DFT are labeled as \textit{ab-initio} methods.

It is well known that extended systems have been mostly studied by using the many-body perturbation theory (MBPT) [6] and Time-Dependent Density-Functional-Theory (TDDFT)[7]. The different treatment of correlation and nonlinear effects marks the range of applicability of the two approaches. The real-time TDDFT makes possible to investigate nonlinear effects like second harmonic generation[8] or hyperpolarizabilities of molecular systems[9]. However the standard approaches used to approximate the exchange-correlation functional of TDDFT treat correlation effects only on a mean-field level. As a consequence, while finite systems—such as molecules—are well described, in the case of extended systems—such as periodic crystals and nano-structures—the real-time TDDFT does not capture the essential features of the optical absorption [10] even qualitatively. On the contrary MBPT allows to include correlation effects using controllable and systematic approximations for the self-energy $\Sigma$, that is a one-particle operator non-local in space and time.

2. The equilibrium and out-of-equilibrium dynamics in an \textit{ab-initio} framework

In recent years, the MBPT approach has been merged with DFT by using the Kohn-Sham [7] Hamiltonian as zeroth-order term in the perturbative expansion of the interacting Green’s functions. This approach is parameter free and completely \textit{ab-initio}, [10] and in this paper will be addressed as \textit{ab-initio}-MBPT (Ai-MBPT) to mark the difference with the conventional MBPT.

In practice the Kohn–Sham equation is first solved:

\[
\left[-\frac{\nabla^2}{2} + V_{KS} \{\{R\}\} (r)\right] \phi_{nk} (r) = \epsilon_{nk} \phi_{nk} (r),
\]

with

\[
V_{KS} \{\{R\}\} (r) = V_{\text{ion}} \{\{R\}\} (r) + \int dr' v (r, r') n (r') + V_{xc} ([n], r).
\]

$V_{KS}$ is, thus, written in terms of the bare Coulomb interaction $v (r_1, r_2) \equiv |r_1 - r_2|^{-1}$, the electronic density $n (r)$, the exchange-correlation potential $V_{xc} ([n], r)$ and the total ionic potential $V_{\text{ion}} \{\{R\}\} (r)$:

\[
V_{\text{ion}} \{\{R\}\} (r) = \sum_I V_{\text{ion}} (r - R_I),
\]

with $R_I$ the position of the generic atom $I$. All the simulations presented in this work have been carried on by using the Quantum Espresso[11] and the Yambo[12] by using pseudo-potential approach written in a plane-waves basis. In this approach $V_{\text{ion}} (r - R_I)$ is a pseudo-potential defined only for the valence electrons (8 in the case of bulk Silicon) in order to restrict the electronic dynamics to valence electron only. Therefore core–electrons are not considered in the present work.
By plugging into Eq.3 the precise atomic structure of a given material, the solution of Eq.1 provides a basis for the electronic levels suitable to rewrite the Many–Body problem. Indeed any Green’s function \( G(1,2;\beta) \) can be easily rewritten as a matrix in the KS basis

\[
G(1,2;\beta) = \sum_{n'k} [G_k(t_1,t_2;\beta)]_{n'n'} \phi^*_{n\beta_k}(r_1) \phi_{n'\beta_k}(r_2),
\]

where \( 1 = (r_1,t_1) \) and \( 2 = (r_2,t_2) \) represent global position, time and (eventually) spin indexes and \( \beta = k_BT/h \) the Boltzmann constant and \( T_{ph} \) the equilibrium temperature.

However the \( \beta \)-MBPT is a very cumbersome technique that, based on a perturbative concept, increases its level of complexity with the order of the expansion. As an example, this makes the extension of this approach beyond the linear response regime quite complex, though there have been recently some applications of the \( \beta \)-MBPT in nonlinear optics [13–15].

Another stringent restriction of the \( \beta \)-MBPT is that it cannot be applied when non-equilibrium phenomena take place: for example it cannot be applied to study the light emission after an ultrafast laser pulse excitation. A generalization of MBPT to non-equilibrium situations has been proposed by Kadanoff and Baym[16–19]. In their seminal works the authors derived a set of equations for the real-time Green’s functions, the Kadanoff-Baym equations (KBE’s), that provide the basic tools of the non-equilibrium Green’s Function theory and allowed essential advances in non equilibrium statistical mechanics.

In the equilibrium MBPT, due to the time translation invariance, the relevant variable used to calculate the Green’s functions is the frequency \( \omega \). Instead, out of equilibrium, the time variables acquire a special role and much more attention is devoted to the their propagation properties. The time propagation avoids the explosive dependence, beyond the linear response, of the MBPT on high order Green’s functions. Moreover the KBE’s are non-perturbative in the external field therefore weak and strong fields can be treated on the same footing.

The key ingredient of the KBE’s are the two times lesser/greater Green’s functions\[16–19\] \( G_{n_1n_2k}^{\leq}(t_1,t_2) \) and self–energies \( \Sigma_{n_1n_2k}^{\leq}(t_1,t_2) \). If we consider the time evolution of \( G^{<} \) only on the macroscopic axis \( (t_1 = t_2 = T) \) we get the starting form of the KBE’s I will further analyze in the later sections of this paper:

\[
\frac{i}{\hbar} \frac{\partial}{\partial T} G_k^{<}(T,T;\beta) = \left[ h_k^{KS} + V_k^{Hartree} \right] n(T;\beta) + \Sigma_k^{>}(T;\beta) + U_k(T;\beta) G_k^{<}(T,T;\beta) + S_k^{<}(T;\beta).
\]

In Eq.5 all quantities are matrices in the space of KS states (see Eq.4), e.g., \( \Sigma_{n_1n_2k}^{<} \), \( \beta = (k_BT_{el})^{-1} \) where \( T_{el} \) is the phonon bath temperature, and I have already introduced the fundamental ingredients that will be discussed in the following of this paper:

- The equilibrium KS Hamiltonian, \( h^{KS} \).
- The Hartree potential \( V^{Hartree} \). The ionic and kinetic part is already embodied in \( h^{KS} \).
- The static self–energy \( \Sigma^{>}. \) This is responsible for introducing static e–e correlations that, if properly defined [20] reduce the KBE, in the linear–response regime, to the well–known Bethe–Salpeter equation[6].
- The interaction with the external, arbitrary electric field. The only assumption here is the field wavelength is long compared with the unit cell size (optical limit).
- The interaction kernel \( S^{<}. \) This is the most important part of the equation that embodies dynamical e–e and e–p scatterings. It will be defined and carefully analyzed in Sec.4.

One of the first attempts to apply the KBE’s for investigating optical properties of semiconductors was presented in the seminal paper of Schmitt-Rink and co-workers. [21] Later
the KBE’s were applied to study quantum wells, [22] laser excited semiconductors, [23] and luminescence [24]. However, only recently it was possible to simulate the Kadanoff-Baym dynamics in real-time. [25–29]

Regarding the out-of-equilibrium carrier dynamics there is a vast literature [18] where the KBE’s have been solved using several kind of approximations. I do not want to give here an exhaustive list of all the works in this field, but a key aspect of most of them is that they rely on very strong approximations for what regards the underlying electronic structure and microscopic dielectric properties of the materials used. For example a large number of works use few (two or four) bands models like in Refs.[30–35]. In all these works the electronic levels are approximated as well as the screening of the electron–electron interaction.

In this work I want to go well beyond these approximations by solving the KBE’s in a fully \textit{ab–initio} framework. I will refer to this novel methodology as \textit{ab–initio} Non–Equilibrium Green’s function (\textit{Ai–NEGF}) theory.

3. The equilibrium self–energies

In order to discuss the physics involved in the dynamics of carriers taken out-of-equilibrium it is useful to introduce the properties of the two self–energies that will be used to describe the e–e and e–p coupling. In the next two subsections, indeed, I will introduce the approximations used and the key definitions of the different ingredients needed to build the self–energies. I will later rewrite these equilibrium self–energies in the framework of the \textit{Ai–NEGF} theory.

The e–e interaction will be treated in the so called \textit{GW} approximation[36] where the self–energy is expanded at the first order in the dynamically screened interaction. This approximation has provided a priceless tool to evaluate accurately the electronic levels of a wide class of materials [10].

Since its first application to semiconductors[37] the \textit{GW} self-energy has been shown to correctly reproduce quasi-particle energies and lifetimes for a wide range of materials[36]. Furthermore, by using the static limit of the \textit{GW} self-energy as a scattering potential of the Bethe-Salpeter equation (BSE)[6], it is possible to calculate response functions including electron-hole interaction effects.

In contrast to the e–e case the e–p coupling has received only minor attention in the \textit{ab–initio} community. And this is somehow astonishing if we consider that the e–p coupling is well known to play a key role in several physical phenomena. For example, beside inducing the well–known superconducting phase of several materials [38], it affects the renormalization of the electronic bands[39–43].

Despite the development of more powerful and efficient computational resources the actual calculation of the effects induced by the e–p coupling in realistic materials remains a challenging task. In addition to the numerical difficulties, it has been assumed, for a long time, that this interaction can yield only minor corrections (of the order of meV) to the electronic levels. As a consequence the majority of the \textit{ab–initio} simulations of the electronic and optical properties of a wide class of materials are generally performed by keeping the atoms frozen in their crystallographic positions. More importantly it is well–known that phonons are atomic vibrations and, as a such, can be easily populated by increasing the temperature. This naive observation is \textit{de-facto} used to associate the effect of the electron–phonon coupling to a temperature effect that vanishes as the temperature goes to zero. However this is not correct as the atoms possess an intrinsic spatial indetermination due to their quantum nature, that is independent on the temperature. These quantistic oscillations are taken in account by the e–p coupling when \( T \to 0 \) in the form of a zero–point–motion effect.
The phonon propagator: $D_{q\lambda}^{(0)}(\omega)$

The dynamically screened interaction $W$.

(a) The equilibrium self–energy: $\Sigma_{e-e,eq}$

(b) The out–of–equilibrium self–energy: $\Sigma_{e-e}$

(c) The Fan self–energy: $\Sigma_{e-p}$

**Figure 1.** The self–energy diagrams at the equilibrium and out–of–the–equilibrium used to account for e–e and e–p coupling. The kind of Green’s functions used in the diagrams are shown. Diagram (a) refers to the equilibrium case only, diagram (b) to the out–of–the–equilibrium case while diagram (c) can be used both at and out–of–the equilibrium. This is because in this work the phonons are assumed to be in thermal equilibrium with the external temperature. This means that no phonon dynamics is included. For the e–p case I consider the first order expansion in the atomic displacements corresponding to the Fan self–energy (diagram (c)). In the e–e case, instead, I first consider an equilibrium GW self–energy (diagram (a)) and the corresponding out–of–equilibrium version (diagram (b)) . In the diagram (c) the independent particle polarization corresponds to the dynamical part of the diagram. Indeed the two vertical wiggled lines correspond to a statically screened microscopical e–e interaction (see Eq.(33)).

3.1. The GW approximation for the electronic self–energy

The Feynman diagram corresponding to the GW approximation (GWA) in the equilibrium is showed in the panel (b) of Fig.1. It corresponds to the lowest order contribution to the self–energy expressed as a Taylor expansion in the screened interaction $W(1,2)$:

$$\Sigma_{e-e}^{eq}(1,2) = iW(1^+,2)G_1(1,2).$$

In the present case we will use a spinless formulation where spin indexes are summed out. In Eq.(6) $1 = (r_1,t_1)$ and $2 = (r_2,t_2)$ represent global position and time. In all diagrams showed in Fig.1 the time lies on the real–time axis. The out–of–equilibrium version of the GWA will be introduced in the next section.

**Figure 2.** The equation for the screened interaction $W$ within the RPA.
In the GWA the screened interaction is written as:

\[ W(1,2) = v(1,2) + \int d^3 q v(1,3) \hat{\chi}(3,4) W(4,2), \]  

(7)

with \( v(1,2) \equiv v(r_1, r_2) \delta(t_1 - t_2) \) is the bare Coulomb interaction and \( \hat{\chi} \) is the irreducible electronic response function [6]. An approximation for \( W \) follows by choosing \( \hat{\chi} \) to be calculated in the independent–particle approximation which corresponds, for \( W \), to the random–phase–approximation (RPA), where

\[ \hat{\chi}(1,2) \approx \chi_0(1,2) = -i G_0(1,2) G_0(2,1). \]  

(8)

The actual evaluation of the space dependence appearing in Eq.8 is treated in reciprocal space, by summing on the lattice vectors \( \mathbf{G}, \mathbf{G}' \) and integrating over the Brillouin Zone (BZ):

\[ \chi(r_1, r_2; t) = \int_{BZ} \frac{d \mathbf{q}}{(2\pi)^d} \sum_{\mathbf{G}, \mathbf{G}'} e^{i(\mathbf{q} + \mathbf{G}) \cdot r_1} \chi_{\mathbf{G}, \mathbf{G}'}(\mathbf{q}, t) e^{-i(\mathbf{q} + \mathbf{G}') \cdot r_2}. \]  

(9)

It follows that, within the Random–Phase approximation (RPA):

\[ \chi_{\mathbf{G}, \mathbf{G}'}(\mathbf{q}, \omega) = \chi_0^{0}(\mathbf{q}, \omega) + \sum_{\mathbf{G}''} \chi_0^{0}(\mathbf{q}, \omega) \frac{4\pi}{|\mathbf{q} + \mathbf{G}'|} \chi_{\mathbf{G}, \mathbf{G}'}(\mathbf{q}, \omega). \]  

(10)

Eq.10 is used together with the definition of the single–particle Green’s function

\[ G_0(r_1, r_2; \omega) = 2 \sum_{n} \sum_{\mathbf{k} \in BZ} \phi_{n\mathbf{k}}(r_1) \phi^*_{n\mathbf{k}}(r_2) G_{n\mathbf{k}}(\omega), \]  

(11)

with

\[ G_{n\mathbf{k}}(\omega) = \frac{f_{n\mathbf{k}}}{\omega - \epsilon_{n\mathbf{k}} - i\delta} + \frac{1 - f_{n\mathbf{k}}}{\omega - \epsilon_{n\mathbf{k}} + i\delta}; \]  

(12)

to write the final expression of the GW self–energy. In the present work we are mainly interested in the carrier dynamics and, in order to keep the notation as compact as possible, we restrict our attention on the imaginary part only of \( \Sigma^{e-e}_{\text{el}} \). This, indeed, defines the quasi–particle linewidth as

\[ \Gamma^{e-e, eq}_{n\mathbf{k}} = 3 \left[ \Sigma^{e-e, eq}_{n\mathbf{k}}(\epsilon_{n\mathbf{k}}) \right], \]  

(13)

where we have used the simplest on–the–mass–shell approximation[36]. By doing simple algebraic operations it is possible to rewrite \( \Gamma^{e-e, eq}_{n\mathbf{k}} \) as:

\[ \Gamma^{e-e, eq}_{n\mathbf{k}} = \frac{2\pi}{N\Omega} \sum_{m} \sum_{\mathbf{q}, \mathbf{G}, \mathbf{G}'} \left\{ \tilde{\rho}_{nm}(\mathbf{k}, \mathbf{q}, \mathbf{G}) \right\} \left[ \tilde{\rho}_{nm}(\mathbf{k}, \mathbf{q}, \mathbf{G}') \right]^* \text{Im} \left[ W_{\mathbf{G}, \mathbf{G}'}(\mathbf{q}, \epsilon_{n\mathbf{k}} - \epsilon_{m\mathbf{k} - \mathbf{q}}) \right] \]

\[ \left( 2 - f_{m\mathbf{k} - \mathbf{q}} \right) \theta(\epsilon_{n\mathbf{k}} - \epsilon_{m\mathbf{k} - \mathbf{q}}) - f_{m\mathbf{k} - \mathbf{q}} \theta(\epsilon_{m\mathbf{k} - \mathbf{q}} - \epsilon_{n\mathbf{k}}) \right\}, \]  

(14)

The term \((\text{electron scattering})\) represents an electron with energy \( \epsilon_{n\mathbf{k}} \) that decays into an empty state with lower energy \( \epsilon_{m\mathbf{k} - \mathbf{q}} \) dissipating the corresponding energy difference in positive energy
e–h pairs (represented by $W_{G,G'}(q,\epsilon_{nk} - \epsilon_{mk-q})$). The \textit{(hole scattering)} channel is equivalent to the \textit{(electron scattering)} with the only difference that the hole jumps into a full state with higher energy.

In Eq.14 we have defined

$$\hat{\rho}_{nm}(k, q, G) = \langle nk|e^{i(q+G)\cdot r_1}|mk - q \rangle = \int dr u^*_n(k)(r) u_{mk-q}(r) e^{i(q+G)\cdot r}, \quad (15)$$

where $u_{nk}(r)$ is the periodic part of the Bloch function. We have also introduced the dynamically screened interaction $W_{G,G'}(q,\omega)$ defined as

$$W_{G,G'}(q,\omega) = \frac{(4\pi)}{|q+G|^2}\delta_{G,G'} + \frac{(4\pi)}{|q+G|^2}\chi_{G,G'}(q,\omega) \frac{(4\pi)}{|q+G|^2}. \quad (16)$$

\textbf{3.2. The electron–phonon Fan self–energy}

The phonon contribution to the equilibrium self-energy comes from the oscillations of the atoms around their equilibrium positions. The vibrational states of the system are described, by using the well–known extension of DFT, Density Functional Perturbation Theory (DFPT)[44, 45] where the change in the ionic potential due to a generic atomic displacement is taken into account, self–consistently, with the solution of the KS equation. In the DFPT framework the bare ionic potential is screened by the electronic polarization and the interaction Hamiltonian between electrons and phonons $\hat{H}_{el-at}$ is given by

$$\hat{H}_{el-at} = \int_{\text{crystal}} dr \hat{\rho}(r) \hat{V}_{scf}([R]) (r). \quad (17)$$

In Eq.(17) the self–consistent ionic potential, calculated at the first order in the atomic displacements, is given by[44, 45]

$$\hat{V}_{scf}([R]) (r) = \int dr_1 \left[ \delta(r - r_1) + \int dr_2 f_{Hxc}(r,r_2) \chi(r_2,r_1) \right] V_{ion}([R]) (r_1), \quad (18)$$

where $\chi$ is the TDDFT response function calculated using the $f_{Hxc}$ kernel. In this work I have used the Local Density Approximation for $V_{xc}$ and $f_{Hxc}$. If we now consider a configuration of lattice displacements $u_{Is}$, $H$ can be expressed as a Taylor expansion

$$\hat{H}_{el-at} - \hat{H}_{el-at} = \sum_{I\alpha} \frac{\partial \hat{V}_{scf}([R]) (r)}{\partial R_{I\alpha}} \hat{u}_{I\alpha}, \quad (19)$$

where $\alpha$ is the Cartesian coordinate. Any overlined operator $\overline{O}$ is evaluated with the atoms frozen in their crystallographic equilibrium positions. The link with the perturbative expansion is readily done by transforming Eq. (19) from the space of the lattice displacements to the space of the canonical lattice vibrations by means of the identity[46]:

$$\hat{u}_{I\alpha} = \sum_{q\lambda} (2N_q M_{I}\omega_{q\lambda})^{-1/2} \xi_\alpha(q\lambda|I) e^{i(q\lambda|I)} \left( \hat{b}_{-q\lambda}^\dagger + \hat{b}_{q\lambda} \right), \quad (20)$$

where $N$ is the number of cells (or, equivalently the number of q–points) used in the simulation and $M_I$ is the atomic mass of the $I$–th atom in the unit cell. $\xi_\alpha(q\lambda|I)$ is the phonon polarization vector and $\hat{b}_{-q\lambda}^\dagger$ and $\hat{b}_{q\lambda}$ are the bosonic creation and annihilation operators.
By inserting Eq. (20) into Eq. (19) we get

$$\hat{H} - \hat{H} = \frac{1}{\sqrt{N_q}} \sum_{\text{kin}} g_{n' \lambda}^{q} e^{\dagger_{n'k+q} c_{nk}} (\hat{b}_{-q}^{\dagger} \lambda + \hat{b}_{q}^{\lambda})$$

(21)

In Eq. (21) we have introduced the \((g_{n' \lambda}^{q})\) electron–phonon matrix elements. This is obtained by rewriting \(V_{scf}\) and by making explicit its dependence on the atomic positions:

$$\hat{V}_{scf} \left( [R] \right) (r) = \sum_{Ia} \hat{V}_{scf}^{(s)} (r - R_{Ia}).$$

(22)

By using Eq.22 I get an explicit form for the electron–phonon matrix elements

$$g_{n' \lambda}^{q} = \sum_{s \alpha} \langle \hat{b}_{n'}^{(s)} \lambda (\delta_{n'}^{q} k + q) | \frac{\partial \hat{V}_{scf}^{(s)} (r)}{\partial R_{s \alpha}} (nk) \rangle \xi_{\alpha} (q \lambda | s).$$

(23)

We have used the short form \(R_{s \alpha} = R_{I \alpha I} |_{I = 0}\).

Note that, in principle, the interaction Hamiltonian, Eq.(21), can be also used to obtain a diagrammatic representation of the phonon self-energy. If the electronic and phononic problems is solved at the same time, however, the exchange–correlation effects introduced within the DFPT would lead to severe double counting problems [47]. In the present case this problem is avoided by using bare DFPT phonon modes without taking in account additional Many–Body corrections.

The actual calculation of the Fan diagram is straightforward. The Fan diagram is similar to the GW one where the screened electronic interaction \(W\) is replaced by a phonon propagator of wavevector \(q\) and branch \(\lambda\) [48]. By applying the finite temperature diagrammatic rules it is possible to define the Fan self-energy operator \(\Sigma_{nk}^{e-p} (\omega)\), recovering the expression originally evaluated by Fan[49] for the phonon mediated quasi–particle linewidth

$$\Gamma_{nk}^{e-p, eq} (\beta) = \Im \left[ \Sigma_{nk}^{e-p, eq} (\epsilon_{nk}, \beta) \right],$$

(24)

with

$$\Gamma_{nk}^{e-p, eq} (\beta) = \left| \left[ \Sigma_{nk}^{e-p, eq} (\epsilon_{nk}, \beta) \right]|_{h} \right|_{e} + \left| \left[ \Sigma_{nk}^{e-p, eq} (\epsilon_{nk}, \beta) \right]|_{ph} \right|_{h} + \left| \left[ \Sigma_{nk}^{e-p, eq} (\epsilon_{nk}, \beta) \right]|_{ph} \right|_{h} =$$

$$\frac{\pi}{N} \sum_{n' \lambda} g_{n' \lambda}^{q} \left( 1 - f_{n'q-k} \right) \delta (\epsilon_{n'k-q} - \epsilon_{nk} + \omega_{q}) +$$

$$\frac{\pi}{N} \sum_{n' \lambda} g_{n' \lambda}^{q} \left( f_{n'q-k} \delta (\epsilon_{n'k-q} - \epsilon_{nk} - \omega_{q}) +$$

$$\frac{\pi}{N} \sum_{n' \lambda} g_{n' \lambda}^{q} \left( N_{q \lambda} (T) \left[ \delta (\epsilon_{n'k-q} - \epsilon_{nk} + \omega_{q}) + \delta (\epsilon_{n'k-q} - \epsilon_{nk} - \omega_{q}) \right] \right),$$

(25)

with \(N_{q \lambda} (T)\) is the Bose function distribution of the phonon mode \((q, \lambda)\) at temperature \(T\). The e–p lifetime is splitted in Eq.25 in three different contributions corresponding to the scattering of an electron/hole \(\Gamma_{nk}^{e-p, eq} (\beta) |_{h} \) with the phonon bath and an explicitly temperature term, \(\Gamma_{nk}^{e-p, eq} (\beta) |_{ph} \). The Fan self-energy can be also derived by using the framework of the Heine–Allen–Cardona (HAC) theory [39, 40, 50]. The HAC approach is based on the static Rayleigh–Schrödinger perturbation theory done using the \(\hat{u}_{I \alpha I}\) displacement operators as scalar variables.
Figure 3. Quasiparticle linewidths of bulk Silicon calculated by using the GW approximation for the e–e scattering (green boxes) and the Fan approximation for the e–p scattering (red circles). The two gray areas denote the energy regions (as large as twice the electronic indirect gap $2E_g \approx 1.2$ eV) where, by simple energy conservation arguments, the electronic linewidth are zero by definition. In these energy conditions the e–p contribution is stronger and the corresponding linewidths are larger then the e–e ones. The quadratic energy dependence of the e–e linewidths inverts this trend at about five times the indirect gap.

3.3. Electron–phonon versus electron–electron contribution to the equilibrium electronic lifetimes

From Eq.14 and Eq.25 we can deduce a simple physical property of the equilibrium self–energies. By simple energetic conditions it is clear that in a system with a direct or indirect gap $E_g$ the e–e linewidths $\Gamma_{\text{e–e}}^{\text{e–e},\text{eq}}$ are exactly zero whenever $\epsilon_{nk} < \epsilon_{\text{CBM}} + 2E_g$ or $\epsilon_{nk} > \epsilon_{\text{VBM}} - 2E_g$, with CBM referring to the conduction band minimum and VBM to the valence band maximum. These two energy conditions define the energy ranges represented by gray areas in Fig.3.

On the contrary the e–p linewidths, thanks to the continuum of phonon frequencies, have no zero energy regions. It follows that in the case of low energy electrons or hole the e–p linewidths largely dominate on the e–e part. Only above (or below) the energy forbidden regions the e–e linewidths start growing quadratically, following the Fermi liquid expected behavior, and cross the e–p part when $\epsilon_{nk} = -4$ eV (valence bands) and 6 eV.

4. An out–of–equilibrium formulation of the carrier dynamics including e–e and e–p scatterings

Now that the equilibrium self–energies have been introduced and defined we can move in the out–of–equilibrium regime by giving a precise form to the interaction kernel $S_k(T)$. First of all this is formally defined [16–19] as

$$ S_k^\leq (T; \beta) = \int_{-\infty}^{T} d\tau \left[ \Sigma_k^\leq (T, \tau; \beta) G_k^\leq (\tau, T; \beta) + G_k^\geq (T, \tau; \beta) \Sigma_k^\geq (\tau, T; \beta) \right] . \quad (26) $$
The key point in Eq.26 is that the definition of $\mathbf{S}$ is not closed as it involves two--times functions and both the greater and lesser Green’s functions. This means that one should consider the two--time evolution of the KBE together with the equation of motion for the $\mathbf{G}_k^{\Sigma}$ and, as we will shortly see, for the $\mathbf{G}_k^{(r/a)}$ Green’s functions.

As a two--time evolution of the KBE is, at the present stage, computationally not feasible for the real–time dynamics of complex solids and nanostructures we want to explore a different path based on several key ansatz and approximations.

For the e–e case the extension of the GWA, Eq.7, to the out–of–equilibrium regime is well–known [17]. The diagrammatic form is represented in the frame (b) of Fig.1. What I want to do here is the rewrite this extension in a ab–initio framework introducing some key approximations. First of all I split $\mathbf{S}$ and $\mathbf{S}$ in the e–e and e–p parts:

$$\Sigma_k^{(e-e)}(t_1,t_2;\beta) = \Sigma_k^{(e-e)}(t_1,t_2;\beta) + \Sigma_k^{(e-p)}(t_1,t_2;\beta), \quad (27)$$

$$S_k^{(e-e)}(T;\beta) = S_k^{(e-e)}(T;\beta) + S_k^{(e-p)}(T;\beta). \quad (28)$$

The GWA for $\Sigma^{(e-e)}$ out–of–equilibrium reads:

$$\Sigma_{m_1m_2k}^{(e-e)}(t_1,t_2;\beta) = -\frac{i}{2N\Omega} \sum_{m_1,m_2} \sum_{G_1,G_2} \sum_{q} \{\hat{\rho}_{m_1m_1}(k,q,G_1) [\hat{\rho}_{m_2m_2}(k,q,G_2)]^* \times$$

$$W_{G_1,G_2}^{\Sigma}(q,t_1,t_2;\beta) G_{m_1m_2k-q}^{\Sigma}(t_1,t_2;\beta)\}, \quad (29)$$

with the screened interaction $W^{\Sigma}$ written as

$$W_{G_1,G_2}^{\Sigma}(q,t_1,t_2) = \int_{-\infty}^{t_1} dt_1 \int_{-\infty}^{t_2} dt_2 \sum_{G_3,G_4} W_{G_1,G_3}^{(r)}(q,t_1,t_1;\beta) \times$$

$$L_{G_3,G_4}^{\Sigma}(q,t_1,t_2;\beta) W_{G_4,G_2}^{(a)}(q,t_2,t_2;\beta). \quad (30)$$

In Eq.30 $W^{(r/a)}$ are the retarded/advanced screened interactions and $L^{\Sigma}$ is the independent particle polarization diagram represented by the closed loop in the diagram (b) of Fig.1:

$$L_{G_1,G_2}^{\Sigma}(q,t_1,t_2;\beta) = -\frac{i}{2N\Omega} \sum_{m_2,m_3} \sum_{m_2',m_3'} \{\hat{\rho}_{m_3m_3'}(p,q,G_1) [\hat{\rho}_{m_2'm_2}(p,q,G_2)]^* \times$$

$$G_{m_3m_3'p}^{\Sigma}(t_1,t_2;\beta) G_{m_2'm_2p-q}^{\Sigma}(t_2,t_2;\beta). \quad (31)$$

Before analyzing in detail the temporal dependence of $\Sigma^{(e-e)}$ it is instructive to introduce the out–of–equilibrium e–p self–energy. The extension out–of–equilibrium of the FAN self–energy is straightforward[17]:

$$\Sigma_{m_1m_2k}^{(e-p)}(t_1,t_2;\beta) = -\frac{i}{N} \sum_{m_1,m_2} \sum_{q,\lambda} (g_{m_1m_1k}^{\lambda} g_{m_2m_2k}^{q} D_{\lambda}^{\Sigma}(q,t_1,t_2;\beta) G_{m_1m_2k-q}^{\Sigma}(t_1,t_2;\beta)\}, \quad (32)$$

with $D_{\lambda}$ the phonon propagator that I will shortly define.

Now I introduce the complete set of approximations I will use to transform Eq.29 in a simpler set of equations of motion to describe, ab–initio, the carrier dynamics in realistic materials. These approximation are guided by simple physical arguments introduced with the aim of
limiting the computation effort. This is a crucial requirement to make the simulations feasible. At the same time those approximations represent a first step in the device of a sound and complete framework for the $\Delta r$-NEGF approach and their validity will be carefully analyzed in the further development of this field.

The approximations are:

(i) The screened interaction entering Eq.30 are assumed to be static. I introduce this semi-static approximation (SSA) of the out-of-equilibrium version of the GWA being inspired by the well-known Coulomb-Hole plus Screened-Exchange approximation (COHSEX)[10]. The SSA reads:

$$ W^{(r/a)}_{G_1G_2}(q,t_1,t_2;\beta) \approx W_{G_1,G_2}(q). $$

This approximation is well motivated in semiconductors and insulators as an electronic gap of 1 eV corresponds to a time scale of 0.66 fs. This a very short time compared to the time scale of the simulations presented in the present work.

(ii) The simulation is done at a fixed temperature $\beta$ that is assumed to not change during the simulation. This approximation corresponds to take the phonon population at the equilibrium and assuming that their energy distribution does never deviate from a Bose distribution. In practice this means that:

$$ D^{<}_{\lambda}(q,t_1,t_2;\beta) \approx (-i) \sum_{\pm} N^{\pm}_{q\lambda}(\beta) e^{\pm i \omega_{q\lambda}(t_1-t_2)}. $$

Note that $D^{>}_{\lambda}(q,t_1,t_2;\beta) = - [D^{<}_{\lambda}(q,t_1,t_2;\beta)]^*$.

(iii) I use the Generalized Baym–Kadanoff ansatz (GBKA) [17–19].

(iv) I neglect the off-diagonal matrix elements of the $G_k$ and $\Sigma_k$ matrices

$$ G^{<}_{nn'lk}(T,T;\beta) \approx i\delta_{nn'} f_{nk}(T;\beta), $$

$$ G^{>}_{nn'lk}(T,T;\beta) \approx i\delta_{nn'} [f_{nk}(T;\beta) - 1], $$

$$ \Sigma^{<}_{nn'lk}(T,T;\beta) \approx \delta_{nn'} \Sigma_{nk}(T;\beta). $$

As it will be showed in the following these approximations will make possible to perform the actual simulations of complex materials at a reasonable computational cost.

4.1. The collision operator in the Generalized Baym–Kadanoff ansatz

The GBKA provides the most important tool to disentangle the complex two-times evolution by rewriting:

$$ G^{<}_{nk}(t_1,t_2;\beta) \approx \langle i \rangle \left[ G^{(r)}_{nk}(t_1-t_2;\beta) G^{<}_{nk}(t_2,t_2;\beta) - G^{<}_{nk}(t_1,t_2;\beta) G^{(a)}_{nk}(t_1-t_2;\beta) \right], $$

that, in the lesser case, acquires a simple form written in terms of the time-dependent electronic occupations:

$$ G^{<}_{nk}(t_1,t_2;\beta) \approx - \left[ G^{(r)}_{nk}(t_1-t_2;\beta) f_{nk}(t_2;\beta) - f_{nk}(t_1;\beta) G^{(a)}_{nk}(t_1-t_2;\beta) \right]. $$

A similar identity holds for the greater Green’s function. By using Eq.36 the KBE’s for the electronic occupations $f_{nk}(T,\beta)$ acquire a closed form if a given analytic form for the advanced/retarded Green’s functions is given. But before discussing the specific form of the $(r/a)$ Green’s functions I use Eq.36 to rewrite the two contributions to $S$. 

11
The algebraic passage to rewrite \( S_{e-p, <}^{(\alpha)} \) by using the GBKA can be found, for example, in Ref.[51] in the simpler case of a Jellium model with e–p interaction treated using a Fröhlich interaction \( g_{nmk}^{\lambda} \approx g_{ql} \) and a single optical phonon mode.

After some involved but conceptually simple algebra it can be shown that

\[
S_{nk}^{e-p, <} (T; \beta) = N^{-1} \sum_{q\lambda} \sum_{s=\pm} e^{i\omega q\lambda T} \left\{ S_{nk}^{q\lambda s} (T; \beta) + \left[ S_{nk}^{q\lambda -s} (T; \beta) \right]^* \right\},
\]

(38)

with

\[
S_{nk}^{q\lambda s} (T; \beta) = (-i) \int_{-\infty}^{T} dt e^{-i\omega q\lambda t} \sum_m |g_{nmk}|^2 \sum_{m} G_{nk}^{(r)}(T - \tau; \beta) \times \left\{ N_{q\lambda}^{<} (\beta) \left[ (2 - f_{mk-q} (\tau; \beta)) f_{nk} (\tau; \beta) - N_{q\lambda}^{s} (\beta) f_{mk-q} (\tau; \beta) (2 - f_{nk} (\tau; \beta)) \right] \times \left[ G_{nk}^{(a)} (\tau; \beta) \right]^* \right\}.
\]

(39)

As it will be clear in the next section the sum of \( S_{nk}^{q\lambda s} \) and of its adjoint in Eq.38 is crucial to keep the time-evolution unitary in such a way that the number of electrons is conserved. The actual time decay of the photo-excited carriers will, then, be connected to the actual memory of the system linked with the \( \tau \) integral appearing in Eq.38 rather that with the imaginary part of the self–energy, as commonly done in equilibrium theories. This detail marks the basic physical difference between the equilibrium QP linewidths and the out–of–equilibrium QP decay time, as it will be clear from the discussion in Sec.5.

In the case of the e–e scattering the application of the GBKA is more involved as it must be applied to three Green’s functions instead of one, like in the e–p case. Nevertheless, once a simple form for \( G^{(r/a)} \) will be given the physical interpretation of the collision operators will be equally simple. By following a procedure similar to the e–p case we find that

\[
S_{nk}^{e-e, <} (T; \beta) = \frac{1}{2 (N!)^2} \sum_{q\lambda} \sum_{m1m2m3} \left| W_{nm1k}^{q} \right|^2 \int_{-\infty}^{T} dt \left\{ \left[ G_{m1k-q}^{(r)} (T - \tau; \beta) G_{m2p-q}^{(r)} (T - \tau; \beta) G_{nk}^{(a)} (T - \tau; \beta) + G_{m1k-q}^{(a)} (T - \tau; \beta) G_{m2p-q}^{(a)} (T - \tau; \beta) \right] \times \left( (2 - f_{m1k-q} (\tau; \beta)) (2 - f_{m3p} (\tau; \beta)) f_{m2p-q} (\tau; \beta) f_{nk} (\tau; \beta) - f_{m1k-q} (\tau; \beta) f_{m3p} (\tau; \beta) (2 - f_{m2p-q} (\tau; \beta)) (2 - f_{nk} (\tau; \beta)) \right) \right\}.
\]

(40)

Eq.38–40 represent the expressions for the time–dependent kernel \( S \) that I will further simplify in the next section. By using this expression for \( S \) the final equation of motion for the electronic occupation reads:

\[
\frac{\partial}{\partial T} f_{nk} (T; \beta) = \frac{\partial}{\partial T} f_{nk} (T; \beta) \bigg|_{coh} - S_{nk}^{<} (T; \beta).
\]

(41)

with

\[
\frac{\partial}{\partial T} f_{nk} (T; \beta) \bigg|_{coh} = - \left[ h_{k}^{KS} + V_{k}^{\text{Hartree}} [n (T; \beta)] + \Sigma_{k}^{\ast} (T; \beta) + U_{k} (T; \beta), G_{nk}^{\ast} (T, T; \beta) \right]_{nn},
\]

(41')
4.2. The Completed Collision Approximation

Thanks to the GKBA the KBE’s can be rewritten as a set of non-linear differential equations for the occupation functions. Nevertheless the whole history of these functions should be integrated numerically in Eqs.39–40. In order to further simplify the theory I introduce two more approximations to the list presented in Sec.2:

(v) The $G^{(r/a)}$ functions are assumed to have a QP form (exponential):

$$G_{nk}^{(r)} (T; \beta) \approx -ie^{-iE_{nk} (\beta)T - \Gamma_{nk}(\beta)T},$$

with $G_{nk}^{(a)} (T; \beta) = \left[ G_{nk}^{(r)} (-T; \beta) \right]^*$. This means that we neglect any deviation from the pure exponential behavior induced by the short–time dynamics\cite{52} and the e–e and e–p correlations\cite{53}. In eq.42 the QP energies $E_{nk}$ and widths $\Gamma_{nk}$ are calculated ab–initio by means of the equilibrium GWA and Fan self–energies. The explicit expressions for $\Gamma_{nk}$ are given by Eq.14 and Eq.25.

(vi) I adopt the so called Completed Collision Approximation (CCA)\cite{51} that makes possible to solve the time–integration appearing in Eqs.39–40 analytically by assuming that the occupation functions vary only slowly on the time scale defined by $G_{nk}^{(r/a)} (T - \tau; \beta)$. This time scale is given by $\Gamma_{nk}(\beta)^{-1}$ and, as shown in Fig.3, it is of the order of 10–20 fs. Under this assumption the $f$ functions can be taken out of the integrals appearing in Eqs.39–40 and the rest of the integral can be done analytically by using Eq.42.

Thanks to the CCA Eqs.39 can be rewritten as:

$$S_{nk}^{q\lambda s} (T; \beta) \approx (-i) \sum m |g_{nmnk}|^2 \left[ N^{q\lambda s}_{q\lambda} (\beta) (2 - f_{mk-q} (T; \beta)) f_{nk} (T; \beta) - N^{q\lambda s}_{q\lambda} (\beta) f_{mk-q} (T; \beta) (2 - f_{nk} (T; \beta)) \right] \times \left\{ \int_{-\infty}^{T} d\tau e^{-i\omega_{q\lambda} \tau} G_{mk-q}^{(r)} (T - \tau; \beta) [G_{nk}^{(a)} (T; \beta)]^* \right\}. \quad (43)$$

By using Eq.42 the $\tau$ integral reads:

$$\int_{-\infty}^{T} d\tau e^{-i\omega_{q\lambda} \tau} G_{mk-q}^{(r)} (T - \tau; \beta) [G_{nk}^{(a)} (T; \beta)]^* = \frac{e^{-i\omega_{q\lambda} T}}{E_{mk-q} (\beta) - E_{nk} (\beta) - i\omega_{q\lambda}}. \quad (44)$$

Thanks to Eq.44 the $S_{nk}^{q\lambda s} (T; \beta)$ functions can be rewritten in terms of three simple lifetimes: $\gamma_{nk}^{(e-p,h)} (T; \beta)$, $\gamma_{nk}^{(e-p,e)} (T; \beta)$ and $\gamma_{nk}^{(e-p,ph)} (T; \beta)$:

$$S_{nk}^{q\lambda s} (T; \beta) \bigg|_{CCA} = -\gamma_{nk}^{(e-p,h)} (T; \beta) [1 - f_{nk} (T; \beta)] + \gamma_{nk}^{(e-p,e)} (T; \beta) f_{nk} (T; \beta) + \gamma_{nk}^{(e-p,ph)} (T; \beta). \quad (45)$$

In order to define and interpret the physical properties of these lifetimes I introduce two functions: $P_{abs}$ and $P_{emit}$.

$$P_{abs}^{nn'kq} (\beta) = \frac{|\Im E_{nk} (\beta)| + |\Im E_{nk-q} (\beta)|}{|\Re (E_{nk-q} (\beta) - E_{nk} (\beta)) - \omega_{q\lambda}|^2 + [|\Im E_{nk} (\beta)| + |\Im E_{nk-q} (\beta)|]^2}, \quad (46)$$

and

$$P_{emit}^{nn'kq} (\beta) = \frac{|\Im E_{nk} (\beta)| + |\Im E_{nk-q} (\beta)|}{|\Re (E_{nk-q} (\beta) - E_{nk} (\beta)) + \omega_{q\lambda}|^2 + [|\Im E_{nk} (\beta)| + |\Im E_{nk-q} (\beta)|]^2}. \quad (47)$$
These two functions represent the energy constrain in the emission ($P^{\text{emit}}$) or absorption ($P^{\text{abs}}$) of a phonon smeared out due to the QP approximation for the retarded Green’s function that. This, indeed, yields a Lorentzian indetermination in the energy conservation. By using these two simple functions the three e–p induced lifetimes can be easily rewritten as

$$\gamma_{nk}^{(\text{e–p},h)} (T; \beta) = \sum_{n'q\lambda} \frac{|g_{n'q\lambda}|^2}{N} P_{n'n'kq}^{(\text{abs})} (\beta) f_{n'k-q} (T; \beta), \quad (48)$$

$$\gamma_{nk}^{(\text{e–p},e)} (T; \beta) = \sum_{n'q\lambda} \frac{|g_{n'q\lambda}|^2}{N} P_{n'n'kq}^{(\text{emit})} (\beta) (2 - f_{n'k-q} (T; \beta)), \quad (48')$$

$$\gamma_{nk}^{(\text{e–p},ph)} (T; \beta) = \sum_{n'q\lambda} \frac{|g_{n'q\lambda}|^2}{N} N_{q\lambda} (\beta) \left[ f_{nk} (T; \beta) - f_{n'k-q} (T; \beta) \right] \left[ P_{nn'kq}^{(\text{abs})} (\beta) + P_{nn'kq}^{(\text{emit})} (\beta) \right]. \quad (48'')$$

The dynamics induced by the e–p kernel on the electronic occupations can be easily visualized by using Eq.45 to simplify Eq.5 for the diagonal matrix elements of $G^<$. If I do not consider the coherent part of the KBE equation I get

$$\frac{\partial}{\partial T} f_{nk} (\beta; T) = \gamma_{nk}^{(\text{e–p},h)} (T; \beta) [2 - f_{nk} (T; \beta)] - \gamma_{nk}^{(\text{e–p},e)} (T; \beta) f_{nk} (T; \beta) - \gamma_{nk}^{(\text{e–p},ph)} (T; \beta). \quad (49)$$

Eq.49 makes the interpretation of the carrier dynamics induced by the e–p coupling straightforward. We start indeed noticing that, simply because of their definition, $\gamma_{nk}^{(\text{e–p},h)}$ and $\gamma_{nk}^{(\text{e–p},e)}$ are always positive. In contrast $\gamma_{nk}^{(\text{e–p},ph)}$ can be positive or negative.

To see why this property is important let’s consider two limiting cases when $f_{nk} (\beta; T = 0) = 0$ or $f_{nk} (\beta; T = 0) = 1$. In the first case, from Eq.49 it follows that, if we consider the zero temperature case,

$$f_{nk} (\beta \to \infty; T = 0) = 0 \Rightarrow \frac{\partial}{\partial T} f_{nk} (\beta \to \infty; T = 0) > 0, \quad (50)$$

$$f_{nk} (\beta \to \infty; T = 0) = 2 \Rightarrow \frac{\partial}{\partial T} f_{nk} (\beta \to \infty; T = 0) < 0. \quad (50')$$

This simply means that an initially completely filled level will be emptied by the e–p interaction while an initially empty level will be filled. The $\gamma_{nk}^{(\text{e–p},ph)}$ is non zero only at finite temperature and works to build up a finite temperature quasi–equilibrium among the carrier occupations. The $\gamma_{nk}^{(\text{e–p},h)}$ lifetimes represent a single scattering of the state $|nk\rangle$ to the state $|nk-q\rangle$ with emission/absorption of a phonon mode ($q\lambda$). This process is similar to the one studied in the equilibrium case and, indeed, if we consider the Lorentzian function appearing in $P^{\text{abs}}$ and $P^{\text{emit}}$ as approximated delta functions we see that the analytic form of $\gamma_{nk}^{(\text{e–p},h)}$ are proportional to two terms appearing in Eq.25. Nevertheless, as we will clearly see in the case of bulk silicon, the dynamics induced by the KBE equation is a cascade process where many phonons are emitted or adsorbed and the final decay time of the electronic levels will be very different from the equilibrium case.

By using approximations (vi) and (vii) the e–e kernel, Eq.40, can be worked out in a way similar to the e–p case. Although the math is more involved the final outcome is quite similar
to the e–p case. Indeed it turns out that
\[
S_{nk}^{e-e,c} (T; \beta) \bigg|_{CCA} = - \xi_{nk} (e,e,h) (T; \beta) [2 - f_{n(k)} (T; \beta)] + \xi_{nk} (e,e) (T; \beta) f_{n(k)} (T; \beta).
\]
(51)

The two additional lifetimes \( \xi_{nk} (e-e,e/h) \) are defined to be:
\[
\xi_{nk}^{e-e,e} (T; \beta) = 4 (N \Omega)^{-2} \sum_{qp} \sum_{m_1 m_2 m_3} \left| W_{q,m_1}^{nm_1,k} \right|^2 R_{m_1 k}^{q} \left( 2 - f_{m_1 k-q} (T; \beta) \right) \left( 2 - f_{m_3 p} (T; \beta) \right) f_{m_2 p-q} (T; \beta),
\]
(52)
and
\[
\xi_{nk}^{e-e,h} (T; \beta) = 4 (N \Omega)^{-2} \sum_{qp} \sum_{m_1 m_2 m_3} \left| W_{q,m_1}^{nm_1,k} \right|^2 R_{m_1 k}^{q} f_{m_1 k-q} (T; \beta) \left( 2 - f_{m_2 p} (T; \beta) \right). \]
(52')

We have introduce the function \( R \) defined as:
\[
R_{nm_1 k}^{q} = \left| \Im E_{m_1 k} (\beta) \right| + \left| \Im E_{m_3 p} (\beta) \right| + \left| \Im E_{m_2 p-q} (\beta) \right| + \left| \Im E_{n k} (\beta) \right|
\]
\[
\{ \left[ \Re (E_{m_1 k-q} (\beta) + E_{m_3 p} (\beta) - E_{m_2 p-q} (\beta) - E_{n k} (\beta)) \right]^2 + \left| \Im E_{m_1 k-q} (\beta) \right| + \left| \Im E_{m_3 p} (\beta) \right| + \left| \Im E_{m_2 p-q} (\beta) \right| + \left| \Im E_{n k} (\beta) \right| \}^{-1}.
\]
(53)

The physical interpretation of the e–e assisted carrier dynamics is very similar to the e–p case. The only difference is that, instead of phonons, the carrier decay is due to the creation of e–h pairs. The basic transition that contributes to the e–e scattering is a four bodies process: the initial electron (or hole) scatters in an e–h pair plus another electron (or hole). If we distinguish between the bath electrons and the optically pumped ones (the carriers) we see that these four bodies can, quite generally, be distributed in both group of states.

Nevertheless in the case of semiconductors and insulators it is meaningful to distinguish between intra–carrier and extra–carrier channels. In the first case the e–h pair is created among the optically pumped carriers while in the second case the pair is created across the gap involving the bath electrons. Of course these two channels, in general, contribute at the same time to the overall dynamics. But, as we will see in Sec.5 their relative strength depends on the carrier energy.

In conclusion the total rate equation governing the carrier dynamics that will be used in the Sec.5 is given by Eq.41 with \( S^c \) defined by Eqs.45 and 51. I coded this equation in the Yambo code[12], in the same numerical framework where the equilibrium GW and Fan self–energies are coded. The existing structure of Yambo has already produced several publications and it provides, therefore, the most appropriate environment to perform out–of–equilibrium simulations in a controllable manner. The e–p matrix elements and phonon modes are calculated, instead, with the Quantum Espresso package[11].

4.3. A detailed balance condition to ensure that the number of electrons is conserved

In the case of e–e scattering the actual possibility of creating high energy electron–hole (e–h) pairs can modify, during the real–time dynamics, the number of carriers that fill the conduction
bands with respect to the pairs initially photo–excited. This process is particularly evident for carriers created with energy larger than \( 2E_g \) below the VBM, or \( 2E_g \) above the CBM.

In the e–p case, instead, the maximum available energy that any carrier can dissipate during the relaxation is the Debye energy that, in the systems I consider in this work, is much smaller than the gap. In this case, indeed, it is important that the theory preserves the correct number of photo–excited electrons during the whole simulation.

I use this example to discuss a methodological aspect embodied in the use of a fully \textit{ab–initio} framework. Indeed the e–p matrix elements \( q_{n'nk}^\lambda \equiv g_{n'nk}^{k-p} \), defined in Eq.23, can be written as

\[
g_{n'nk}^{k-p} = \sum_{sa} (2M_s\omega_{k-p})^{-1/2} e^{i(k-p)m_s} (p'|p|) \frac{\partial V_{\text{scf}}(r)}{\partial R_{sa}} |n\rangle_0 \xi_\alpha (k - p, \lambda | s) .
\]

We can, now, consider the KBE for the two isolate states \(|n\rangle\) and \(|mp\rangle\) that are scattered one into the other by means of the \( S_{n'nk}^{e-p,<} (T; \beta) \). For simplicity I only consider the dynamics induced by the e–p kernel at zero temperature and for a single phonon band \( \mu \). This dynamics can be rewritten in terms of four lifetimes: \( \gamma_{n'k}^{(h)} \), \( \gamma_{mp}^{(e)} \):

\[
\gamma_{n'k}^{(h)} (T) = \frac{g_{n'nk}^{k-p,\mu}}{N} P_{nmk(k-p)\mu}^{(abs)} f_{mp} (T) ,
\]

\[
\gamma_{n'k}^{(e)} (T) = \frac{g_{n'nk}^{k-p,\mu}}{N} P_{nmk(k-p)\mu}^{(emit)} (2 - f_{mp} (T)) ,
\]

\[
\gamma_{mp}^{(h)} (T) = \frac{g_{nmk}^{p-k,\mu}}{N} P_{nmn(p-k)\mu}^{(abs)} f_{nk} (T) ,
\]

\[
\gamma_{mp}^{(e)} (T) = \frac{g_{nmk}^{p-k,\mu}}{N} P_{nmn(p-k)\mu}^{(emit)} (2 - f_{nk} (T)) .
\]

Now we notice that, thanks to simple symmetry arguments it can be shown that

\[
g_{nmk}^{p-k,\mu} = \left[ g_{n'nk}^{k-p,\mu} \right]^* .
\]

Moreover from Eqs.46–47 it follows that \( P_{nmk(k-p)\mu}^{(emit)} = P_{nmn(p-k)\mu}^{(abs)} \) and, consequently, from Eqs.55 we see that:

\[
\frac{\partial}{\partial T} f_{nk} (T) = \frac{g_{n'nk}^{k-p,\mu}}{N} P_{nmk(k-p)\mu}^{(emit)} (2 - f_{mp} (T)) (2 - f_{nk} (T))
\]

\[
- \frac{g_{n'nk}^{k-p,\mu}}{N} P_{nmk(k-p)\mu}^{(abs)} f_{mp} (T) f_{nk} (T) ,
\]

and

\[
\frac{\partial}{\partial T} f_{mp} (T) = \frac{g_{nmk}^{p-k,\mu}}{N} P_{nmk(k-p)\mu}^{(abs)} (2 - f_{mp} (T)) (2 - f_{nk} (T))
\]

\[
- \frac{g_{nmk}^{p-k,\mu}}{N} P_{nmk(k-p)\mu}^{(emit)} f_{mp} (T) f_{nk} (T) .
\]
By summing Eq.57 and Eq.58 one gets

$$\frac{\partial}{\partial T} (2 - f_{mp}(T) - f_{nk}(T)) =$$

$$- \left[ \frac{g_{nmk}}{N} \right]^2 \left[ f^{(abs)}_{nmk}(k-p) + f^{(emit)}_{nmk}(k-p) \right] (2 - f_{mp}(T) - f_{nk}(T)).$$ (59)

But from the boundary condition $(2 - f_{mp}(T = 0) - f_{nk}(T = 0)) = 0$ it follows $\frac{\partial}{\partial T} (2 - f_{mp}(T) - f_{nk}(T)) = 0$. The total number of electrons in the two levels made scattering by the e–p interaction is conserved, as it should be.

Eq.56 thus represents a sort of detailed balance condition that ensures that the fraction of charge lost by a given state $|nk\rangle$ is exactly compensated by the charge added to the $|mp\rangle$ state. In total there is no nonphysical charge lost or increase. Of course the level of accuracy of this detailed balance condition is ensured by the numerical precision under which Eq.56 is satisfied. This precision must be carefully checked and, eventually imposed, as on long–time simulations (of the order of picoseconds) tiny deviations can induce important deviations of the total number of electrons conservation.

5. An application in a paradigmatic solid: bulk silicon

After all the math and approximations introduced in the previous sections it is instructive to apply all the machinery composed by the KBE’s written in the ab–initio framework to perform realistic simulations. The paradigmatic case I consider is bulk silicon. This a quite simple material that, nevertheless, has an interesting property: the minimal gap is indirect.

From the point of view of the carrier dynamics this is an important aspect. Indeed the optical gap is located at the $\Gamma$ point and it is larger ($\sim 3.2 \, eV$) of the indirect gap ($\sim 1.2 \, eV$). Thus if an electron is optically pumped in the conduction states this will populate the energy region surrounding the CBM, that is the $\Gamma_15$ state, and the relaxation is expected to occur along the $\Gamma X$ line (see the upper frame of Fig.4) towards the $X_1$ point.

This has been, indeed, experimentally measured[54] by using Time–Resolved Two–Photon Photoemission (2PPE) and the time–decay of a level measured along the $\Gamma X$ has been found to be around 40 fs.

I do not want, however, to interpret and reproduce the 2PPE experiment but to use bulk silicon to illustrate the relative balance of the e–e and e–p channels in the carrier dynamics on the basis of the initial carrier energy and density. I will also briefly discuss the effect of the scattering with acoustic phonons keeping in mind the large number of simulations done using simple optical–only phonon models.

The simulation is performed by defining an ad–hoc initial condition for the occupations and let the system evolve by means of Eq.5. In the first series of calculations I take

$$f_{nk}(t = 0; \beta) = 2 \quad \text{when} \quad |nk\rangle = |\Gamma_15\rangle.$$ (60)

Two electrons (to account for the two spin channels) are moved by hand in two out of the three states that are energetically degenerate with the $\Gamma_15$ state. This corresponds to an initial carried density of $3.2 \times 10^{21} \, cm^{-3}$. As the $\Gamma_15$ state is at the border of the $2E_g$ threshold its extra–carriers contribution to the e–e lifetime is very small. This is because the energy of the state is not enough to create e–h pairs across the minimum gap. The real–time occupations of three paradigmatic states: $|\Gamma_15\rangle$, $|L_1\rangle$ and $|X_1\rangle$ are showed in Fig.4. In the k–grid I have used the $X_1$ state is the nearest to the CBM.

In Fig.4 the functions $f_{\Gamma_15}(T; \beta)$ (red curves), $f_{L_1}(T; \beta)$ (blue curves) and $f_{X_1}(T; \beta)$ (green curves) are showed comparing the dynamics with only the e–p scattering (continuous lines)
Figure 4. Time dependent occupations (lower frames) $f_{\nu,k}(T;\beta)$ as a function of the $T$ for two external fixed temperature (0 K left frame and 300 K right frame) for three selected states in the band-structure of Silicon (upper frame). At $T = 0$ only the $\Gamma_{15}$ is artificially populated with 4 electrons needed to fill 2 of the 3 energy degenerated states. This explains why the $y$-axis goes from 0 to 4. In the lower frames the dynamics is done including only e–p scattering (continuous lines) and e–e plus e–p scattering (dashed lines). In this case, as shown in Fig.3, the energy of the $|\Gamma_{15}\rangle$ state is below the $2E_g$ threshold. This means, indeed, that at $T = 0$ the e–p scattering is largely dominant over the e–p one. But, as the time increases and the initial charge is fragmented in lower energy states the intra–carriers e–e scattering quickly increases leading to a much faster population of the $X_1$ state.

and with the e–p and the e–e scatterings (dashed lines) included. We immediately see some clear trends. The decay if the $\Gamma_{15}$ is very fast, with an exponential time–dependence with a lifetime of the order of $12 \, fs$. This value is very similar to the value of $16 \, eV$ obtained by calculating $\Gamma_{\Gamma_{15}}^{e-p,eq},(\beta \rightarrow \infty)$ showing that the e–e contribution is practically zero. As the time increases the initial charge is fragmented and lower state are populated. At this point the intra–carriers contribution to the e–e is rapidly increased and the contribution to the filling is largely underestimated.

The message is thus clear. In the very short time the dynamics of carriers with energy below the $2E_g$ threshold is e–p dominated. Nevertheless in this regime the use of the CCA is highly questionable and a more careful full–memory treatment is required. As time increases also in this energy range the intra–carriers contribution to the carrier dynamics makes the e–e scattering dominant compared to the e–p channel.

An important aspect that must be taken in account in the analysis of the time–dependencies showed in Fig.4 is that the overall filling of the $X_1$ and $L_1$ states is enhanced when the e–e scattering is included also because of the promotion of additional electrons from the valence to the conduction by the creation of e–p pairs across the gap. This process, although energetically impossible for the $\Gamma_{15}$ state, is made possible by the energy indetermination caused by the use
of the QP approximation for the retarded Green’s function. Indeed thanks to the Lorentzian line–shape, the \( R \) functions (see Eq.53) can be non–zero for arbitrary values of the energy levels involved in the e–e scattering. This is, in my opinion, a stringent limitation imposed by the QP approximation that can be easily avoided by using more refined approximations or by calculating the equilibrium spectral functions numerically in the GW and Fan approximations.

By looking at the two lower frames of Fig.4 we see that the increase of the temperature from 0 to 300 K does not appreciably change the occupation dynamics in the first 500 fs. This is reasonable as the e–e lifetimes are not explicitly temperature sensitive and, as far as the e–p channel is considered, the Silicon Debye temperature is as large as 645 K. Thus at room temperature only the low–energy acoustic modes are populated. Although those contribute for half of the e–p lifetime the overall effect on the dynamics is marginal.

- **Figure 5.** Time dependent occupations (right frame) \( f_{nk}(T;\beta) \) as a function of the \( T \) for an external vanishing temperature for three selected states in the band–structure of Silicon (left frame). At \( T = 0 \) only the \( \Gamma_{15} \) is artificially populated with 4 electrons needed to fill 2 of the 3 energy degenerated states. In the right frame the dynamics is done only including the e–p scattering but including only optical phonon modes (continuous lines) or all modes (dashed lines). We see that the effect of the acoustic modes is not at all negligible, especially for the \( X_1 \) state filling.

- **Figure 6.** Time dependent occupation as a function of the \( T \) for the set of electronic states with energy 5.76 eV, well above the \( 2E_g \) threshold energy. The simulation is performed with the e–p scattering and including (dashed line) and not including (continuous line) the e–e contribution. For this high energy level the equilibrium e–e and e–p lifetimes are similar and both contribute to the decay of the state. This contrasts the \( \Gamma_{15} \) case where the e–p scattering dominates in the short–time dynamics.
An important issue that can be fully exploited by the present calculations is, indeed, the role played by the acoustic phonon modes that, in model calculations, are often neglected. As shown in Fig. 5, the acoustic modes, even at zero temperature, contribute for more than half of the filling of the $X_1$ state. Their contribution to the $\Gamma_{15}$ state is, instead, minor and it slowly increase as the energy of the state approaches the CBM. This phenomena is connected with the physics of both the e–e and e–p induced relaxation of the carriers. This is indeed a cascade process where the initial $\Gamma_{15}$ electrons jump towards other states by the emission of, initially, a single phonon and a single e–h pair. This elementary process is very fast, of the order of even faster of 10 fs. But in order to reach the $X_1$ state the electron undergoes several of these elementary processes and the overall relaxation is a much slower process.

The last aspect I want to analyze by using bulk Silicon as paradigmatic system is the relative influence of the e–e and e–p scattering channels in the short–time regime for carriers pumped well above the $2E_g$ threshold. To this end I change the boundary condition given by Eq. 60 to be

$$f_{nk}(t=0;\beta)=2 \text{ when } E_{nk}=5.76\text{ eV}$$

(61)

This corresponds to an initial carried density of $1.9 \times 10^{22}\text{ cm}^{-3}$, an order of magnitude larger then in the $\Gamma_{15}$ case. From Fig.3 we see that the equilibrium QP e–e and e–p lifetimes are very similar. And, indeed, the decay of the 5.76 eV state is very much affected by the e–e scattering that, in this case, at short times is dominated by the extra–carriers channel.

6. Conclusions

In conclusion I have presented an $ab$–$initio$ approach to the solution of the Baym–Kadanoff equations in realistic materials. The present approach is based on two major approximations: the generalized Baym–Kadanoff ansatz and the completed collision approximation. These makes the calculation feasible, as shown in the case of bulk silicon and provide a consistent and solid theoretical and numerical framework that can be applied to any kind of material that can be studied using the standard $ab$–$initio$ Many–Body Perturbation Theory techniques. Indeed, a major point of the present approach is that all the presented theoretical and numerical tools has been coded in an existing long–term, stable and highly productive code: the Yambo code[12]. By using bulk silicon as a paradigmatic material I show that, in semiconductors and insulators, the presence of an electronic gap makes the e–p channel dominant over the e–e one in the short–time regime ($T \lesssim 100\text{ fs}$) and for low energy carriers injected or optically pumped above the conduction band minimum but below the $2E_g$ threshold. This is due to the fact that one of the most important kinematic channels in the e–e scattering is the one that involves e–h pairs created across the energy gap, and those pairs are energetically forbidden when the carriers are pumped with energy that is $2E_g$ below the CBM. It is important to stress that on the basis of the relative balance of the e–e and e–p scattering channels presented in this work it is not possible be draw general conclusions for all kind of materials. If on one side I am convinced that, even if the CCA weakens the results obtained in the very–short time scale, the arguments based on the energy of the initial carrier to motivate a potential bigger importance of the e–p channel is reliable for any kind of semiconductor or insulator. Of course the same argument does not apply to metals. On the other hand silicon is not a system known for its strong e–p coupling. There are systems like bulk Diamond[42], carbon nano–structures[43] or super–conductors[38] where the e–p scattering is enormously strong and this can strengthen the e–p channel over the e–e one.

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