## Landauer-Büttiker formula for energy transport in hybrid electron-boson junctions

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We propose a Landauer-Büttiker formula to describe energy transport between weakly coupled hybrid electron-boson nano-junctions. The electron bath is described by effective bosonic baths with possibly nonzero chemical potential. Our theory gives a unified account of current-induced heating/cooling, electroluminescence and thermoelectric transport in nano-junctions. Our results extend the Landauer formalism to hybrid electron-boson systems and shed light on the nature of energy transfer between electron and boson systems.

**Introduction**— The celebrated Landauer-Büttiker formalism has been widely used in the study of particle and energy transport in meso- and nano-scale conductors, carried by particles following different statistics, including electrons[], photons[1], and phonons[2–8]. The energy current between two baths (i=1,2) can be written as the following general form

$$J = \int_0^{+\infty} \frac{d\omega}{2\pi} \hbar \omega T(\omega) [n(\omega, \mu_1, T_1) - n(\omega, \mu_2, T_2)]. \quad (1)$$

Here, the transmission coefficient  $T(\omega)$  describes the transmission probability of particle with energy  $\hbar\omega$  from bath 1 to 2 and vice versa. The two baths are assumed to be in thermal equilibrium, and the corresponding distribution function  $n(\omega, \mu_{\alpha}, T_{\alpha})$  of each bath is determined by the particle statistics. It is the Fermi-Dirac distribution for fermions, and Bose-Einstein distribution for bosons. A difference in the distribution drives an energy current flow between the two thermal baths. This driving force for energy transport could be a chemical potential or temperature bias for fermions. However, for bosons we have  $\mu=0$  in thermal equilibrium, following the textbook argument that bosons without number conservation have zero chemical potential. Thus, for bosonic energy transport the only driving force is temperature.

However, there are a large class of energy transport processes that can not be directly handled by Eq. 1. Hybrid energy transport between fermions and bosons is one case, and baths under nonequilibrium state is another. These kinds of processes are ubiquitous in the study of energy transport in nano-junctions. Examples include electrical-current-driven light emission, Joule heating, current-induced cooling and so on. In this work, we show that, energy transport between steadystate nonequilibrium electrons and bosons can be well described by a Landauer-Büttiker formula between bosonic baths with non-zero chemical potentials. The key observation is that, the electronic system can be treated as bosonic electron-hole pair (EHP) baths with possibly non-zero chemical potential. This is possible since the energy transport is always accompanied by the generation or annhilation of different kinds of EHPs. Our theory thus gives a unified account of different physical

processes and generalizes the Landauer-Büttiker formalism to hybrid energy transport between nonequilibrium baths.

Model— We consider a model system schematically shown in Fig. 1 (a). The *system* composed of an independent set of bosonic degrees of freedom (DOF) taken as a set of harmonic oscillators. The system couples to two kinds of baths. One is an equilibrium boson bath (phbath), modeled by an infinite number of harmonic oscillators. The other is an electronic bath (e-bath), which itself includes a central part and two electrodes (L and R). The e-bath may be driven into a nonequilibrium steady state by a voltage bias applied between electrodes L and R. Without loss of generality, we assume that the system couples only to the central region of the e-bath. Energy transport between e-bath and ph-bath takes place through their simultaneous coupling to the system.

Within the picture of Landauer-Büttiker formalism, we treat the electrons as non-interacting. But they may be driven into nonequilibrium by applying voltage or temperature bias between L and R electrodes. The electrons couples to the 'displacement' of the system harmonic oscillator

$$H_{es} = \sum_{i,j,k} M_{ij}^k c_i^{\dagger} c_j u_k. \tag{2}$$

Here,  $M_{ij}^k$  describes the coupling of the system mode k to the electronic transition between electron states i and j, and  $u_k$  is the 'displacement' operator of the system mode k. We consider the weak coupling case so that we only need to take into account the interaction up to the second order in M. The coupling of system to the phbath is linear between harmonic oscillators and can be treated exactly.

Electron-hole pair bath — Our key observation is that the interaction between the system and electron bath can be modeled by different kinds of reactions between EHPs in the e-bath and the bosonic modes in the system. The creation and annihilation of the bosonic mode is always accompanied by the recombination and creation of EHPs. These processes can be expressed in

the form of reactions

$$e_{\alpha} + h_{\beta} \rightleftharpoons p_n,$$
 (3)

where  $e_{\alpha}$ ,  $h_{\beta}$  and p represent electron in electrode  $\alpha$ , hole in electrode  $\beta$  and bosonic mode n in the system. It is equivalent to the more obvious form

$$e_{\alpha} \rightleftharpoons e_{\beta} + p_n,$$
 (4)

representing inelastic electronic transition from electrode  $\alpha$  to  $\beta$ , accompanied by emission (forward) and absorption (backward) of bosonic mode n.

There are four types of EHPs which we label by the spatial location of the electron  $(\alpha)$  and hole  $(\beta)$  state. They are schematically shown in Fig. 1 (c), and termed EHP-i, with i=1,2,3,4. They are further divided into two groups, where 1, 2 are intra-electrode type (LL,RR), and 3, 4 are inter-electrode type (RL,LR). In additional to energy transfer between e-bath and the system, the generation and recombination of inter-electrode EHPs  $(\alpha \neq \beta)$  also involves electron transport across the system. We take the energy of mode  $p_n \ \varepsilon_n = \hbar \omega_n$  and that of the EHPs to be positive.

The key quantity to describe the EHPs is the couplingweighted EHP power spectrum. It can be written as

$$\Lambda_{mn}^{\alpha\beta}(\omega) = \hbar\omega \left( n_B(\hbar\omega - \mu_{\alpha\beta}, T_e) + \frac{1}{2} \right) \Gamma_{mn}^{\alpha\beta}(\omega), \quad (5)$$

with the coupling weighted density of states (DOS)

$$\Gamma_{mn}^{\alpha\beta}(\omega) = -\sum_{i_{\alpha},f_{\beta}} \langle \psi_{i_{\alpha}}(\varepsilon_{i}) | M^{m} | \psi_{f_{\beta}}(\varepsilon_{f}) \rangle$$

$$\times \langle \psi_{f_{\beta}}(\varepsilon_{f}) | M^{n} | \psi_{i_{\alpha}}(\varepsilon_{i}) \rangle \delta(\varepsilon_{i} - \varepsilon_{f} - \hbar \omega)$$

$$\times (n_{F}(\varepsilon_{\alpha} - \mu_{\alpha}, T_{\alpha}) - n_{F}(\varepsilon_{\beta} - \mu_{\beta}, T_{\beta})) (\hbar \omega)^{-1}$$

$$= -\int \frac{d\varepsilon_{\alpha}}{2\pi} \text{tr}[M^{m} A_{\beta}(\varepsilon - \hbar \omega) M^{n} A_{\alpha}(\varepsilon_{\alpha})]$$

$$\times (n_{F}(\varepsilon_{\alpha} - \mu_{\alpha}, T_{\alpha}) - n_{F}(\varepsilon_{\beta} - \mu_{\beta}, T_{\beta})) (\hbar \omega)^{-1}$$

$$(7)$$

Here,  $n_F$  and  $n_B$  are the Fermi-Dirac and the Bose-Einstein distributions, respectively,

$$n_{F/B}(\varepsilon, T) = \frac{1}{\exp(\beta_B \varepsilon) \pm 1},$$
 (8)

and  $\beta_B = (k_B T)^{-1}$ ,  $\mu_{\alpha\beta} = \mu_{\alpha} - \mu_{\beta}$ . They are fixed by the e-bath temperature  $T_e$  and the chemical potentials of the two electrodes.

Equation 5 follows the normal form of fluctuationdissipation relation for an equilibrium boson bath, albeit with a possibly non-zero chemical potential. The intra-electrode EHPs (i=1,2) are always in equilibrium with  $\mu_{\alpha\alpha} = 0$  and temperature  $T_e$ . But the two interelectrode EHPs (i=3, 4) have opposite chemical potential  $mu_{RL} = -\mu_{LR}$ . They are nonzero when there is a voltage bias applied. This effective model is shown in Fig. 1 (b). Thus, we have shown that the *nonequilibrium* e-bath can be divided into four *equilibrium* EHP baths with different chemical potentials.

Detailed balance and effective temperature.— We now proceed to show that, a slightly modified detailed balance relation applies to each of the EHP baths. To simplify the analysis, we consider one bosonic mode with angular frequency  $\Omega$ . A simple rate equation for the mode population N can be established by considering the forward and backward reaction processes

$$\dot{N} = B(N+1) - AN,\tag{9}$$

where B and A are the reaction rates. They can be calculated by summing over the individual rates  $B = \sum_{\alpha\beta} B_{\alpha\beta}$ ,  $A = \sum_{\alpha\beta} A_{\alpha\beta}$ , with

$$B_{\alpha\beta} = \frac{2\pi}{\hbar} \sum_{i_{\alpha}, f_{\beta}} |\langle \psi_i(\varepsilon_i) | M^m | \psi_f(\varepsilon_f) \rangle|^2$$
 (10)

$$\times n_F(\varepsilon_i)(1 - n_F(\varepsilon_f))\delta(\varepsilon_i - \varepsilon_f - \hbar\Omega), \qquad (11)$$

and  $A_{\alpha\beta}$  is obtained by the replacement  $\hbar\Omega \to -\hbar\Omega$ . As a result, the ratio A/B follows

$$\frac{A_{\alpha\beta}}{B_{\alpha\beta}} = \exp(\beta_B(\hbar\Omega - \mu_{\alpha\beta})). \tag{12}$$

This is a generalized detailed balance equation for the emission and absorption of the bosonic quanta. For ebath in equilibrium ( $\mu_L = \mu_R$ ), we recover the equilibrium detailed balance relation. This means each type of the reaction drives the mode into a Bose-Einstein with a chemical potential  $\mu_{\alpha\beta}$ . This coincides with the chemical potential of the corresponding EHPs, as required by the equilibrium condition of 3:

$$\mu_{\alpha} - \mu_{\beta} = \mu_{p}. \tag{13}$$

Here, we have used the fact that chemical potential of holes is the opposite to that of electrons.

The steady state population of mode is obtained by setting  $\dot{N}=0$  as

$$N = \frac{1}{A/B - 1}.\tag{14}$$

In equilibrium  $(\mu_{\alpha} = \mu_{\beta})$ , we have  $A/B = \exp(\beta_B \hbar \Omega)$ , as required by detailed balance. Thus, N follows the standard Bose-Einstein distribution with temperature  $T_e$  and zero chemical potential. When there is voltage bias applied, the final distribution can not be written as a simple form. Normally, an effective temperature is defined by assuming N follows the Bose-Einstein distribution with zero chemical potential

$$k_B T_{eff} = \frac{\hbar\Omega}{\ln(1+N^{-1})}. (15)$$

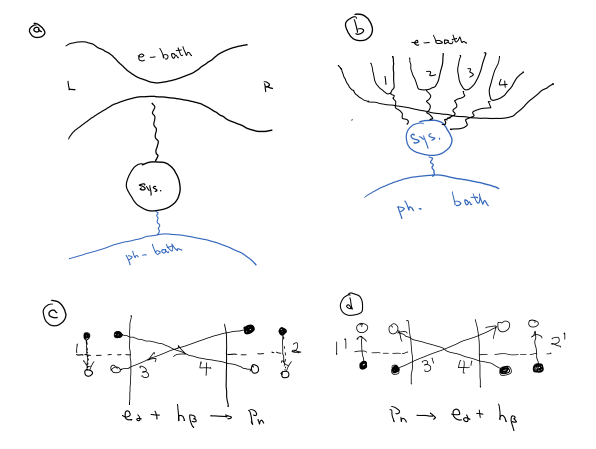


FIG. 1. (a) Schematics of the model we consider. The system consists a set of independent bosonic modes. It couples to an electron bath (e-bath), which is modeled as a conductor including a left (L) and a right (R) electrode, with temperature  $T_e$  and chemical potential  $\mu_L$  and  $\mu_R$ , respectively. The system further couples to an external thermal bath (ph-bath) at temperature  $T_{ph}$ . (b) The electron bath can be treated as four different kinds of electron-hole pair (EHP) baths (1-4), shown in (c). (c-d) Four kinds of EHP recombination (c) and excitation (d) processes. The EHPs are classified according to the spatial location of the electron  $(e_{\alpha})$  and the hole  $(h_{\beta})$ . (1,1') Both are at electrode L, (2,2') Both are at electrode R, (3,3') The electron at electrode R and a hole at electrode L. (4,4') The electron at electrode L and a hole at electrode R.

According to previous discussion, we can equivalently defined an effective chemical potential by assuming N follows the Bose-Einstein distribution at  $T_e$ 

$$\mu_{eff} = \hbar\Omega - k_B T \ln(1 + N^{-1}).$$
 (16)

These two effective parameters are related through

$$T_{eff} = \frac{T_e}{1 - \mu_{eff}/(\hbar\Omega)}.$$
 (17)

Several comments are noteworthy at this point. Firstly, in the presence of voltage bias, if the reverse of process 4 is normally enhanced more than process 3, we have a positive  $\mu_{eff}$  and consequently  $T_{eff} > T_e$ . The result is heating of the bosonic mode. In the limiting case shown in Fig. (2) (a), resonant enhancement may lead to the extreme case of  $\mu_{eff} = \hbar \Omega$ , or  $T_{eff} \to +\infty$ . This marks the instability of the bosonic mode. This case has been analyzed in details in Ref. 10. This instability means

that the harmonic approximation is not applicable any more[12]. In the other limiting case (Fig. 2(b)), process 3 is resonantly enhanced, resulting in  $\mu_{eff} > 0$  or  $T_{eff} < T_e$ . In this regime, the voltage bias is used to cool the bosonic mode below  $T_e$ .

Energy transport.—Within the effective EHP model in Fig. 1 (b), hybrid energy transport between the electrons and the system bosons can be treated as bosonic transport. To the lowest order approximation, we arrive at a Landauer-Büttiker formula for the energy transport from e-bath to the system as a summation of contributions from all the EHP baths

$$J = \sum_{\alpha,\beta} \int_{0}^{+\infty} \frac{d\omega}{2\pi} \hbar\omega \, \mathcal{T}^{\alpha\beta}(\omega)$$
$$\times \left[ n_{B}(\omega - \mu_{\alpha\beta}, T_{e}) - n_{B}(\omega, T_{ph}) \right] \tag{18}$$

where

$$\mathcal{T}^{\alpha\beta}(\omega) = \text{Tr}[\Gamma^{\alpha\beta}(\omega)\mathcal{A}_{ph}(\omega)], \tag{19}$$

is the transmission between the EHP bath and the phbath. Here,  $T_e$  and  $T_{ph}$  are the temperature of the e-bath and ph-bath, respectively. The trace Tr is over system DOF, with  $\mathcal{A}_{ph} = D^r \Gamma_{ph} D^a$  the spectral function of the system due to coupling to the ph-bath. The summation over  $\alpha\beta$  includes contributions from all the four types of EHPs. Each of them contributes to an energy transport channel. In the following we show several applications of this central result.

Heat rectification in hybrid electron-boson system— Firstly, we consider the situation where the ebath and ph-bath are in their own thermal equilibrium with two different temperature  $T_e$  and  $T_{ph}$ . This indicates that  $\mu_{\alpha} = \mu_{\beta}$  and  $T_{\alpha} = T_{\beta} = T_e$ . If we ignore the energy dependence of A within the integral window determined by the difference of the two Fermi functions, the EHP DOS becomes a constant, independent on system parameters,

$$\Gamma_{mn} = \text{tr}[M^m A M^n A], \tag{20}$$

with  $A = A_L + A_R$ . Consequently, the transmission  $\sum_{\alpha\beta} \mathcal{T}^{\alpha\beta}$  does not depend on  $T_e$ . Equation (18) reduces to the Landauer formula for heat transport between two harmonic thermal baths. From the energy transport point of view, the EHPs behave as linear harmonic oscillator thermal baths.

On the other hand, if we consider the energy dependence of  $A(\varepsilon)$ ,  $\Gamma(\omega)$ ,  $\mathcal{T}^{\alpha\beta}$  will depend on  $T_e$ . Energy transport becomes nonlinear. In this case, heat rectification is possible, i.e.,  $J(\Delta T) \neq J(-\Delta T)$ , with  $\Delta T = T_e - T_{ph}$ . We thus find a necessary condition for heat rectification in a hybrid electron-boson system: the electron DOS in the thermal window near the chemical potential has to be energy dependent[13, 14]. For normal metal electrode, the energy scale of electrons is much larger than thermal energy, leading to a flat DOS within the thermal window. The energy dependence of A can be engineered by changing the electronic states of the central part. For example, discrete energy levels of a molecular junction or quantum dot can be used. Figure 2 shows two limiting cases. We have two electronic levels which couple to the left and right electrodes, respectively. The left level lies below  $\mu_L$  and the right level lies above  $\mu_R$ . Depending on their relative positions, one of the inter-electrode EHPs couples strongest to the bosonic system. This setup has been studied in Ref. 10.

Thermoelectric transport: current from coldspot.— Another problem we can look at is the thermoelectric transport. When there is a temperature difference between  $T_{ph}$  and  $T_e$ , in addition to the energy transport between system and e-bath, an electrical current may also be induced between left and right electrodes[15, 16]. In our EHP picture, this is realized through coupling of the bosonic mode with two interelectrode EHPs. Since they contribute to two electri-

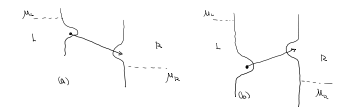


FIG. 2. Two limiting cases of electronic heating (a) and cooling (b). For the e-bath, we have a filled electronic level that couples to the left electrode with chemical potential  $\mu_L$ , and an empty level that couples to the right electrode. Electron transport between the two states is mediated by the bosonic mode in the system. (a) Heating of bosonic mode due to resonant recombination of EHPs. (b) Cooling of bosonic mode due to resonant generation of EHPs.

cal current with opposite directions, in order to get a non-zero electrical current, their coupling to the bosonic mode should be different[16]. Here, we consider the case where  $T_{ph} < T_e$  and  $\mu_L = \mu_R$ . The temperature difference between e-bath and ph-bath generates a heat current flow from the e-bath to the ph-bath. At a result of the heat transport, electron transport between L and R electrode takes place. Previously, electrical current generated from a phonon hot-spot  $(T_{ph} > T_e)$  has been considered[15]. The situation we consider here is somewhat counter-intuitive, since the electrical current is generated by a cold-spot. This demonstrates the decoupling of heat and charge transport as an advantage of thermoelectricity in hybrid nano-junctions.

Electronic cooling of bosonic mode.— The presence of voltage bias between the two electrodes changes the initial and final electron states of the EHP excitation. Thus, the EHP DOS can be tuned by voltage bias. More importantly, the inter-electrode EHPs acquire a non-zero chemical potential, determined by the electrochemical potential difference between the two electrodes. We assume  $\mu_L > \mu_R$  without loss of generality. The EHP-4 has a chemical potential of  $eV = \mu_L - \mu_R > 0$ , while EHP-3 gets a chemical potential with opposite value -eV. Change of the chemical potential breaks the equilibrium in the reaction, and drives the energy transport between e-bath and the system. If we assume  $T_e = T_{ph}$ , the energy transport is through the two inter-electrode EHP channels

$$J = \int_0^{+\infty} \frac{d\omega}{2\pi} \hbar\omega \left[ \mathcal{T}^{LR}(\omega) [n_B(\omega - eV, T_e) - n_B(\omega, T_{ph})] - \mathcal{T}^{RL}(\omega) [n_B(\omega + eV, T_e) - n_B(\omega, T_{ph})] \right]$$

Direction of energy flow depends on the relative magnitude of two fluxes. It can be engineered by tuning the electronic band structure, or more specifically, the transition probability. Figure 2 shows two limiting cases. In Fig. 2 (a), process 4 is enhanced due to resonant inelastic electron tunneling when the separation between the two

DOS peaks is resonant with energy of the bosonic mode. Consequently, energy flows from e-bath to the system. In Fig. 2 (b), the reverse of process 3 is resonantly enhanced, resulting in energy flow in the opposite direction. Electronic cooling becomes possible using this resonantly enhanced process.

## Conclusions-

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