



Gate-modulated thermopower of disordered nanowires: II. Variable-range hopping regime

Riccardo Bosisio, Cosimo Gorini, Geneviève Fleury, Jean-Louis Pichard

► To cite this version:

Riccardo Bosisio, Cosimo Gorini, Geneviève Fleury, Jean-Louis Pichard. Gate-modulated thermopower of disordered nanowires: II. Variable-range hopping regime . New Journal of Physics, Institute of Physics: Open Access Journals, 2014, 16, pp.095005.
 <10.1088/1367-2630/16/9/095005>.
 <cea-01138312>

HAL Id: cea-01138312 https://hal-cea.archives-ouvertes.fr/cea-01138312

Submitted on 1 Apr 2015

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Gate-modulated thermopower of disordered nanowires: II. Variable-Range Hopping Regime

Riccardo Bosisio, Cosimo Gorini, Geneviève Fleury and Jean-Louis Pichard‡

Service de Physique de l'État Condensé (CNRS URA 2464), IRAMIS/SPEC, CEA Saclay, 91191 Gif-sur-Yvette, France

Abstract. We study the thermopower of a disordered nanowire in the field effect transistor configuration. After a first paper devoted to the elastic coherent regime (Bosisio R., Fleury G. and Pichard J.-L. 2014 New J. Phys. 16 035004), we consider here the inelastic activated regime taking place at higher temperatures. In the case where charge transport is thermally assisted by phonons (Mott Variable Range Hopping regime), we use the Miller-Abrahams random resistor network model as recently adapted by Jiang et al. for thermoelectric transport. This approach previously used to study the bulk of the nanowire impurity band is extended for studying its edges. In this limit, we show that the typical thermopower is largely enhanced, attaining values larger that $10 k_B/e \sim 1 \,\mathrm{mV}\,\mathrm{K}^{-1}$ and exhibiting a non-trivial behaviour as a function of the temperature. A percolation theory by Zvyagin extended to disordered nanowires allows us to account for the main observed edge behaviours of the thermopower.

PACS numbers: 72.20.Ee 72.20.Pa 84.60.Rb 73.23.-b 73.63.Nm

‡ Corresponding author: jean-louis.pichard@cea.fr

1. Introduction

The conversion of temperature to voltage differences or its inverse, enabling respectively waste heat recovery or cooling, is the purpose of a thermoelectric device. In linear response the device efficiency is controlled by its dimensionless figure of merit $ZT = S^2 GT/\Xi$, with T the temperature, S the Seebeck coefficient or thermopower and G, Ξ respectively the electrical and thermal conductances. As ZT increases, the efficiency moves closer to the Carnot limit. The stronger the particle-hole asymmetry in a system is, the higher S will be. An ideal thermoelectric device should then exploit to the maximum such asymmetry, while at the same time ensuring a poor thermal and a good electrical conductance [1]. Whereas the former requirement is necessary to increase efficiency, the latter is needed for enough electric (cooling) power to be extracted from a heat engine (Peltier refrigerator). From this perspective, semiconductor nanowires appear as very promising central building blocks of flexible, efficient and environmentally friendly thermoelectric converters [2, 3, 4, 5, 6, 7, 8]. Whereas their thermoelectric properties can be easily tuned by gates [5, 8], the phononic contribution to thermal transport Ξ_{ph} is suppressed due to the reduced dimensionality [3, 4], and a good power output could be achieved by stacking them in parallel [9, 3, 6]. Furthermore Si-based devices, already under intense investigation [10, 11, 12, 9, 13, 14, 3, 15, 4, 6], exploit an abundant and non-polluting resource.

Most existing works concentrate either on highly doped samples [10, 12, 9, 3, 7, 6] or on the thermal conductivity of undoped wires [11, 14, 16, 15, 4]. On the other hand recent studies by Jiang *et al.* [17, 18] have rekindled the interest for systems in which electronic transport takes place via phonon-assisted hopping between localised states, of which disordered nanowires with low carrier density are a paradigmatic realization. Whereas in our first paper [19] we focused on the low-temperature coherent regime, in this work we extend the approach reviewed in Refs. [17, 18] in order to investigate band-edge transport in the (activated) hopping regime. Two simple physical mechanisms have in this regime a synergy which is ideal for thermoelectric conversion [20, 21]: (i) a strongly broken particle-hole symmetry due to the Fermi level lying close to the band edge; (ii) a wide energy window around the Fermi level made available for transport by the phonons. In other words, the phonons lend the carriers the energy necessary for them to hop through the system, but of the latter only one species, either electrons or holes, has available states and thus actually propagates.

The general setup we have in mind is sketched in Fig. 1: A disordered semiconductor nanowire (green) connected to two metallic contacts (yellow) and deposited on an insulating substrate (blue). A heater (grey) and an applied bias voltage can induce a temperature and an electrochemical potential difference between the two contacts. A back gate (dark grev), placed below the substrate. allows to shift the impurity band of the nanowire by means of a gate voltage. This way, the transport of charges and heat can be studied when the Fermi potential of the setup probes either the bulk of the band or its edges. Fig. 1 depicts the more commonly used field effect transistor (FET) configuration [5]. Another possibility would be to cover only the nanowire with a top gate (see e.g. Ref. [22]). Putting a back gate is easier, but large gate voltages (a few hundreds volts) are necessary for shifting the impurity band, while few volts are sufficient if one uses a top gate. The nanowire itself could be (i) lightly doped, with electrons localised around distant impurity states, or (ii) highly doped but strongly depleted, or (iii) made of an amorphous semiconductor. A crucial feature of such wires is that their length L should be much longer than the localisation length ξ of their electron states, such that their electrical resistance becomes exponentially large when the temperature is lowered below a few Kelvin degrees. A crude modelling of such setup is sketched in Fig. 2: A purely 1D disordered chain with $L \gg \xi$, connected to two electron reservoirs and to a phonon bath (represented by the substrate), and coupled to a gate used to modulate its carrier density. Each site of the chain corresponds to an electronic state localised by disorder or

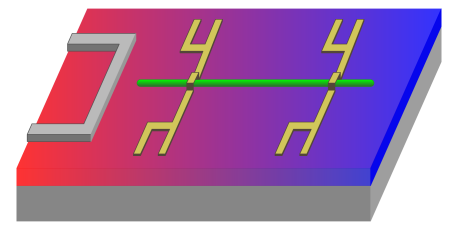


Figure 1. Nanowire in the field effect transistor (FET) device configuration: The nanowire (green) with two metal contacts (yellow) is deposited on an insulating substrate (blue). A heater (grey) makes the left side of the setup hotter (red) than its right side. A back gate (dark grey) is put below the substrate.

bound to impurity sites. Though such a model is strictly 1D, it should allow us to describe also quasi-1D wires [23, 24, 25] as long as their transverse sizes remain negligible compared to the typical hopping length (which we will define later).

In this work we are primarily interested in the thermopower S of a single nanowire, as studied in many recent experiments [5, 6, 7, 8]. We start in Sec. 2 by introducing model and methods employed, moving on to discuss the nanowire electrical conductance in Sec. 3 and its thermopower in Sec. 4, before concluding in Sec. 5. Various technical details, skimmed over in the main body for ease of reading, are gathered in the appendices.

2. Model and method

As sketched in Fig. 2, we consider a disordered nanowire of length L in which all available electronic states are exponentially localised at positions x_i , with a localisation length $\xi_i \ll L$. We assume each state i is either empty, or occupied by a single electron, but cannot be doubly occupied owing to a strong on-site Coulomb repulsion[26]. The energy levels E_i of the localised states are distributed within a band of width $2E_B$ and $\nu(E)$ denotes their density of states (DOS) per unit length at energy E. They can be shifted as a whole by an external gate voltage V_g . The nanowire is attached at its ends to two metallic contacts held at electrochemical potentials μ_L and μ_R and temperatures T_L and T_R . It is also coupled to a phonon bath at temperature T_{ph} which provides the energy for electrons to hop between localised states. We focus on the situation in which the temperature T is the same in all reservoirs ($T_L = T_R = T_{ph} \equiv T$) and consider linear response, assuming the difference in electrochemical potentials between left and right leads to be small ($\mu_L = \mu + \delta \mu \gtrsim \mu_R \equiv \mu$).

2.1. Identification of the different transport mechanisms and of their temperature scales

Transport through the nanowire happens as follows. Since there is a continuum of available states in the leads, we assume that charge carriers, let us say electrons, enter or leave the nanowire by

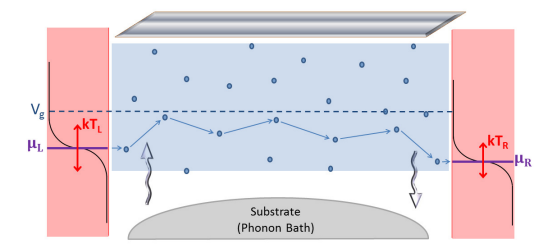


Figure 2. Variable Range Hopping (VRH) transport for a disordered nanowire in a FET configuration: Two ohmic contacts are connected by a 1D disordered chain of length L where the electron states are localised. The contacts are thermalised at temperatures $T_L(T_R)$ with electrochemical potentials $\mu_L(\mu_R)$ respectively. The electronic states (blue dots of coordinates (x_i, E_i)) are localised in regions of size $\xi_i \ll L$. Their centres x_i are taken at random along the chain, with energies E_i distributed inside an impurity band of width $2E_B$ (shaded light blue region). A top gate (in grey at the top of the figure) allows to shift the impurity band. The gate potential V_g sets the center of the band (dashed line). In linear response, the carriers are injected from the left (right) contacts inside the electronic states localised near the edges of the chain, in a window of energies of order $k_B T_L(k_B T_R)$ around $\mu_L(\mu_R)$. Inside the chain, the carrier propagation is thermally assisted by phonons (wavy arrows), which allow a carrier to do hops of variable range between localised states at different energies. The phonon bath at temperature T_{ph} is represented by the substrate upon which the chain is deposited.

elastic tunneling processes, without absorbing or emitting phonons§. Inside the nanowire they have the possibility to hop either to localised states at higher energies by absorbing phonons, or to localised states at lower energies by emitting them. Determining precisely the favoured electronic paths is a complicated task. The proper way to tackle this issue is to map the hopping model to an equivalent random resistor network [27] and then to reduce it to a percolation problem [26]. Such microscopic approaches are needed for giving precise quantitative predictions, but Mott's original argument [28, 29] gives the main ideas: Assuming the localisation lengths and the density of states to be constant within a certain window of energies Δ to be explored ($\xi_i \approx \xi$, $\nu(E) \approx \nu$), the electron transfer from one localised state to another separated by a distance x and an energy $\delta E \propto 1/(\nu x^D)$ (D = 1 for us) results from a competition between the elastic tunneling probability $\propto \exp{-(2x/\xi)}$ to do a hop of length x in space and the Boltzmann probability ($\propto \exp{-(\delta E/k_BT)}$) to do a hop of δE in energy. Short hops are favoured by the former but are too energy-greedy for the latter, since localised states close in space are far in energy. This competition gives rise to an optimal electron hopping length, the Mott hopping length, which reads

$$L_M = \sqrt{\frac{\xi}{2\nu k_B T}} \tag{1}$$

§ Phonon absorption and emission in the electrodes could be straightforwardly taken into account. However, it should not add any new physics and we neglect it.

in one dimension. L_M is a decreasing function of the temperature, which allows us to define two characteristic temperature scales: the *activation temperature*

$$k_B T_x = \frac{\xi}{2\nu L^2} \tag{2}$$

at which $L_M \simeq L$ and the *Mott temperature*

$$k_B T_M = \frac{2}{\nu\xi},\tag{3}$$

at which $L_M \simeq \xi$. At low temperatures $T < T_x$, L_M exceeds the system size and transport through the nanowire occurs via elastic coherent tunneling (see Ref. [19]). Above T_x , transport becomes inelastic, and remains coherent at scales smaller than L_M only. The regime of intermediate temperature $T_x < T < T_M$ is known as the variable-range hopping (VRH) regime. As sketched in Fig. 2, electronic transport in this regime is achieved via several jumps of length $\approx L_M$ (with $\xi < L_M < L$). As it can be proven using a microscopic approach based on random resistor networks and percolation theory [26, 20, 21], the VRH conductance can be simply expressed in terms either of L_M , T_M or the hopping energy Δ ,

$$G \propto \exp\left\{-\frac{2L_M}{\xi}\right\} = \exp\left\{-\sqrt{\frac{T_M}{T}}\right\} = \exp\left\{-\frac{\Delta}{k_B T}\right\},\tag{4}$$

where (it will be of prime importance later on)

$$\Delta = k_B \sqrt{T_M T} \tag{5}$$

defines the width of the energy interval around μ inside which are located all states contributing to transport. Let us underline that if $T_x < T \ll T_M$, Δ becomes much larger than $k_B T$, the relevant energy interval for transport in the coherent regime $(T < T_x)$. At large temperatures $T > T_M$, L_M becomes of the order of or even smaller than the localisation length ξ , and one enters the nearest-neighbour hopping (NNH) regime where transport is simply activated between nearest neighbour localised states. Actually, in 1D, the crossover from VRH to simply activated transport is expected to take place at temperatures lower than T_M . The reason is the presence of highly resistive regions in energy-position space, where electrons cannot find empty states at distances $\sim \Delta, L_M$. These regions can be circumvented in 2D or 3D but not in 1D, where they behave as "breaks" in the percolating path: electrons are topologically constraint to cross them by thermal activation, making the temperature dependence of the overall resistance simply activated [30, 31]. The critical temperature T_a that marks the onset of this simply activated behaviour is given implicitly by the relation [32]

$$L = \frac{\xi}{2} \sqrt{\frac{T_M}{2T_a}} \exp\left\{\frac{T_M}{2T_a}\right\}.$$
 (6)

Below T_a , the probability of having such breaks in the nanowire can be neglected.

In Fig. 3 the temperatures T_x , T_M and T_a are given as a function of the gate voltage V_g , taking for the disordered nanowire an Anderson model where the L random site potentials are shifted by $V_q \parallel$, the electrochemical potential μ being fixed in the reservoirs.

5

^{||} We assume the gate acting only along the nanowire, which corresponds to using a top gate. A FET configuration with a back gate should behave similarly, the field effect in the metallic contacts being negligible.

6

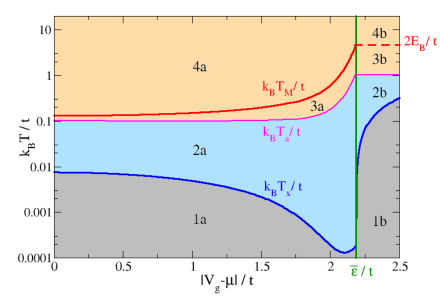


Figure 3. Gate dependence of the temperature scales separating the different regimes of electronic transport in a disordered nanowire: Elastic regime (grey), inelastic VRH regime (blue) and simply activated regime (red)). By varying the gate voltage V_g , one scans the impurity band, starting from its center (when $V_g - \mu = 0$) towards its edges (approximately for $|V_g - \mu| = \bar{\epsilon}$) and ending up eventually outside the band (when $|V_g - \mu| \gtrsim \bar{\epsilon}$). The scales T_x , T_a and T_M defined in Sec. 2.1 have been plotted for the Anderson model introduced in Sec. 2.3, with W = t and L = 200.

Still following Mott's approach, we consider $\xi_i \approx \xi$ and $\nu(E) \approx \nu$ (both evaluated at μ), thus neglecting their variations within Δ . The shape of the curves is a consequence of the explicite energy dependence of the localisation length ξ and of the DOS ν , which is detailed in Sec. 2.3. Approaching an impurity band edge $(\pm E_B)$, both ξ and ν decrease rapidly, inducing a large increase of T_M and T_a that must be eventually cut-off when T_M exceeds the bandwidth $2E_B$. Indeed, when $T \to T_M = 2E_B, \ \Delta \to 2E_B$ and the range of states available for hopping transport reaches its limit. More explicitly, we estimate this to happen at an energy scale $|\mu - V_g| \approx \bar{\epsilon} \approx 2.2t$ for the set of parameters considered in Fig. 3.

When Eqs. (2), (3), and (6) cease to be valid, we will use a simplified model introduced by Zvyagin for estimating the temperatures T_x , T_M and T_a . In this model, the DOS drops abruptly from a constant to 0 at $\bar{\epsilon}$. This yields that, when $|\mu - V_g| \gtrsim \bar{\epsilon}$, T_M and T_a do not vary anymore and keep their values at $\bar{\epsilon}$, while the activation temperature T_x gives the energy that electrons need in order to jump inside the band: $k_B T_x \approx |\mu - V_g| - \bar{\epsilon}$. We will show later that the edge behaviours numerically obtained using the Anderson model are well described by this simplified model.

As a summary, let us now discuss the regimes of electronic transport corresponding to each region of the temperature diagram established in Fig. 3. Standard VRH regime takes place in region (2a), at intermediate temperatures, when μ lies inside the impurity band. According to Mott law in 1D, the average logarithm of the resistance behaves there as $(T_M/T)^{1/2}$. In Sec. 3, we will see how this statement has to be revisited in the vicinity of the band edges, and how to take into account the energy dependency of ξ . At higher temperatures, transport is simply activated (the temperature dependence of the logarithm of the resistance $\propto T^{-1}$). This is due either to the presence of a very resistive link in the best conducting path that dominates the resistance (region (3a)), or simply to the fact that the thermal energy k_BT is so high that transport occurs via hops between nearest neighbour states, no matter how far in energy they are (region (4a)). On the contrary at lower temperatures, in region (1a), $L \leq L_M$ and transport ceases to be thermally activated to become elastic and coherent through the whole nanowire. The thermopower in this regime has been studied in Ref. [19]. If now μ lies outside the impurity band, electrons need to absorb energy in order to enter the band. In region (1b), k_BT is too small for that (the only way for electrons to cross the nanowire is then to tunnel directly from one reservoir to the other, which results in a exponentially vanishing conductance). At higher temperatures, in regions (2b), (3b) and (4b), electrons can be thermally activated. Once they have entered the nanowire, they hop from site to site according to the mechanism prevailing in regions (2a), (3a) and (4a) respectively.

2.2. Formulation in terms of a random resistor network

We follow the approach used in Refs. [17, 18] for studying thermoelectric transport in the hopping regime. It consists in solving the Miller-Abrahams resistor network [27] which was first introduced for describing charge transport in weakly doped crystalline semiconductors and later on extended to non crystalline Anderson insulators. The nodes are given by the localised states. Each pair of nodes i, j is connected by an effective resistor, which depends on the transition rates Γ_{ij}, Γ_{ji} induced by local electron-phonon interactions. In addition, one needs to connect this network to the leads, if one wants to calculate the charge and heat currents flowing through it. Usually (and actually, we did not find a reference where this is not the case) one assumes for calculating these transition rates that $\xi_i = \xi(E_i) \equiv \xi$ (evaluated at μ) for the localisation lengths of the different states, which can be done if the variations of the ξ_i are negligible within Δ . Here we need to go beyond such an approximation, since we are interested in band edge transport, where those variations cannot be neglected. The procedure is summarized below.

Let us consider a pair of localized states i and j of energies E_i and E_j . Assuming no correlations between their occupation numbers, the (time-averaged) transition rate from state i to state j is given by the Fermi golden rule as [18]

$$\Gamma_{ij} = \gamma_{ij} f_i \left(1 - f_j \right) \left[N_{ij} + \theta (E_i - E_j) \right], \tag{7}$$

where f_i is the average occupation number of state i and $N_{ij} = [\exp\{|E_j - E_i|/k_BT\} - 1]^{-1}$ is the phonon Bose distribution at energy $|E_j - E_i|$. The presence of the Heaviside function accounts for the difference between phonon absorption and emission [26]. γ_{ij} is the hopping probability $i \to j$ due to the absorption/emission of one phonon when i is occupied and j is empty. Assuming that the energy dependence of ξ can be neglected, in the limit $x_{ij} \gg \xi$ one obtains

$$\gamma_{ij} \simeq \gamma_{ep} \exp(-2x_{ij}/\xi) \,. \tag{8}$$

Here $x_{ij} = |x_i - x_j|$ is the distance between the states, whereas γ_{ep} , containing the electron-phonon matrix element, depends on the electron-phonon coupling strength and the phonon density of states. Since it is weakly dependent on E_i , E_j and x_{ij} compared to the exponential factors, it is assumed to be constant. Under the widely used approximation [26, 33, 34, 20] $|E_{ij}| \gg k_B T$, Eq. (7) reduces to:

$$\Gamma_{ij} \simeq \gamma_{ep} \, e^{-2x_{ij}/\xi} \, e^{-(|E_i - \mu| + |E_j - \mu| + |E_i - E_j|)/2k_B T} \,. \tag{9}$$

Hereafter, we will go beyond these standard approximations by considering the exact expression (7) for Γ_{ij} , and by taking

$$\gamma_{ij} = \gamma_{ep} \left(\frac{1}{\xi_i} - \frac{1}{\xi_j}\right)^{-2} \left(\frac{\exp\{-2r_{ij}/\xi_j\}}{\xi_i^2} + \frac{\exp\{-2r_{ij}/\xi_i\}}{\xi_j^2} - \frac{2\exp\{-r_{ij}(1/\xi_i + 1/\xi_j)\}}{\xi_i\xi_j}\right), \quad (10)$$

for γ_{ij} . Eq. (10) takes into account the energy dependence of $\xi(E)$ and is derived in Appendix A.

The tunneling transition rates between each state i and the leads α ($\alpha = L$ or R) are written in a similar way as

$$\Gamma_{i\alpha} = \gamma_{i\alpha} f_i \left[1 - f_\alpha(E_i) \right] \tag{11}$$

where

$$\gamma_{i\alpha} \simeq \gamma_e \exp(-2x_{i\alpha}/\xi_i).$$
 (12)

In the above equations $f_{\alpha}(E) = [\exp\{(E - \mu_{\alpha})/k_BT\} + 1]^{-1}$ is lead α 's Fermi-Dirac distribution, $x_{i\alpha}$ denotes the distance of state *i* from lead α and γ_e is a rate quantifying the coupling between the localized states and the leads (taken constant for the same reason as γ_{ep}).

Then, the net electric currents flowing between each pair of localized states and between states and leads are obtained by

$$I_{ij} = e\left(\Gamma_{ij} - \Gamma_{ji}\right) \tag{13a}$$

$$I_{i\alpha} = e \left(\Gamma_{i\alpha} - \Gamma_{\alpha i} \right) \qquad \alpha = L, R \tag{13b}$$

e < 0 being the electron charge. The linear response solution of this random resistor network problem is reviewed in Ref. [18]. Details of the calculation of the charge currents and heat currents are summarized in Appendix A for the Peltier configuration we consider, where the temperature is T everywhere and the reference (equilibrium) electrochemical potential is that of the right reservoir $(\mu \equiv \mu_R)$. In this case the electrical conductance G, Peltier coefficient II and thermopower S are determined within the Onsager formalism by the charge (I_L^e) and heat (I_L^Q) currents exchanged with the left reservoir:

$$G = \frac{I_L^e}{\delta \mu/e},\tag{14a}$$

$$\Pi = \frac{I_L^Q}{I_L^e},\tag{14b}$$

$$S = \frac{\Pi}{T} = \frac{1}{T} \frac{I_L^Q}{I_L^e}.$$
 (14c)

In the last equation, the Kelvin-Onsager symmetry relation [35] $\Pi = ST$ has been used for deducing the thermopower. Notice that a different choice of reference could be adopted (see for instance Ref. [18]) without affecting the transport coefficients G, Π and S.

2.3. Anderson model for the localised states

The set of energies E_i and localisation lengths ξ_i are required as input parameters of the random resistor network problem. To generate them we use the Anderson model. The disordered nanowire is modeled as a 1D lattice of length L with a lattice spacing a set equal to one, described by a $L \times L$ tight-binding Hamiltonian:

$$\mathcal{H} = -t \sum_{i=1}^{L-1} \left(c_i^{\dagger} c_{i+1} + \text{h.c.} \right) + \sum_{i=1}^{L} (\epsilon_i + V_g) c_i^{\dagger} c_i , \qquad (15)$$

8

9

where c_i^{T} and c_i are the electron creation and annihilation operators on site *i* and *t* is the hopping energy. In the following all energies will be expressed in units of *t*. The disorder potentials ϵ_i are (uncorrelated) random numbers uniformly distributed in the interval [-W/2, W/2]. The constant potential V_g is added to take into account the presence of an external top gate, allowing to shift the whole nanowire impurity band.

By diagonalizing the Hamiltonian (15), we find the energies E_i of the localised states. They are distributed with the DOS $\nu(E)$ in the interval $[V_g - E_B, V_g + E_B], \pm E_B$ being the band edges of the model at $V_g = 0$. In the limit $L \to \infty$, $E_B = 2t + W/2$. To generate the localisation lengths ξ_i , we neglect sample-to-sample fluctuations and assume that ξ_i is given by the typical localisation length $\xi(E_i)$ at energy E_i , characterizing the exponential decay of the average logarithm of the elastic conductance $(\ln G \sim -2L/\xi)$. The DOS $\nu(E)$ and localisation length $\xi(E)$ are shown in Fig. 4; their energy dependence is analytically known in the large size and small disorder limits, both in the bulk of the band and close to the edges (see Refs. [36, 19]). Obviously, if μ lies close to the band edges and/or if the available energy window Δ around μ is not small compared to t, the energy dependency of $\nu(E)$ and $\xi(E)$ cannot be neglected. This explains why we need to go beyond the approximation of constant DOS and localisation length when scanning the impurity band with the gate voltage.

Solving the Anderson model gives us the full set of localised states: their energy levels E_i , their localisation lengths $\xi_i = \xi(E_i)$ and their positions along the disordered chain. However, to speed up the procedure of building a basis of localised states, we simply assign the levels E_i to random positions x_i between 0 and L along the chain (with a uniform distribution). This approximation is conventional in numerical simulations of VRH transport (see [37, 32, 18] among others) ¶.

Hereafter, we will study disordered chains with a disorder strength W = t, which is sufficiently small for using weak disorder expansions [19] and sufficiently large for ensuring $L \gg \xi_i$ at relatively small sizes. For $V_g = 0$ and $L \approx 1000$, the spectrum edges of the disordered nanowire are found at $E_B \approx 2.35 t$, which is smaller than 2.5 t, the value characterizing the limit $L \to \infty$. Such finite size effects are a consequence of the infinitely small tails of the asymptotic DOS $\nu(E)$ shown in Fig. 4: States of energy close to 2.5 t can only exist in infinitely long chains.

3. Electrical Conductance

3.1. Background

The electrical conductance of one-dimensional conductors in the VRH regime has been much studied in the literature, both experimentally [38, 39, 40, 41, 42] and theoretically [30, 37, 32, 31, 43, 44]. In particular, the validity of Mott law for the typical conductance in 1D

$$\ln G(T) \sim -\alpha \sqrt{\frac{T_M}{T}},\tag{16}$$

with $\alpha \approx 1$, was a subject of controversy for a long time since strictly speaking, Mott's argument leading to Eq. (16) does not hold in 1D. It was shown that due to the presence of "breaks", the prefactor α is actually also a function of the temperature and system length [32, 31]. Nevertheless, the *T*- and *L*-dependency of α turns out to be so weak that at low temperatures α is almost

[¶] By doing this we lose a feature of Anderson model, namely that states which are close in energy are distant in space, and as a consequence our model may overestimate the hopping between certain pairs of states. However this should not play an important role if L is sufficiently large, $L \gg \xi(\mu)$. In this case states which are accidentally taken close both in space and in energy should be not only rare but, more importantly, can merely be seen – regarding percolation – as one small localised cluster, i.e. as a single new effective localised state. The reason is that the optimal percolation path is eventually determined by the most resistive links. Thus, we can always reformulate the problem in order to end up in a situation in which neighbouring states are far away in energy.

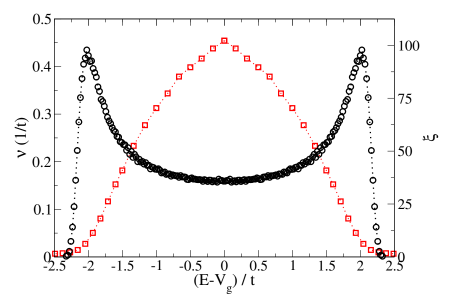


Figure 4. Density of states per site ν (O) and localisation length ξ (\Box), as a function of energy E for the 1D Anderson model (15) with disorder amplitude W/t = 1. The points correspond to numerical data (obtained in the large length limit with L = 1600). Analytical expressions describing $\nu((E - V_g)/t)$ and $\xi((E - V_g)/t)$ are given in Ref. [19].

constant and Mott law is recovered. Taking the proper $\alpha(L,T)$ into account allows an analytical description of the crossover from Mott law to the activated behaviour, $\ln G(T) \sim T^{-1}$, above T_a (see Sec. 2.1) but the refinement thus introduced is too small to be clearly evidenced by numerical simulations and even less by experimental measurements.

Another limitation of Mott's standard argument and of subsequent, more elaborate percolation-based ones is the initial assumption of a constant DOS and a constant localisation length around μ . As long as $\nu(E)$ is slowly varying in the energy window $|E - \mu| < \Delta$ (still keeping ξ constant), Eq. (16) is expected to hold, but it lacks justification in the case of strongly varying DOS. In particular, Eq. (16) has to be revised when transport through the system occurs at energies around the impurity band edges. This question was tackled by Zvyagin in Refs. [20, 21], by approximating the DOS by a step-like function. If one considers the lower band edge, the approximated DOS reads

$$\nu(E) \simeq \nu_0 \,\theta(E - \epsilon_c),\tag{17}$$

where ϵ_c plays the role of an effective band edge. Though three-dimensional systems were considered in Refs. [20, 21], a similar approach can be extended to our 1D model setting $\epsilon_c = V_g - \bar{\epsilon}$, where $\bar{\epsilon}$ is the 1D effective edge introduced in Sec. 2.1 for $V_g = 0$. The idea is that when μ lies outside the impurity band, electrons need an activation energy $\epsilon_c - \mu$ in order to "jump" inside it to find available states. This entails an extra term in Eq. (16), which in 1D becomes

$$\ln G(T) \sim -\frac{E_A}{k_B T} - \bar{\alpha} \sqrt{\frac{T_M}{T}},\tag{18}$$

with $E_A \sim \epsilon_c - \mu$ and $\bar{\alpha}$ differing from α by some numerical factors [45, 21].

3.2. Numerical results

We have investigated numerically how the typical conductance of a disordered nanowire depends on the temperature when the applied gate voltage is varied. For the model described in Sec. 2.3, we have solved the random resistor network problem and calculated the conductance G via Eq.(14a). This procedure has been iterated over many random configurations of the energy levels E_i in order to extrapolate the *typical* logarithm of the conductance $[\ln G]_0$, defined as the median of the resulting distribution $P(\ln G)^+$.

In Fig. 5, $[\ln G]_0(T)$ is plotted for two values of V_q , corresponding to the bulk and the lower edge of the band. In both cases we show that low temperature data exhibit Mott law $T^{-1/2}$ behaviour (red dashed curve), while at higher temperatures they are well fitted by an activated T^{-1} law (green dashed curve). Eq. (18) with adjusted values for E_A and $\bar{\alpha}$ describes the crossover between the two regimes (full blue line). More precisely, when μ lies inside the band (Fig. 5(a)), the validity range of Mott law $(k_B T/t \lesssim 0.05)$ is consistent with the required hypothesis of weakly varying DOS. Indeed, below such temperatures, the energy window $\Delta = k_B \sqrt{T_M T}$ of accessible states around μ is so small ($\Delta \lesssim 0.2$ using for T_M the value given in Fig. 3) that the DOS can be considered as weakly energy dependent ($\Delta \partial_E \ln \nu(E) |_{\mu} \approx 0.3 < 1$). This justifies the validity of Eq. (16) in such a regime. Note that the onset of activated behaviour at $k_B T \approx 0.05 t$ is also in rough agreement with the predicted value of $k_B T_a \approx 0.1 t$ in Fig. 3. On the other hand, when μ lies in a region where the DOS is exponentially small (Fig. 5(b)), there is no more reason to use Mott law to describe our data, even if it appears to be well fitted by Eq. (16) at low temperatures. The point is that other power law formula, $[\ln G]_0 \sim T^{\beta}$, could be used to fit our data in this narrow temperature range. Thus, one cannot use the apparent suitability of Eq. (16) to support the validity of Mott law in this regime. Outside the band the correct framework for analysis is provided by Eq. (18). The activated contribution to the conductance is always present, which explains why in Fig. 5(b) the T^{-1} fit starts to be accurate much below the temperature $k_B T_a/t \approx 0.95$. Finally, at very high temperatures (typically larger than t), the typical conductance is found in both cases to decrease with temperature. This is due to the fact that in the limit $T \to \infty$, the factors $f_i(1-f_j)$ and $f_j(1-f_i)$ on one hand, and $f_i(1-f_\alpha)$, $f_\alpha(1-f_i)$ on the other, converge to the same value. Hence, the opposite rates Γ_{ij} , Γ_{ji} and Γ_{iL} , Γ_{Li} tend to level out, which results in a vanishing net current and a divergent resistance. An expansion of the Fermi functions to the next order in inverse temperature yields I_{ii} , $I_{i\alpha} \sim T^{-1}$, which explains the linear decay at high T of $[\ln G]_0$ versus $\ln T$ in Fig. 5 (not marked).

4. Thermopower

4.1. Background

The thermopower is a measure of the average energy $\langle E - \mu \rangle$ transferred by charge carriers from the left lead to the right one. In the low temperature coherent regime [19], transport takes place near the Fermi energy. Hence, in linear response with respect to the bias voltage between the two leads, the thermopower depends on the electron-hole asymmetry at μ . On the contrary, in the VRH regime, all states in the energy window $|E - \mu| < \Delta$ contribute. Since $\Delta \gg k_B T$ when $T \ll T_M$, the thermopower benefits from the contribution of states far below and above μ , despite being in linear response. When the gate voltage is adjusted in order to probe the impurity band edges, the electron contribution dominates over the hole one (or vice-versa), yielding an enhanced thermopower.

 $^+$ More details concerning the distributions of the logarithm of the conductance for 1D systems in VRH regime can be found in Refs. [32, 46]

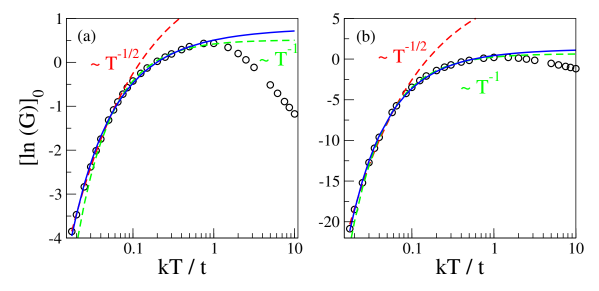


Figure 5. Typical value of the logarithm of the conductance $[\ln G]_0$ as a function of T for $\mu = 0$ and two values of the gate voltage: (a) $V_g = 1.9t$ inside the band and (b) $V_g = 2.3t$ at the edge of the band. In both cases, at low temperatures, numerical data (points) are well fitted by a $T^{-1/2}$ fit (red dashed lines), evolving to a T^{-1} behaviour as T increases (green dashed lines). Full blue lines correspond to Eq.(18), which describes the crossover between the two regimes. Parameters: L = 200, W = t and $\gamma_e = \gamma_{ep} = t$.

To study the thermopower in the VRH regime *, we use the approach introduced by Zvyagin in [20, 21]. The starting point is the percolation theory of hopping transport, according to which transport through the system is achieved via percolation in energy-position space. The average $\langle E - \mu \rangle$ is calculated by averaging the energy over the sites composing the percolation cluster, and the thermopower is given by

$$S = \frac{\langle E - \mu \rangle}{e T} = \frac{1}{e T} \frac{\int dE \left(E - \mu \right) \nu(E) p(E)}{\int dE \, \nu(E) \, p(E)},\tag{19}$$

where p(E) is the probability that a state of energy E belongs to the percolation cluster. The latter quantity is supposed to be proportional to the average number of bonds $N_b(E)$, given by

$$N_b(E) = \int dx \int dE' \,\nu(E') \,\theta\left(\sqrt{\frac{T_M}{T}} - \frac{2x}{\xi} - \frac{|E-\mu| + |E'-\mu| + |E-E'|}{2k_B T}\right),\tag{20}$$

under the assumptions leading to Eq. ((9)) (μ inside the band, low temperature and energy independent localisation length $\xi(E) = \xi(\mu)$) [26, 21]. The Heaviside function θ accounts for the existence of a percolating path, and restricts the energy range of integration to the window

^{*} We stress that the usual Mott formula for the thermopower, $S = (\pi^2 k_B^2 T/(3e)) \partial_E \ln \sigma|_{\mu}$ (σ being the electrical conductivity), does not apply in the VRH regime, as pointed out by Mott himself in [29]. Indeed, this formula has been derived by averaging $\langle E - \mu \rangle$ within the standard Boltzmann formalism, not suitable in the VRH regime where $\Delta \gg k_B T$.

 $[\mu - \Delta, \mu + \Delta]$. After integrating over the single spatial variable x (in 1D), one gets

$$p(E) \propto \theta \left(\Delta - |E - \mu|\right) \times \int_{\mu - \Delta}^{\mu + \Delta} dE' \nu(E') \left(1 - \frac{|E - \mu| + |E' - \mu| + |E - E'|}{2\Delta}\right) \theta \left(\Delta - |E - E'|\right).$$
(21)

Note that if μ lies outside the impurity band, electrons need to jump inside the latter by thermal activation before accessing the percolation cluster. In that case, Eqs. (20) and (21) have to be modified accordingly, by replacing μ by the energy ϵ_c of the closest band edge and by changing the energy range of integration to $[\epsilon_c, \epsilon_c + \Delta]$ (lower band edge) or $[\epsilon_c - \Delta, \epsilon_c]$ (upper band edge).

Eqs. (19) and (21) enable us to calculate the thermopower once the DOS $\nu(E)$ is known. Following Zvyagin's works [20, 21], we discuss below a few extreme cases where the DOS takes a simple form. Contrary to those works focused on three-dimensional bulk materials, we derive expressions for the thermopower of nanowires in the 1D case. Despite the simplicity of our approach, we will see in the next subsection that it enables us to qualitatively capture the typical behaviour of the thermopower and the role of the gate (see Sec. 2).

Let us first consider the case where (i) the DOS can be approximated by its first order expansion $\nu(E) \approx \nu(\mu) + (E - \mu) \partial_E \ln \nu(E)|_{\mu}$ in the interval $[\mu - \Delta, \mu + \Delta]$, and (ii) ν is expected to vary slowly at the scale of Δ , *i.e.* $\Delta \partial_E \ln \nu(E)|_{\mu} \ll 1$. Using Eqs. (19) and (21), one finds

$$S \approx \frac{k_B}{e} \left(\frac{k_B T_M}{4}\right) \partial_E \ln \nu(E)|_{\mu} .$$
⁽²²⁾

This shows that the thermopower should be temperature independent when the assumptions above are fulfilled, which is always the case at very low temperatures (bottom part of region (2a) in Fig. 3). Note that the same hypothesis for the DOS lead to the standard Mott formula (16) for the conductance: Eq. (22) describes the thermopower when Eq. (16) holds for the conductance.

Let us now consider the case where the impurity band edges are explored, say the lower one. In analogy to the previous section, using a rough step-like model for $\nu(E)$ provides useful insight. Using Eq. (17) for the DOS and Eq. (19), one gets for the thermopower

$$S = \frac{k_B}{e} \left(\frac{\epsilon_c - \mu}{2k_B T} + \frac{\Delta(T)}{2k_B T} \right) \quad \text{if} \quad \epsilon_c < \mu \quad \text{and} \quad \mu - \epsilon_c < \Delta,$$
(23a)

$$S = \frac{k_B}{e} \left(\frac{\epsilon_c - \mu}{k_B T} + \frac{\Delta(T)}{2k_B T} \right) \quad \text{if} \quad \epsilon_c > \mu \,, \tag{23b}$$

assuming $\mu(E) = 1$ in the energy window $|E - \mu| < \Delta [0 < E - \epsilon_c < \Delta]$ and 0 elsewhere. Similar formulas can be deduced by symmetry if the upper band edge is explored. The resulting thermopower behaviour as a function of temperature turns out to be rich. Indeed, depending on the position of μ with respect to the (bottom) edge ϵ_c of the DOS, and depending on the magnitude of Δ , the thermopower can be an increasing or decreasing function of T. If μ lies outside the impurity

We have also calculated the thermopower beyond this approximation, by plugging Eq. (17) for $\nu(E)$ into Eq. (21) for p(E). Instead of Eqs. (23a) and (23b), we find respectively

$$S = \frac{k_B}{e} \left[\frac{5(\epsilon_c - \mu)}{8k_B T} + \frac{3\Delta(T)}{8k_B T} + O\left(\frac{\epsilon_c - \mu}{k_B T}\right) \right],\tag{24a}$$

$$S = \frac{k_B}{e} \left[\frac{\epsilon_c - \mu}{k_B T} + \frac{3\Delta(T)}{8k_B T} \right] . \tag{24b}$$

The two sets of equations are obviously very similar. At a qualitative level of analysis, it is meaningless to favour one over the other.

band, the thermopower (in unit of k_B/e if not otherwise specified) is found to be a monotonically decreasing function of the temperature (see Eq. (23b)). On the other hand, if μ lies inside the band, close to the edge ϵ_c of the DOS, the thermopower increases with the temperature, reaches a maximum (at $k_BT = (\epsilon_c - \mu)^2/(16k_BT_M)$) and then starts to decrease (see Eq. (23a)).

Let us finally address the large temperature limit $(k_B T \gtrsim 2E_B)$, corresponding to region (4b) and the upper part of region (4a) in Fig. 3. In that case, all impurity band states are involved in thermoelectric transport, with $p(E) \approx 1$. As a consequence, the thermopower temperature behaviour is merely $S \sim T^{-1}$. Assuming a constant DOS, one gets

$$S = \frac{k_B}{e} \left(\frac{V_g - \mu}{k_B T} \right) \,. \tag{25}$$

4.2. Numerical results

For the model introduced in Sec. 2.3, we now study the thermopower by solving numerically the random resistor network (see Appendix A).

Fig. 6(a) gives the distribution P(S) of the thermopower S in the VRH regime, when the impurity band center (red curve) and lower edge (blue curve) are probed at μ . While the thermopower distribution is symmetric around a vanishing average value at the band center, it is shifted away from 0 and gets skewed close to the band edges. Such features can be easily understood: The level distribution becomes highly asymmetric with respect to μ when one probes the lower band edge with a positive gate voltage V_g . Consequently, an electron entering the nanowire from the left lead around μ finds more states *above* its energy than below. It has therefore a tendency to absorb energy in order to move to regions of higher DOS, before releasing it at the right side of the nanowire, as illustrated in Fig. 2. Recalling that $S = \langle E - \mu \rangle / (eT)$, one can thus explain why P(S) is shifted and skewed at finite V_q . Let us notice that such a skewness cannot be seen in the low-temperature coherent regime [19], where transport only involves electrons at energies very close to μ ; In that case, distributions are found to be shifted with V_q but always symmetric. Another important message of Fig. 6(a) is that for both values of V_g the thermopower distribution turns out to be independent of the nanowire length L. This is consistent with the observation that the thermopower is governed by the edges of the nanowire in the hopping regime, as recently pointed out in Ref. [18].

We then investigate the typical thermopower behaviour as a function of temperature and gate voltage, by extracting the median S_0 of the distribution P(S) for different sets of parameters. The temperature dependence of S_0 is shown in Fig. 6(b) for different values of the gate voltage, which have been chosen for scanning the vicinity of the lower band edge. The main observation is that our model predicts a huge enhancement of the thermopower around the band edges. Values larger that $10 k_B/e$ are obtained by properly tuning the strength of the gate voltage in the VRH regime. Other features of those curves are worth emphasizing:

- (i) S_0 is always positive in unit of k_B/e , hence negative in V K⁻¹ (since e < 0). This is expected since transport is due to electrons near the lower band edge, the sign of the thermopower reflecting the sign of the charge carriers.^{††}
- (ii) At low temperatures the typical thermopower can either increase or decrease with the temperature depending on the gate voltage. Roughly speaking, it increases inside the band and decreases outside, in agreement with the theoretical predictions (23a) and (23b), obtained assuming a step-like model for the DOS close to the band edge ϵ_c . Moreover, the position

[†]The occurrence of negative S_0 is nevertheless possible not far from the lower band edge, as soon as Δ is sufficiently small and the DOS slope at μ becomes strongly negative. In our model, such a negative slope occurs close to the band edges, as shown in Fig. 4.

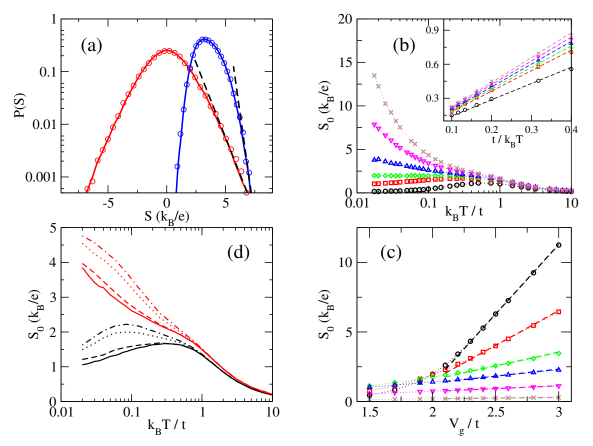


Figure 6. In all panels, unless specified, L = 200, $\mu = 0$, W = t and $\gamma_e = \gamma_{ep} = t$. (a) Thermopower distributions in the VRH regime, when μ lies in the bulk (left red curve, $V_g = 0$, W = 4t) or close to the edge (right blue curve, $V_g = 2.2t$, W = t) of the impurity band. Data are given for L = 200 (full lines) and L = 400 (circles). The straight dashed lines underline the exponential behaviour of the tails $\sim \exp\{-cS\}$ predicted in Ref. [18]. In both cases, $k_BT = t$. (b) Main panel: Typical thermopower as a function of T around the (lower) band edge. From the bottom to the top, the various curves correspond to $V_g/t = 1.5$ (O), 1.9 (\Box), 2.0 (\diamond), 2.1 (\triangle), 2.2 (∇) and 2.3 (\times). Dotted lines are guides to the eye. Inset: zoom at very large temperatures $k_BT \gtrsim E_B$. The fits $f(V_g)/T$ (dashed lines) confirm the expected behaviour $S_0 \sim T^{-1}$ (Eq. (25)). (c) Typical thermopower as a function of V_g , for $k_BT/t = 0.1$ (O), 0.2 (\Box), 0.5 (\diamond), 1.0 (\triangle), 2.5 (∇) and 10.0 (\times). At large V_g (when μ lies outside the band), dashed lines are linear fits with slope t/k_BT (Eq. (23b)). Dotted lines are guides to the eye to the eye. (d) Typical thermopower as a function of T, for electron-phonon coupling strength $\gamma_{ep}/t = 1$ (full line), 0.5 (dashed line), 0.1 (dotted line) and 0.05 (mixed line), at $V_g = 1.9t$ (black curves, bottom set) and $V_g = 2.1t$ (red curves, top set).

of the crossover between the two behaviours is found around $V_g - \mu \approx 2t$, a value consistent with our previous estimation of the (lower) band edge position of the Anderson model at $\epsilon_c \approx V_q - 2.2t$ (see Sec. 2.1).

- (iii) At high temperature (typically larger than the bandwidth), the curves converge to a T^{-1} behaviour, as shown in the inset of Fig. 6(b). The crude estimation (25) turns out to be satisfactory in this regime.
- (iv) In the low temperature limit and in the case where μ lies inside the band, the typical thermopower S_0 is expected to saturate, according to Eq. (22). Such a saturation is not observed in Fig. 6(b). Two reasons can be invoked. The first one is that Eq. (22) was actually derived under the assumption of a constant localisation length $\xi_i \approx \xi(\mu)$ while the numerical results reported here were obtained going beyond this approximation, by taking into account the energy dependency of the different localisation lengths ξ_i of sites *i*. In Appendix B, we show that under the assumption $\xi_i \approx \xi(\mu)$, S_0 indeed saturates at low temperature. The other possibility is simply that the saturation appears at lower temperatures, which are not reachable numerically because of round-off errors.
- (v) For high values of V_g , the typical thermopower seems to diverge as the temperature is lowered. It is obvious that the thermopower eventually decreases below a certain temperature, since all curves in Fig. 6(b) are known to drop down to zero in the zero-temperature limit (linearly with T and with a positive slope) [19].

In Fig. 6(c), we show how the typical thermopower depends on the gate voltage, for different values of the temperature. Approaching the edge of the impurity band, we see that S_0 increases, the effect being more pronounced at low temperatures. Outside the band, the behaviour of S_0 with V_g is perfectly well fitted by the formula $S_0 = (k_B/e)\left[\frac{V_g}{k_BT} + f(T)\right]$, as illustrated by the straight lines in Fig. 6(c). This linear enhancement of S_0 with V_g , as well as its range of validity, is consistent with the prediction (23b) and our initial estimation $\epsilon_c \approx V_g - 2.2t$ for the position of the lower band edge. Note however that Eq. (23b) does not capture the y-intercept $f(T) \approx 0.89 - 1.94/(k_BT)$ of the linear fits. On the other hand the fact that S_0 keeps increasing even outside the impurity band, when the conductance drops exponentially, may seem in contrast with recent experimental observations [5]. We think the explanation lies in the fact that, when the nanowire is almost completely depleted by V_g , the probability for an electron at μ to tunnel inside the band becomes extremely small, and so do the electrical and heat currents; consequently, they may be too hard to measure. Nonetheless their ratio, which gives the thermopower, remains formally well defined and finite.

We conclude our analysis by discussing the order of magnitude of our numerical results. In panels (a), (b) and (c) of Fig. 6, data was obtained taking $\gamma_e = \gamma_{ep} = t$ as input parameters of the model. In panel (d) we investigate how the typical thermopower depends on the choice of these parameters, finding that S_0 does not vary by more than 50% when the ratio γ_e/γ_{ep} is increased or decreased by an order of magnitude. Remarkably, at the lowest studied temperatures (in the VRH regime) and around the band edges, the typical thermopower is found to reach very large values of the order of $10 (k_B/e) \sim 1 \,\mathrm{mV \, K^{-1}}$. It is worthwhile to note that, despite the simplicity of the model, the order of magnitude of these results is comparable to recent measurements of thermopower in semiconducting nanowires [8, 47, 48, 5], showing strong thermoelectric conversion at the band edges.

5. Discussion and conclusion

We have studied thermoelectric transport in a disordered nanowire in the field effect transistor configuration, focusing on intermediate to high temperatures. More precisely, T was high enough

for inelastic processes (phonon-assisted hopping between localised states) to be dominant, but still such that $k_BT < 2E_B$, with $2E_B$ the spread in available nanowire states. Transport in this regime is typically of variable range hopping type [28, 29]. We have extended the Miller-Abrahams random resistor network model [27] to deal with band-edge transport, and performed accurate numerical analysis based on the 1D Anderson model. The thermopower shows remarkable gate- and temperature-dependent behaviour, whose features can be understood within a suitable generalization of Zvyagin's analytical treatment of VRH transport [20, 21]. In particular we have shown them to be largely independent of fine system details such as electron-phonon interaction strength or the specific form of the DOS. Our results are in line with numerous experimental observations [8, 47, 48, 5], confirming the great thermoelectric conversion potential of band-edge transport. Notice in particular that semi-quantitative agreement with observations was reached, though we stress that our treatment's strength lies in its general applicability rather than in its high precision – the latter being heavily dependent on fine details of each particular setup, such as materials involved, doping level/type, geometry and so on.

Let us now comment on certain limitations of our work. First, interactions have been neglected, except for the requirement of single-occupation of any given localised state [26]. Whereas this is appropriate in some cases, it is by no means a universally valid assumption. Indeed, numerous delicate issues related to the role of interactions in activated transport are discussed in [34] and references therein. Secondly, we have ignored phonon-drag effects, which is however a much safer bet. It is well known that the latter can play a prominent role in standard band transport – i.e. when electronic states are delocalised – but are irrelevant when transport is due to hopping between localised states[20, 21]. Finally, we used the Anderson model which is a single band model and hence neglected the possibility of temperature activated transport via other bands. This amounts to assuming that $k_BT < E_{act}$, where E_{act} is the interband spacing. E_{act} depends on the considered material, ranging from tens of Kelvin degrees for weakly doped crystalline materials, to hundreds of Kelvin degrees in amorphous materials [34].

Acknowledgments

This work has been supported by CEA through the DSM-Energy Program (project E112-7-Meso-Therm-DSM). We acknowledge stimulating discussions with Y. Imry, R. Jalabert, F. Ladieu, K. Muttalib and A. Parola.

Appendix A. Solution of the random resistor network

In linear response we assume that on each localised state the electron occupation is characterized by a *local distribution* [34, 18]:

$$f_i = f_i^0 + \delta f_i, \tag{A.1}$$

where f_i^0 is the Fermi distribution at equilibrium (i.e., evaluated at the reference values μ and T), and δf_i is the correction induced by the (small) applied bias $\delta \mu$. Linearizing Eqs.(13), and making use of Eqs. (7), (8), and (11), the hopping currents between each pair of localised states, and the tunnelling currents from/to the electrodes can be written in terms of "local potentials" U_i 's:

$$I_{ij} = G_{ij}(U_i - U_j),$$

$$I_{iL(R)} = G_{iL(R)}(U_i - U_{L(R)}),$$
(A.2)

where

$$G_{ij} = \frac{e^2}{k_B T} \gamma_{ij} f_i^0 (1 - f_j^0) (N_{ij} + 1/2 \mp 1/2),$$

$$G_{iL(R)} = \frac{e^2}{k_B T} \gamma_{iL(R)} f_i^0 (1 - f_i^0),$$

$$U_i = \frac{k_B T}{e} \delta f_i / [f_i^0 (1 - f_i^0)],$$

$$U_{L(R)}(E_i) = \frac{k_B T}{e} \delta f_{L(R)} / [f_i^0 (1 - f_i^0)].$$
(A.3)

In the above expressions, in case of double signs, the upper (lower) sign refers to $E_j > E_i$ ($E_j < E_i$). At steady state, according to Kirchoff's conservation law, the net electric current throughout every node *i* must vanish:

$$\left(\sum_{j\neq i} I_{ij}\right) + I_{iL} + I_{iR} = 0.$$
(A.4)

By plugging Eqs. (A.2), we end up with a set of L equations (one for every node i) to calculate the L local potentials U_i 's, which can be written conveniently in the matrix form:

$$\sum_{j} A_{ij} U_j = z_i, \tag{A.5}$$

where

$$A_{ij} = -G_{ij} \qquad (\text{for } i \neq j),$$

$$A_{ii} = \sum_{k \neq i} G_{ik} + G_{iL} + G_{iR},$$

$$z_i = G_{iL} \ (\delta \mu_L / e) \qquad (A.6)$$

In writing the expression for z_i , we exploited the fact that $\delta \mu_R = \delta T_R = 0$, having chosen to set the right terminal as reference (see Sec. 2).

Once the system is solved and the U_i are known, all the I_{ij} 's and $I_{iL(R)}$ can be calculated via Eqs. (A.2). The electric and heat current can be computed by summing the outgoing contributions from the left (right) lead toward *every* states in the system:

$$I_{L}^{e} = -\sum_{i} I_{iL} = \sum_{i} I_{iR},$$

$$I_{L(R)}^{Q} = \sum_{i} \left(\frac{E_{i} - \mu_{L(R)}}{e}\right) I_{L(R)i}.$$
(A.7)

Appendix B. Calculation of the hopping probability

Miller and Abrahams[27, 34] described how to calculate the hopping probability γ_{ij} between two donors *i* and *j* in a 3D semiconductor, mediated by the absorption or emission of a phonon. When the distance between the donors is large, they obtain for γ_{ij} an expression which depends on the (weak) overlap between the donor wavefunctions and on the mutual electrostatic effect between them:

$$\gamma_{ij} \propto \left| \langle \psi_i | \frac{e^2}{\kappa |\boldsymbol{r} - \boldsymbol{r}_i|} | \psi_j \rangle - \langle \psi_i | \psi_j \rangle \langle \psi_i | \frac{e^2}{\kappa |\boldsymbol{r} - \boldsymbol{r}_j|} | \psi_i \rangle \right|^2.$$
(B.1)

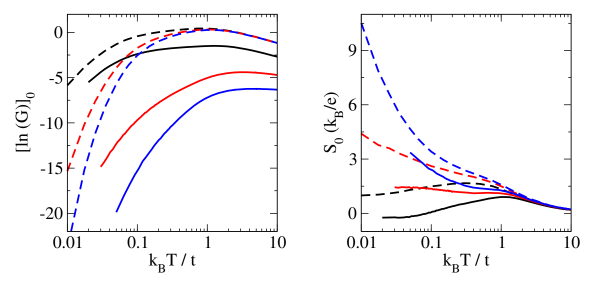


Figure B1. Typical logarithm of electrical conductance (left) and typical thermopower (right) of a disordered nanowire as a function of temperature, and for different values of the applied gate voltage. Full lines refer to the approximation in which all $\xi_i = \xi(\mu)$, while dashed lines refer to the improved theory in which the energy dependence of $\xi_i = \xi(E_i)$ is taken into account for evaluating the transition rates. In each set, from the top to the bottom (left panel) or reversely (right panel), the curves correspond to $V_g = 1.9t$ (black), $V_g = 2.1t$ (red) and $V_g = 2.2t$ (blue). Other parameters: L = 200, W = t, $\mu = 0$, $\gamma_e = \gamma_{ep} = t$.

If the donor wavefunctions ψ_i and ψ_j are characterized by the same decay length ξ , Eq. (B.1) can be simplified [27, 34]

$$\gamma_{ij} \propto \exp(-2|\boldsymbol{r}_i - \boldsymbol{r}_j|/\xi). \tag{B.2}$$

If the decay lengths of ψ_i and ψ_j are different ($\xi_i \neq \xi_j$), a rigourous evaluation of γ_{ij} from Eq. (B.1) may be complicated, but the key point is that it will always be proportional to the overlap $\langle \psi_i | \psi_j \rangle$. Hence, we can write it in the form

$$\gamma_{ij} \propto \left| \langle \psi_i | \psi_j \rangle \right|^2 \sim \left| \mathcal{C}_i \exp(-r_{ij}/\xi_i) + \mathcal{C}_j \exp(-r_{ij}/\xi_j) \right|^2, \tag{B.3}$$

where $r_{ij} = |\mathbf{r}_i - \mathbf{r}_j|$ is the distance between *i* and *j*, and the coefficients C_i and C_j depend on ξ_i, ξ_j and r_{ij} . The explicit form of these coefficients will take into account all details concerning the wavefunction overlap $\langle \psi_i | \psi_j \rangle$. In 1D the calculation becomes simpler and leads to Eq. (10). Extending a theory originally developed for lightly doped cristalline semiconductors (where the decay length is the donor Bohr radius) to Anderson insulators (where the decay length becomes the localisation length), Ambegaokar *et al.*[26] have used Eq. (B.2) for describing the hopping probability. For similar reasons, we use Eq. (10) in our numerical calculations, for both *G* and *S*, taking for ξ_i and ξ_j the localisation length of two Anderson localised states.

In order to estimate the difference between taking $\xi(\mu)$ or $\xi(E)$ when computing the transition rates (Eqs.(7) and (11)), we have calculated the typical logarithm of the conductance and the typical thermopower as functions of the temperature in the two cases: Fig. B1 shows that there is no qualitative difference between the curves computed using $\xi(\mu)$ (full lines) and $\xi(E)$ (dashed lines). The main effect of taking into account the localisation length energy dependence is that, according to Eq.(10), all transitions toward the more delocalised states around the band center are favoured. This leads to a much better conductance especially at low temperatures, where the difference could be of several orders of magnitude; on the other hand, the effect on the thermopower is weaker.

References

- [1] Chen G, Dresselhaus M S, Dresselhaus G, Fleurial J P and Caillat T 2003 Int. Mater. Rev. 48 45
- [2] Hicks L D and Dresselhaus M S 1993 Phys. Rev. B 47 16631
- [3] Curtin B M, Fang E W and Bowers J E 2012 J. Electron. Mat. 41 887
- [4] Blanc C, Rajapbour A, Volz S, Fournier T and Bourgeois O 2013 Appl. Phys. Lett. 103 043109
- [5] Brovman Y M, Small J P, Hu Y, Fang Y, Lieber C M and Kim P 2013 arXiv:1307.0249
- [6] Stranz A, Waag A and Peiner E 2013 J. Electron. Mat. 42 2233
- [7] Karg S, Mensch P, Gotsmann B, Schmid H, Kanungo P D, Ghoneim H, Schmidt V, Björk M T, Troncale V and Riel H 2013 J. Electron. Mat. 42 2409
- [8] Roddaro S, Ercolani D, Safeen M A, Suomalainen S, Rossella F, Giazotto F, Sorba L and Beltram F 2013 Nano Lett. 13 3638
- Hochbaum A I, Chen R, Delgado R D, Liang W, Garnett E C, Najarian M, Majumdar A and Yang P 2008 Nature 451 163
- [10] Tilke A, Pescini L, Erbe A, Lorenz H and Blick R H 2002 Nanotechnology 13 491
- [11] Bourgeois O, Fournier T and Chaussy J 2007 J. Appl. Phys. 101 016104
- [12] Boukai A I, Bunimovich Y, Tahir-Kheli J, Yu J K, Goddard W A and Heath J R 2008 Nature 451 168
- [13] Galli G and Donadio D 2010 Nat. Nanotech. 5 701
- [14] Heron J S, Bera C, Fournier T, Mingo N and Bourgeois O 2010 Phys. Rev. B 82 155458
- [15] He Y and Galli G 2012 Phys. Rev. Lett. 108 215901
- [16] Hu M and Poulikakos D 2012 Nano Lett. 12 5487
- [17] Jiang J H, Entin-Wohlman O and Imry Y 2012 Phys. Rev. B 85 075412
- [18] Jiang J H, Entin-Wohlman O and Imry Y 2013 Phys. Rev. B 87 205420
- [19] Bosisio R, Fleury G and Pichard J L 2014 New J. Phys. 16 035004
- [20] Zvyagin I P 1973 Phys. Stat. Sol. (b) 58 443
- [21] Zvyagin I P 1991 Hopping Transport in Solids (ed. by M. Pollak and B. I. Shklovskii (North-Holland, Amsterdam))
- [22] Poirier W, Mailly D and Sanquer M 1993 Phys. Rev. B 59 10856
- [23] Tilke A, Blick R H, Lorenz H, Kotthaus J P and Wharam D A 1999 Appl. Phys. Lett. 75 3704
- [24] Dayen J F, Wader T L, Rizza G, Golubev D S, Cojocaru C S, Pribat D, Jehl X, Sanquer M and Wegrowe J E 2009 Eur. Phys. J. Appl. Phys. 48 10604
- [25] Rodin A S and Fogler M M 2010 Phys. Rev. Lett. 105 106801
- [26] Ambegaokar V, Halperin B I and Langer J S 1971 Phys. Rev. B 4 2612
- [27] Miller A and Abrahams E 1960 Phys. Rev. 120 745
- [28] Mott N F 1969 Phil. Mag. 19 835
- [29] Mott N F and Davis E A 1979 Electronic Processes in Non Crystalline Materials (Clarendon, Oxford, (2nd ed.))
- [30] Kurkijärvi J 1973 Phys. Rev. B 8 922
- [31] Raikh M E and Ruzin I M 1989 Sov. Phys. JETP 68 642
- [32] Serota R A, Kalia R K and Lee P A 1986 Phys. Rev. B 33 8441
- [33] Pollack M 1972 J. Non-Cryst. Solids 11 1
- [34] Shklovskii B and Efros A 1984 Electronic Properties of Doped Semiconductors (Springer-Verlag, Berlin)
- [35] Callen H 1985 Thermodynamics and an Introduction to Thermostatics (John Wiley and Sons, New York)
- [36] Derrida B and Gardner E 1984 J. Physique 45 1283
- [37] Lee P A 1984 Phys. Rev. Lett. 53 2042
- [38] Kwasnick R F, Kastner M A, Melngailis J and Lee P A 1984 Phys. Rev. Lett. 52 224
- [39] Webb R A, Hartstein A, Wainer J J and Fowler A B 1985 Phys. Rev. Lett. 54 1577
- [40] Ladieu F, Mailly D and Sanquer M 1993 J. Phys. I France 3 2321
- [41] Hasko D G, Cleaver J R A, Ahmed H, Smith C G and Dixon J E 1993 Appl. Phys. Lett. 62 2533
- [42] Han M Y, Brant J C and Kim P 2010 Phys. Rev. Lett. 104 056801
- [43] Ladieu F and Bouchaud J P 1993 J. Phys. I France 3 2311
- [44] Rodin A S and Fogler M M 2011 Phys. Rev. B 84 125447
- [45] Grant A J and Davis E A 1974 Sol. State Comm. 15 563
- [46] Rodin A S and Fogler M M 2009 Phys. Rev. B 80 155435
- [47] Moon J, Kim J H, Chen Z, Xiang J and Chen R 2013 Nano Lett. 13 1196
- [48] Curtin B M, Codecido E A, Krämer S and Bowers J E 2013 Nano Lett. 13 5503