Nonlinearity exponent in low dimensional nanotubes and nanowires

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A scaling formalism is used to analyze non-Ohmic conductance-voltage data in nanowires of an isolated potassium manganese oxide K0.27MnO2·0.5H2O and a poly(3,4-ethylenedioxythiophene) (PEDOT) and nanotubes of conducting polypyrrole at different temperatures. This scaling analysis provides a single voltage scale for non-Ohmic conduction which scales with Ohmic conduction with an exponent xL called nonlinearity exponent. A detailed analysis reveals that the onset exponents xL are different in different systems. Non-Ohmic conduction and the nonlinearity exponents in these low-dimensional nanostructures are analyzed within the framework of scaling approach and existing theoretical models.

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1. Introduction

Quasi-one-dimensional nanostructures such as nanotubes [1–7], nanowires [6–9], and nanofibres [3–5,10] are being studied extensively due to their importance in fundamental research and potential applications in nanoscale devices [11,12]. These systems are widely used as sensors in molecular electronics [13], as field effect transistors in nanoelectronics [14,15] and as chemical and bio-sensors in bio-nanotechnology and medicine [16]. These low-dimensional materials especially nanowires and nanotubes are prepared with chemical doping and are known to possess structural inhomogeneity having ordered conduction regions separated by small insulating barriers. This complex micro-structure of the low dimensional systems affects the various physical properties in particular the electrical transport [17,18]. In order to explore the salient features of the electrical transport and their possible applications, a number of experimental [4,6,7] and theoretical [19–22] studies have been made in these low dimensional nanostructures. However, its systematic description mainly in the non-Ohmic region is still lacking.

In low-dimensional systems, several physical properties show interesting behavior in contrast to their bulk counterpart. For example, electrical conductivity [23] and elastic constants [24] increase sharply with the decrease in diameter of the conducting polymer nanowires and nanotubes, and magneto-resistance of isolated polymer nanowires/tubes at low-temperatures is much smaller than that of the bulk pellets or films [25]. A number of conduction mechanisms such as variable range hopping due to Mott (M-VRH) [19] and its modification by Efros and Shklovskii (ES-VRH) [22] in the presence of coulomb interaction and fluctuation induced tunneling (FIT) due to Sheng et al. [20] have been considered to describe the electrical transport behavior in the low-dimensional materials [6,16,26,27] at low field strengths. In M-VRH model, an electron near the Fermi level hops between localized states separated by certain physical distances and having energy differences by absorbing phonons. The Ohmic conductance Σ0 is obtained by optimizing the electron hopping paths using the relation:

Σ0(T) = Σm exp \left[-\left(\frac{T_0}{T}\right)^m\right] \tag{1}

Here Σm is the conductance pre-factor, T is the temperature and the exponent is m = 1/(d + 1) with d is the dimensionality of the system under consideration. The characteristic temperature T0 is given by T0 = 4π/3N(EF)Kbd^3 where N(EF) is the density of states at the Fermi level, Kβ is the Boltzmann constant and d is the localization length. The conduction behavior at low field strengths in three as well as two dimensional systems is well described by this M-VRH model.
p-toluensulfonate (PTS)-doped PPy films [28] are reported to yield $m = 1/3$ suggesting two-dimensional nature. The exponent $m$ in single crystals of PTS-doped polydiacetylene (PDA) [29] is found to be $\sim 0.65 - 0.70$ apparently demonstrating quasi-one-dimensional nature of the system. In ES-VRH model [22], electron–electron interaction opens up a coulomb gap in the density of states at the Fermi level and the value of $m$ becomes 1/2 which is independent of $d$. In this model, $T_0$ is given by

$$T_0 = \frac{2.8 e^{d}}{4 \pi e \varepsilon \kappa_{a} d}, \quad (2)$$

where $\varepsilon$ is the dielectric constant. PEDOT [30] and Polypropylene (PPy) films [31] tend to follow ES-VRH particularly at low temperatures with $m = 1/2$.

Non-Ohmic conduction is a common feature of the low-dimensional nanotubes and nanowires [6,16,26,27]. For small applied voltage $V$, the conductance $G(V)$ at a fixed temperature remains almost constant to its zero-voltage Ohmic value $G_0$. However with an increase in $V$, $G(V)$ increases from $G_0$ showing the non-Ohmic characteristics. The value of $V$ at which $G(V)$ deviates from $G_0$ for the first time is known as the onset voltage $V_0$ and the corresponding current is known as the onset current $I_0$ [32]. Such non-Ohmic conduction has been observed in films [32], composites [33], conducting polymers [34] and manganites [35,37]. This common feature of the non-Ohmic conduction in different types of disordered systems is expressed using the scaling relation [34–37]:

$$\frac{\Sigma(V)}{\Sigma_0} = g \left( \frac{V}{V_0} \right). \quad (3)$$

where $g(\frac{V}{V_0})$ is a scaling function. For $V \leq V_0$, $g(\frac{V}{V_0}) \approx 1$ and this corresponds to the fact that the conductance remains almost constant to $\Sigma_0$. At voltages $V \gg V_0$, $g(\frac{V}{V_0}) \approx 1$. Thus, the voltage $V_0$ at a fixed temperature separates the Ohmic regime from the non-Ohmic regime along the voltage axis. It has been shown by several authors that the scaling description given by Eq. (3) is applicable to various disordered systems like amorphous and doped semiconductors [34], films [34], composites [33,36], conducting polymers [34] and lightly doped manganites [35,37]. This onset voltage $V_0$ scales with zero-voltage Ohmic conductance $\Sigma_0$ [34–36] as

$$V_0 = A_T \Sigma_0^{\alpha}, \quad (4)$$

where $A_T$ is a constant. The value of $A_T$ is determined by fixing the voltage scale $V_0$. $x_T$ is the nonlinearity exponent, also known as the onset exponent. It has been shown in Refs. [34,35] that $T$ does not enter explicitly in Eq. (3) but does so through the parameter $\Sigma_0$ which depends on temperature. Thus, the voltage scale $V_0$ for nonlinearity is determined solely by the Ohmic conductance $\Sigma_0$.

Experimental results [34,35,37] show that disordered systems possess a single voltage scale $V_0$ and the non-Ohmic conduction takes place upon the application of only a few volts across the samples. Similar scaling description of AC conduction has been recently observed by Nandi et al. [38] in various disordered systems.

Several theoretical models such as variable range hopping under field [39], extended fluctuation induced tunneling (FIT) [20,40] and multi-step tunneling through n-localized states due to Glazman and Matveev (GM) [21] have been widely used for the analysis of non-Ohmic conduction in disordered systems. M-VRH is basically a zero voltage model to provide the conductance of disordered systems through hopping via localized states. At finite electric field, the accessible states are aligned to the occupied states allowing charge carriers to move along the field via phonon-assisted hopping. With further increase in field, the number of the aligned states increases near the onset field $F_0$ and the conductivity increases. At higher electric field strengths $F$, the energy $eF_0$ gained by an electron may become comparable to $K_B T$ and leads to deviation from Ohmic behavior. $R_0$ is the mean hopping length and $e$ is the electronic charge. Theories [41,42] predict two characteristic field scales $F_1$ and $F_0$ corresponding to hopping and localization lengths respectively such that the non-Ohmic conductivity at intermediate field strengths $F < K_B T/e a$ is given by

$$\sigma(T, F) = \sigma(T, 0) \exp \left( \frac{eF}{K_B T} \right). \quad (5)$$

$\sigma(T, 0) = \sigma_0$ is the Ohmic conductivity and $L$ is a length related to the hopping length $R_0$. Clearly, the field $F_1 = K_B T/eL$ could be identified as the onset field scale $F_0$ as discussed above. In the limit of large field strengths $F \gg K_B T/ea$, theories [42] agree with ‘activationless’ hopping at least qualitatively. In such situation the energy gained by an electron is large enough to hop without absorbing any phonons. As a result, the conductivity becomes independent of temperature and is given by

$$\sigma \sim \exp \left( - \frac{F_0}{F} \right)^m, \quad (6)$$

Here $m$ and $T_0$ are same as in Eq. (1). $\lambda$ is a numerical constant equal to unity [42] when $m = 1/4$.

Glazmann and Matveev [21] considered the process of multi-step indirect tunneling through n-localized states in a disordered system and proposed the following expression for conductance through tunnel barriers of an amorphous material under external bias $V$ ($eV > K_B T$): $p_n = n - 2/(n + 1)$:

$$\Sigma = \Sigma_d + \sum_{n=1}^{N} \Sigma_n V^{p_n} = \Sigma_0 + \Sigma_1 V^{1.33} + \Sigma_2 V^{2.5} + \Sigma_4 V^{3.5} + \Sigma_5 V^{4.75} + \Sigma_6 V^{5.71} + \ldots. \quad (7)$$

Here $\Sigma_d = \Sigma_4 + \Sigma_1$, and $\Sigma_2$ accounts for the direct tunneling and $\Sigma_1$ for the elastic resonant tunneling via one localized state. Each term (for $n > 1$) in the series arises out of the events when a number of localized states happen to be arranged physically as well as energetically in such a way that an electron can traverse a sample length via multi-step inelastic tunneling. Each term may be said to constitute a separate channel of conduction. Thus, the macroscopic nonlinearity in this GM-model results from two contributions: appearance of new channels with increasing bias and nonlinearity of each such channel. This multi-step tunneling model has been widely used to explain relevant data in various tunnel junctions [43], lightly doped conducting polymer [34], and manganites [35,44,45]. Following the procedure adopted by Talukdar et al. [34], Eq. (7) can be made compatible with the scaling given in Eq. (3) by assuming that for $n > 2$, the coefficients satisfy the following relation

$$\Sigma_n = c_n \Sigma_0 V_{0}^{\alpha_n}. \quad (8)$$

Here $c_n$ are constants and can be determined if coefficients $\Sigma_n$ and onset voltage $V_0$ are known. Using this expression, Eq. (7) can be written as

$$\Sigma \Sigma_0 = 1 + c_1 \left( \frac{V}{V_0} \right)^{1.33} + c_2 \left( \frac{V}{V_0} \right)^{2.5} + c_4 \left( \frac{V}{V_0} \right)^{3.5} + c_5 \left( \frac{V}{V_0} \right)^{4.75} + c_6 \left( \frac{V}{V_0} \right)^{5.71} + \ldots \quad (9)$$

Eq. (9) is thus really a scaling function and implies the existence of a single bias scale $V_0$ as in Eq. (3).

In this report, the experimental results of electrical transport in a number of low-dimensional nanowires and nanotubes over a wide range of temperatures are presented both in Ohmic and non-Ohmic regimes. Attempts are made to test the validity of the scaling function $g(x)$ given by Eq. (3) and to extract the onset exponents $x_T$ defined in Eq. (4). Attention is also paid to find the nature
of the voltage dependence of the conductance both at Ohmic and Non-Ohmic regimes. Experimental details are given in Section 2. In Section 3, the data are presented, analyzed and discussed by scaling approach. Finally, we conclude the results in Section 4.

2. Experiments

Crystalline nanowires of K$_{0.27}$MnO$_2$·0.5H$_2$O were synthesized by the hydrothermal method. Analytical grade KMnO$_4$, MnCl$_2$, and KOH were obtained from Shanghai Chemical Co., and used without further purification. In a typical synthesis procedure, 1 mmol KMnO$_4$ and 1.5 mmol MnCl$_2$ were dissolved in 50 ml deionized water, then 40 g KOH was added into the solution under drastic stirring as a mineralizer. Finally, the solution was poured into a 100 ml Teflon vessel, sealed in a stainless steel autoclave, and maintained at 265 °C for 10 h under autogenous pressure. After the autoclave was cooled to room temperature, the final product was filtered off and washed sequentially with deionized water and absolute ethanol, then dried in ambient air at 60 °C for 2 h. More details can be found in [26,25]. Scanning electron microscopy (SEM) images of the nanowires shows that the diameter of the resulting nanowires is about 50–200 nm, and the length can be up to hundreds of micrometers [25]. Polypyrrole (PPy) nanotubes used in this work were prepared by using the typical template-free self-assembly method [6,27]. Pyrrole monomer (0.002 mol) and p-toluene sulfonic acid (PTSA, 0.001 mol) were mixed as dopant in 10 ml of distilled water. The mixture reacted and formed a transparent solution of PTSA–pyrrole salt and was cooled in an ice bath. Then an aqueous solution of ammonium persulphate (0.002 mol in 5 ml of distilled water) cooled in advance was added slowly into the PTSA–pyrrole salt solution. After reacting for 15 h in the ice bath, the precipitate was filtered out from the solution and washed sequentially with distilled water, ethanol, and ether several times. Finally the solution was dried for 24 h at room temperature in dynamic vacuum. PEDOT nanowires with a diameter of about 92 nm were prepared by using the template method according to a typical synthesis procedure as in Ref. [27].

Two Pt microleads in a K$_{0.27}$MnO$_2$·0.5H$_2$O nanowire and four Pt microleads in PPy nanotube and PEDOT nanowires were fabricated by the focused-ion beam deposition technique (Dual-Beam 235 Focused-Ion Beam System from FEI Company; the working voltage of the system was 30 kV and the focused-ion beam current was 0.5H$_2$O nanowire system. The panels (a), (b), (c) and (d) of Fig. 3 show the results of different samples of nanostructured systems. For clarity, the data of PPy-HQSA nanotube and PEDOT nanowire have been shifted upward by a factor of 10. Solid lines are linear fits to the data.

3. Results and discussion

Fig. 1 shows the variation of zero-voltage Ohmic conductance $\Sigma_0$ with temperature of four samples: K$_{0.27}$MnO$_2$·0.5H$_2$O nanowire of diameter 148 nm, PPy nanotube of diameter 160 nm, PPy-HQSA nanotube and PEDOT nanowire. The Ohmic conductance data of PPy-HQSA nanotube and PEDOT nanowire are multiplied by 10 to show them separately in the figure. The solid lines are fits to $\Sigma_0(T)$ data of all the samples with the variable range hopping given by the Efros–Shklovskii (ES) law: $\ln\Sigma_0 \sim T^{-1/2}$. These fits are extended over several orders of magnitude of Ohmic conductances $\Sigma_0$, e.g. three orders of magnitude in K$_{0.27}$MnO$_2$·0.5H$_2$O nanowire and five orders of magnitude in PPy nanotube. Excellent linearity of the fits to each curve indicates that within the measured range of temperature all the samples obey ES-type variable range hopping conduction.

The voltage dependence of conductance of a K$_{0.27}$MnO$_2$·0.5H$_2$O nanowire of diameter 148 nm and a PPy nanotube of diameter 160 nm are shown in Fig. 2(a) and (b) respectively. The data collapse into a master curve by data collapse method. The scaled data of the panels (a) and (c) correspond to the $\Sigma(V)$–$V$ data of the Fig. 2(a) and (b) respectively. The data collapse method for obtaining the master curve is discussed in detail for K$_{0.27}$MnO$_2$·0.5H$_2$O nanowire system. The $\Sigma(V)$–$V$ curve at 100 K is considered appropriate for starting the scaling as the conductance possesses predominantly Ohmic and minimum non-Ohmic regime. The conductance at 100 K was divided by its Ohmic value $\Sigma_0$ and the voltage is kept unaltered (i.e. divided by 1). This value (=1) of voltage is considered as the onset voltage $V_0$ corresponding to the temperature 100 K. In fact for the first $\Sigma(V)$–$V$ curve at a
particular temperature, any arbitrary choice for $V_0$ could be adapted as far as data collapse is concerned. For the next lower temperature (80 K), the conductance was divided by $\Sigma_0$ as before but the voltage was divided by a factor in such a way that this ratio becomes reliable due to fluctuation. These values of $\Sigma_0$ are shown in column III of Table 1.

The last section dealt with the results of the variation of dc conductance as a function of voltage both in linear and nonlinear regimes in low-dimensional $K_{0.27}$MnO$_2$-0.5H$_2$O nanowire, PEDOT nanowire and PPy nanotube. The fibril chain-like constituents [5] make these systems structurally inhomogeneous. These polymeric chains are ordered in some regions, and disordered in some other regions. This complex structure consists of long conducting pathways separated by small insulating barriers. Electrons in the conduction regions interact through Coulomb repulsion and open up a soft gap in the density of localized states at the Fermi level resulting ES-VRH conduction as seen in Fig. 1. This interaction in nanostructures is enhanced due to the confinement of electrons and is almost 20–40 times larger than that for bulk samples [5]. At sufficiently low temperatures, the zero bias anomaly [25] and smooth crossover from M-VRH to ES-VRH [5] confirm that the Coulomb gap arising out of the electron–electron interaction indeed influences the hopping transport in the low-dimensional nanostructures. The diameter of the nanowires and nanotubes (of the order of 50–200 nm) is much larger than the localization length (less than 15 nm) of the electrons confirming the three-dimensional character of the system in regard to the motion of the electrons. This indicates that temperature dependence given by Eq. (1) with $m = 1/2$ is due to ES-VRH not the one-dimensional M-VRH.

The $\Sigma(V)−V$ characteristics of all the systems under consideration at different $T$ reveal a common feature that the increase in conductance $\Delta \Sigma (=\Sigma(V)−\Sigma_0)$ increases with decrease in temperature. Here $\Sigma(V)$ is the conductance at the highest voltage corresponding to a particular temperature. This $\Delta \Sigma$ is different in different systems in the same range of $T$. For example, $\Delta \Sigma$ is nearly three orders of magnitude in PPy nanotube but it is only a factor of two in $K_{0.27}$MnO$_2$-0.5H$_2$O nanowire. Different values of $\Delta \Sigma$ indicate the different degree of nonlinearity in different systems.

The non-Ohmic conduction in these nano-structured systems can be qualitatively understood in the following way: due to the presence of defects, nano-tubes and nano-wires do not tend to be single crystal over the whole length of the sample and can be viewed as an one-dimensional array of small crystalline grains separated by multiple tunneling junctions. At low bias, conduction occurs through these one-dimensional conducting arrays resulting Ohmic character of conductance. With the increase in bias, the rate of tunneling through these multiple tunnel junctions increases and as a result conductance also increases with voltage. As the temperature increases, electrons from different conducting regions start hopping giving larger conductance. The tunnel junctions of relatively larger separation start contributing in the conduction at
higher bias. This indicates that $V_0$ increases with $T$ as pointed out by the orientation of the dotted lines in Fig. 2(a) and (b). Such non-Ohmic conduction has also been observed in several low-dimensional nanostructures e.g. p-toluensulfonate (PTS)-doped polydiacetylene (PDA) single crystal [29], individual iodine-doped polyacetylene nanofibre [40], Au-PPy-Au and Pt-PPy-Au nanowires [9], polyaniline nanotube [7], CdS nanorope [7], etc. PTS-doped PDA single crystals [29] are quasi-one-dimensional in nature, consisting of weakly coupled linear parallel chains of covalently bonded carbon atoms and obey VRH conduction with $m = 1/2$. The non-Ohmic behavior of conductivity in this system was also found to be enhanced with the lowering of temperature. Individual
iodine-doped polycacetylene nanofibre [40] follows activated-type conduction at high temperatures but Zener-type tunneling at low temperatures. This system also shows nonlinear conductance characteristics at different temperatures. Gence et al. [9] observed electrical nonlinearity in Au-PPy-Au and Pt-PPy-Au nanowires in the temperature range from room temperature down to 0.5 K. The low temperature zero-voltage conductance data of the single nanowire of diameter 70 nm and multi-nanowire samples embedded in polycarbonate were described by 3 d M-VRH conduction mechanism [9]. The nonlinearity in these systems was successfully explained by the hopping relations given by Eqs. (5) and (6) respectively in the intermediate and high field strengths signifying the existence of two field scales in the systems corresponding to hopping and localization lengths.

The panels (a), (b), (c) and (d) of Fig. 3 indicate that the \( \Sigma(V) \)–\( V \) data of these nanostructured systems at different temperatures do indeed collapse into a master curve. The maximum value of the normalized conductance \( \Sigma(V)/\Sigma_0 \) varies from system to system: by a factor of two in \( \text{K}_0.27\text{MnO}_2\) 0.5H\(_2\)O nanowire and PEDOT nanowire, four orders of magnitude in PPy nanotube and one order of magnitude in PPy-HQSA nanotube. But the normalized voltage \( V/V_0 \) is extended over at least five orders of magnitude. This does signify that the description of non-Ohmic conduction by the scaling function (3) is valid in these nanostructured systems. Scaled conductance versus scaled voltage data shown in Fig. 3(b), (c) and (d) were fitted with the expression:

\[
\frac{\Sigma(V)}{\Sigma_0} = 1 + C_1 \left( \frac{V}{V_0} \right)^{x_1} + C_2 \left( \frac{V}{V_0} \right)^{x_2} \tag{10}
\]

where \( C_1, C_2, x_1, \) and \( x_2 \) are the fitting parameters. These parameters have values (0.0452, 0.0, 0.66 and 0.0), (0.033, 4.5 \times 10^{-4}, 1.03 and 2.48) and (0.056, 0.0, 0.63 and 0.0) respectively for PEDOT nanowire, PPy nanotube and PPy-HQSA nanotube. The values of the parameters \( c_2 \) and \( x_2 \) are different in different systems. For example, \( x_2(=1.030) \) in PPy nanotube is much larger than in PEDOT nanowire (\( x_1 = 0.66 \)) and in PPy-HQSA (\( x_1 = 0.63 \)) nanotube signifying the fact that the electron–electron interaction in this nanotube is much stronger than the other two systems. \( x_2(=2.48) \) in PPy nanotube is very close to the power of the third term (=2.50) of the GM expression given in Eq. (9). This indicates that at higher voltages, the conduction channels mainly consist of three localized states. With the increase in voltage, these channels increase in number and provide paths for the flow of current causing the non-Ohmic conduction. These facts reflect that at lower voltages, the electron–electron interaction is prominent whereas at higher voltages three-step tunneling dominates the non-Ohmic conduction. The smaller value of the coefficient \( c_2(=4.5 \times 10^{-8}) \) indicates that the number of such three-step channels is small. Within the measured range of voltages, the second term of Eq. (10) is not required while fitting the scaled conductance versus voltage data of PEDOT nanowire and PPy-HQSA nanotube. This indicates that multi-step tunneling has no contribution to conductance in these two systems and the increase in normalized conductance is small as observed in the scaled plots shown in the panels (b) and (d) of Fig. 3. But in PPy nanotube, the multi-step tunneling is operative at higher voltages indicating huge \( (\sim 10^8) \) increase in normalized conductance.

Insets of the panels (a), (b), (c) and (d) of Fig. 3 show the log–log plot of the onset voltages \( V_0 \) corresponding to each temperature against the Ohmic conductances \( \Sigma_0 \). Fits to the data with Eq. (4) provide the values of \( x_T \) which are all positive as shown in Table 1. This indicates that \( V_0 \) increases with temperature \( T \) in these nanowire and nanotube systems. Further it is observed that in nanowire systems, \( x_T \) is relatively higher compared to the value of \( x_T \) in nanotubes. This indicates that a sample of nanowire becomes non-Ohmic at a voltage higher than that required for a sample of nanotube of comparable conductance. The values of \( x_T \) for two samples of PEDOT nanowire of diameters 92 and 95 nm indicate that similar-sized PEDOT nanowires have similar values of \( x_T \). Except in PPy nanotube of diameter 160 nm, the values of \( x_T \) were found to be large compared to the value in composites [32], lightly doped conducting polymers [34] and manganites [35,37]. This indicates that nanowires/nanotubes possess strong electrical nonlinearity. In lightly doped conducting polymer \( x_T \) was probed experimentally along different paths by varying conductance by several variables e.g. temperature \( T \), disorder \( D \) and magnetic field \( B \) [34] and was found to be positive, negative or even zero. The values of \( x_T \) of two such systems like PTS-doped PDA single crystals and individual iodine doped polycacetylene nanofibre are shown in Table 1. These values of \( x_T \) were successfully explained by the model of multi-step tunneling via n-localized states introduced by Glazman and Matveev [21]. It should be mentioned here that all these nanowires/nanotubes possess a single voltage scale \( V_0 \) within the measured range of voltage and temperature. The existence of such a single voltage scale \( V_0 \) has also been experimentally verified by Talukdar et al. [34] in several lightly doped conducting polymers available in bulk, film and single crystal forms. This experimental observation is contrary to the theoretical prediction of two voltage scales [41,42,46] corresponding to hopping and localization lengths in disordered systems. The field dependent conductance data of the nanostructured systems are successfully described by tunneling mechanism influenced by electron–electron interaction through the multiple tunneling junctions.

4. Conclusion

A comprehensive quantitative description of the variation of conductance of low-dimensional nanotubes and nanowires is presented over the temperature range from room down to 20 K and voltage range up to 6 V. Within the measured range of temperature, Ohmic conductance follows ES-type variable range hopping signifying the influence of electron–electron interaction on the conduction mechanism. The \( \Sigma(V) \)–\( V \) curves were found to be non-Ohmic and properly described by a single voltage scale \( V_0 \) corresponding to each temperature. This onset voltage sets an intrinsic length scale which leads to the associated scaling properties of various electrical quantities. The onset exponents \( x_T \) are all positive and different in different systems indicating that the microscopic details play a significant role in setting the non-Ohmic conduction in these nanostructures. The phenomenology of scaling, existence of a single voltage scale and the nonlinearity exponent \( x_T \) need to be supported by proper theoretical understanding.

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Table 1

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<th>Systems</th>
<th>P</th>
<th>( x_T )</th>
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<tr>
<td>( \text{K}_0.27\text{MnO}_2) 0.5H(_2)O nanowire of diameter 148 nm</td>
<td>T</td>
<td>3.98 ± 0.104</td>
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<tr>
<td>PEDOT nanowire of diameter 95 nm</td>
<td>T</td>
<td>2.94 ± 0.18</td>
</tr>
<tr>
<td>PEDOT nanowire of diameter 92 nm</td>
<td>T</td>
<td>2.85 ± 0.096</td>
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<tr>
<td>PPy nanotubes of diameter 160 nm</td>
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<td>0.88 ± 0.04</td>
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<tr>
<td>PPy-HQSA nanotube</td>
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<tr>
<td>PDA single crystal [34]</td>
<td>T</td>
<td>0.51 ± 0.02</td>
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<tr>
<td>Polyaniline nanofibre [34]</td>
<td>T</td>
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