The Relationship Between Material Properties and Transparent Heater Performance for Both Bulk-Like and Percolative Nanostructured Networks

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ABSTRACT: Transparent heaters are important for many applications and in the future are likely to be fabricated from thin, conducting, nanostructured networks. However, the electrical properties of such networks are almost always controlled by percolative effects. The impact of percolation on heating effects has not been considered and the material parameter combinations which lead to efficient performance are not known. In fact, figures of merit for transparent heaters have not been elucidated, either in bulk-like or percolative systems. Here, we develop a simple yet comprehensive model describing the operation of transparent heaters. By considering the balance of Joule heating versus power dissipated by both convection and radiation, we derive an expression for the time-dependent heater temperature as a function of both electrical and thermal parameters. This equation can be modified to describe the relationship between temperature, optical transmittance and electrical/thermal parameters in both bulk-like and percolative systems. By performing experiments on silver nanowire networks, systems known to display both bulk-like and percolative regimes, we show the model to describe real systems extremely well. This work shows the performance of transparent heaters in the percolative regime to be significantly less efficient compared to the bulk-like regime, implying the diameter of the nanowires making up the network to be critical. The model allows the identification of figures of merit for networks in both bulk-like and percolative regimes. We show that metallic nanowire networks are most promising, closely followed by CVD graphene with networks of solutionprocessed graphene and carbon nanotubes being much less efficient.

Keywords: percolation, radiative, convective, thermal, graphene, nanotubes

The past few years have seen a considerable amount of research devoted to the study of nanostructured transparent conducting materials. The aim of this research is to use nanomaterials to replace traditional transparent conductors such as indium tin oxide (ITO).¹ While transparent conducting oxides have provided good service for many years² they face a number of difficulties which make them unsuitable for next-generation applications. For example, the rising price of indium has made ITO increasingly expensive. In addition, all transparent metal oxides are brittle³ and expensive to deposit over large areas. This makes them inappropriate for many future uses given the anticipated shift to large area, flexible display technology.

A number of different nanomaterials have been extensively tested in this space. Probably the most studied are solution processed networks of carbon nanotubes,⁴⁻⁶ graphene nanosheets^{7, 8} and metallic nanowires (NWs)⁹⁻¹⁷ as well as vapour grown graphene films.¹⁸ These materials have been used as electrodes in a range of applications including light emitting diodes, solar cells and transparent capacitors.^{19, 20} However, more recently, attention has turned to using nanostructured transparent conductors as transparent heaters.²¹⁻³⁰ Transparent heaters are simply conducting films which are thin enough to be transparent but can be heated up on application of a voltage. For a given combination of electrical and thermal properties the steady state temperature increase is set by balance of Joule heating and heat dissipation and can be controlled *via* the voltage. Such devices are important for a number of applications from defogging of windows or mirrors²¹ to performance optimisation of liquid crystalline displays *via* temperature control³¹ to art conservation.³²

One of the most commonly used, commercially available transparent heater materials is ITO. In fact, one of the earliest patents describing transparent, conducting doped tin oxide films explicitly described de-icing aircraft windscreens as a potential application.³³ However, ITO transparent heaters suffer the same problems that ITO faces in other transparent conducting applications, namely cost, flexibility and areal scaling. As with other transparent conducting applications, a number of researchers have turned to nanostructured materials, particularly carbon nanotubes,^{23, 24, 26, 28, 29} graphene,^{18, 25, 34} silver nanowires^{22, 27, 30, 35} and hybrid systems.^{36, 37} The results have been very promising with reported temperature increases of up to 140 K for 4W input power.²⁹

However, there is a considerable amount of work required before it becomes clear what the true capabilities of the various nanomaterials are. In addition, it is very difficult to compare the performance of the various materials studied. For example, no well-defined figure of merit (FoM) is generally reported to facilitate benchmarking and performance comparison. Part of the problem is the lack of a comprehensive theoretical framework to analyse transparent heater behaviour. Only one paper discusses the power balance in any detail, driving an approximate relationship between temperature increase and time, current, voltage and heat transfer parameters (radiative losses are neglected).²¹ No papers discuss the relationship between temperature increase, current, voltage and heat transfer parameters with transmittance, which we view as critically important. Theoretical understanding of such relationships would not only facilitate understanding of the heating mechanisms but would allow the definition of a FoM which would allow ranking of materials and selection of most promising materials for further study.

Here we address these problems by developing a comprehensive yet simple theoretical framework which describes the relationships between temperature increase, transparency and both electrical and thermal properties. This framework can be applied to standard transparent heater materials such as ITO in a straightforward way by considering standard relationships between transmittance and sheet resistance. However, parallel experimental studies using networks of AgNWs show that such an approach can only be applied to nanostructured networks which are relatively thick. In thinner networks, connectivity effects become important, and the electrical properties become limited by percolation theory. Here we have developed an additional theoretical framework which is appropriate to the percolation regime. We show that experimental results for the steady state temperature increase for AgNW networks of different transparencies clearly display two regimes which are perfectly described by the normal (i.e. bulk) and percolative theoretical frameworks. Access to theoretical models allows us to define FoMs appropriate to both for normal and percolative transparent heaters. By analysing literature results, we can show that AgNW networks, closely followed by CVD graphene films, are by far the best performing nanostructured transparent heaters.

RESULTS AND DISCUSSION

The aim of this work is to develop an understanding into the factors which limit the performance of transparent heaters. We will do this by developing a simple but comprehensive model describing the dependence of network temperature on current, voltage, sheet resistance and transmittance both in the steady state and time-dependent regimes. This model will be applied to networks whose electrical properties are bulk-like and those limited

by percolative effects. We will test the validity of this model by comparison with experiments.

The model systems we will use in this work are networks of silver nanowires. These systems have been well studied by a range of authors and are reasonably well understood.^{9, 11-} ^{14, 38-42} Before developing a mathematical model to describe transparent heaters, we will fully characterise the opto-electric properties of spray deposited AgNW networks allowing them to be used as an appropriate model system. We use spray-casting¹⁴ to deposit AgNW networks of various thickness (i.e. various nanowire densities) onto PET substrates (substrate thickness ~135 μ m). From the amount of AgNW deposited, we roughly estimate the nanowire densities to range from ~10 to 150 mg/m² giving average thicknesses of ~15 to 300 nm (assuming a porosity of ~95%).⁹ This procedure results in films of various transparencies which appear uniform to the naked eye. Examples of networks with transparencies, T_R, of 97% and 57% are shown in figure 1 A and B respectively. Closer examination using SEM or He ion microscopy (figure 1 C-E) show such networks to consist of arrays of nanowires which are randomly arranged in the plane of the network. We measured the transmittance and sheet resistance of a wide range of networks as shown in figure 1F. The thickest networks had (Rs, T_R) combinations of (8.2 Ω/\Box , 57%). These values increased smoothly to (2×10⁷ Ω/\Box , 98.5%) for the thinnest networks. As a benchmark, we note that the network with transmittance of 90.5% had a sheet resistance of 53 Ω/\Box . This compares reasonably favourably to the literature for metal nanowire networks⁴³ although a number of papers have reported lower sheet resistances for $T_R \approx 90\%$.³⁸ However, the results presented here are far superior to reported values for transparent conductors of almost all other nanomaterials.⁴³

For thin conducting films, the optical transmittance (at a given wavelength) can be related to the sheet resistance, R_s , by^{4, 42, 44}

$$T_{R} = \left(1 + \frac{Z_{0}}{2R_{s}} \frac{\sigma_{Op}}{\sigma_{DC,B}}\right)^{-2}$$
(1)

where Z_0 is the impedance of free space (377 Ω). (We note that for films of nanostructured objects such as nanowires this expression holds so long as the film thickness >2.33 D, where D is the nanowire diameter, see below⁴²). Here the ratio of bulk D.C. to optical conductivity, $\sigma_{DC,B}/\sigma_{Op}$ can be considered a figure of merit with high values giving the desired properties (high T_R coupled with low R_s). It has been pointed out that equation 1 is not strictly applicable

for many nanostructured films because it assumes the interaction of light solely with free carriers which is not strictly true in the visible region.⁴⁵ However, this expression generally describes experimental data for relatively thick films rather well and has the advantage that values of $\sigma_{DC,B}/\sigma_{Op}$ are known for a range of nanostructured thin films.⁴³ In addition, because the optical conductivity in this expression can be shown to be proportional to the Lambert-Beer absorption coefficient, α ,⁴² we feel is it acceptable to use once $\sigma_{DC,B}/\sigma_{Op}$ is treated as a figure of merit rather than a physical property. It is worth noting that the entire analysis described below can be performed equally well using an expression which is analogous to equation 1 but is based on the Lambert-Beer law (*i.e.* $T = e^{-\alpha/\sigma_{DC,B}R_s}$).⁴²

We can test applicability of equation 1 to our data by plotting $T_R^{-1/2} - 1$ versus R_s/Z_0 (figure 1G). Here a straight line on a log –log plot with slope of -1 is characteristic of bulk behaviour. This is indeed the case, allowing us to obtain $\sigma_{DC,B}/\sigma_{Op}=70$ (using this value, we have plotted equation 1 as a dotted line on figure 1G for comparison). This value is smaller than values reported for other metallic nanowire networks(83 $<\sigma_{DC,B}/\sigma_{Op}<453$, see a recent review for tabulated data⁴³), probably due to source to source variations in the nanowires (or perhaps the organic stabilising coating).

However, equation 1 only fits the data for networks with $R_s/Z_0<0.12$, with the data diverging for thinner networks. This is relatively common phenomenon^{9, 46-48} and has been attributed to percolation effects.⁴² For such thin networks, a new relationship between T_R and R_s has been proposed:⁴²

$$T_{R} = \left[1 + \frac{1}{\Pi} \left(\frac{Z_{0}}{R_{s}}\right)^{1/(n+1)}\right]^{-2}$$
(2)

Where n is the percolation exponent and Π is known as the percolative figure of merit:

$$\Pi = 2 \left[\frac{\sigma_{DC,B} / \sigma_{Op}}{\left(Z_0 t_{\min} \sigma_{Op} \right)^n} \right]^{1/(n+1)}$$
(3)

Here, t_{min} is the transition thickness, below which the DC conductivity becomes thickness dependent (*i.e.* equation 2 applies for t $< t_{min}$, while equation 1 applied for t $> t_{min}$). Analysis of these equations shows that large values of Π coupled with low values of *n* are desirable to achieve low R_s and high T_R .⁴² Furthermore, we showed empirically that networks of nanowires have values of t_{min} which scale closely with the wire diameter, *D*: $t_{min} \approx 2.33D$.⁴² In fact equation 2 fits the high R_S/Z₀ data in figure 1F very well, giving fit values of Π =26 and n=5.6 (again, these values have been used to plot equation 2 on figure 1G for comparison). This value of Π is somewhat below the median value of 31.7 for metallic nanowire networks, probably due to the low value of $\sigma_{DC,B}/\sigma_{Op}$.⁴³ In addition, the percolation exponent is higher than typical values found for metallic nanowire networks,⁴³ suggesting the networks to be somewhat disordered.¹⁴

The data in figure 1 shows that the AgNW networks studied here both resemble and behave similarly to networks previously described in a number of papers. Critically, the optical and electrical properties of thick networks behave as expected for bulk-like films while thinner networks are described by percolation theory. In fact it is known that such percolative behaviour is almost always found for nanostructured transparent conductors.⁴³ In addition, the technologically relevant regime around $T_R=90$ almost always falls in the percolative regime, at least for solution processed networks.⁴³ This is important as it means that any comprehensive understanding of nanostructured heaters will have to incorporate percolation theory in some form.

Heating behaviour – time dependence

In order to assess the performance of these networks as transparent heaters, we drove a fixed current through the networks (inter-electrode separation, l=2 cm, electrode width, w=2 cm), measuring the surface temperature as a function of time. This was carried out for a number of networks of different thicknesses (and so different transmittances and sheet resistances) using a range of current values. In all cases the data were perfectly reproducible with no irreversible temperature effects observed. Examples of the resultant temperature *versus* time data are shown in figure 2A for a network with T_R=61% for a number of different applied currents. In all cases the temperature increased monotonically with time before eventually saturating. The saturation temperature depended on both the applied current and the network thickness (*i.e.* transmittance).

To quantitatively analyse this data it is necessary to develop a model which relates the time evolution of the temperature to the applied current and a parameter representing the network thickness *e.g.* the transmittance or the sheet resistance. We note that elements of such a model have been presented by Bae *et al.*²¹ However, a full description such as that described here has not been reported. We can develop such a model by considering the

energy balance between heating and dissipation. During current flow, the power dissipated by Joule heating in the AgNW network is given by $P_{in} = I^2 R$ where R is the network resistance. Some of the dissipated power goes to increasing the temperature, T, of both the nanowire network and (*via* conduction) the substrate while the remainder is lost *via* radiation and convection at both the nanowire network surface and the opposite surface of the substrate. Then, making the approximation that the instantaneous temperature is the same everywhere in both the network and the substrate, we can write a power balance equation:

$$I^{2}R = (m_{1}C_{1} + m_{2}C_{2})\frac{dT(t)}{dt} + A(h_{1} + h_{2})(T(t) - T_{0}) + \sigma A(\varepsilon_{1} + \varepsilon_{2})(T(t)^{4} - T_{0}^{4})$$
(4)

Here T(t) and T_0 are the instantaneous sample temperature and the ambient temperature, A is the area of the film (assumed equal to the substrate area) and σ is the Stefan Boltzmann constant. The subscripts 1 and 2 refer to the network and the substrate respectively such that m_1 and m_2 are the masses and C_1 and C_2 are the specific heat capacities of network and substrate. Similarly, h_1 and h_2 are the convective heat-transfer coefficients and ε_1 and ε_2 are the emissivities of the sides of the sample associated with the network and substrate respectively. The term on the left of equation 4 is the dissipated electrical power while the first term on the right describes the portion of that power used to raise the temperature of both network and substrate. The second term on the right is an approximate representation of the energy lost by convection while the third term on the right represents the net energy lost radiatively (taking into account the thermal radiation absorbed from the environment).

Unfortunately no simple analytical solution exists for this differential equation. However, we can simplify it somewhat by noting that for small temperature rises (*i.e.* $T(t) - T_0 < 40K$) we can apply a Taylor expansion to give $T(t)^4 - T_0^4 \approx 4T_0^3(T(t) - T_0)$. This allows us to approximate the energy balance expression as $(m_1C_1 + m_2C_2)\frac{dT(t)}{dt} + A[(h_1 + h_2) + 4T_0^3\sigma(\varepsilon_1 + \varepsilon_2)](T(t) - T_0) - I^2R \approx 0$ (5)

This equation can be solved analytically to give

$$T(t) \approx T_0 + \frac{1}{\alpha} \frac{I^2 R}{A} \left[1 - exp\left(-\frac{\alpha}{C_2 m_2 / A} t \right) \right]$$
(6)

where $I^2 R / A$ is the areal power density. We note that the network is much less massive than the substrate leading to the approximation: $m_1 C_1 \ll m_2 C_2$. In addition, we introduce the symbol α for the quantity

$$\alpha = (h_1 + h_2) + 4(\varepsilon_1 + \varepsilon_2)\sigma T_0^3 \tag{7a}$$

where we refer to α as the heat transfer constant.

We have fitted equation 6 to the experimental data for T(t). We found very good fits in all cases as illustrated in figure 2A. From the fits we can extract the time constant $\tau = m_2 C_2 / (A\alpha)$ which we plot *versus* network transmittance (*i.e.* a measure of network thickness or density) in figure 2B. While the data is somewhat scattered, all values cluster between 60 and 100 s, of the same order of magnitude as time constants reported by other researchers.^{21, 22, 25-27, 29, 30, 36, 37}

From the fits, we can independently extract α . This is plotted *versus* the network transmittance in figure 2C. This shows the heat transfer constant to lay in the range 25-50 Wm⁻²K⁻¹ depending on the network thickness. While the emissivity of the AgNW network is not known, the emissivity of PET is known to be reasonably high; $\varepsilon_2 \approx 0.9$. This means the radiative contribution to α from the PET substrate side of the system is $\approx 4\varepsilon_2 \sigma T_0^3 \approx 6$ Wm⁻ 2 K⁻¹. Roughly extrapolating the experimental data to T_R=100% suggests that for PET alone, α ~40 Wm⁻²K⁻¹ (for both sides). Combining this with the radiatiative heat loss from PET implies that the convective heat-transfer coefficient of PET is $h_2 \sim 14 \text{ Wm}^{-2}\text{K}^{-1}$, close to that of glass (hglass~10 Wm⁻²K⁻¹).²¹ Thus, depending on the network thickness, heat loss from the AgNW-coated surface contributes ~5-30 Wm⁻²K⁻¹ to the observed value of alpha (with the PET substrate side contributing ~20 Wm⁻²K⁻¹). The maximum radiative contribution to α from the AgNW-coated surface is $4\sigma T_0^3 \approx 6$ (*i.e.* if $\varepsilon_1=1$). This means that convective heat loss from the AgNW-coated surface is likely to dominate the heat loss process with values of $h_1 \sim 0.24 \text{ Wm}^{-2} \text{K}^{-1}$ depending on the network thickness. This is consistent with values of h=8-92 Wm⁻²K⁻¹ previously observed for nanostructured or metallic films.²¹ In addition, it confirms that both convection and radiation are significant heat loss pathways. This is different to graphene films where heat loss is almost completely due to convection because of the low emissivity of graphene.²¹

We can begin to understand the dependence of α on transmittance by noting that heat convection and radiation are interfacial phenomena. On the network side of the sample, for very low nanowire coverage, the solid-air interface has contributions from both PET and AgNWs. Under these circumstances, h₁ and α_1 will depend on the convective heat-transfer coefficients and emissivities of both PET and nanowires as well as the area fraction of surface coated with nanowires, f_{NW}. At low coverage and so high transmittance, we can make the approximations that $h_1 = f_{NW}h_{NW} + (1 - f_{NW})h_{PET}$ and $\varepsilon_1 = f_{NW}\varepsilon_{NW} + (1 - f_{NW})\varepsilon_{PET}$ while of course $h_2 = h_{PET}$ and $\varepsilon_2 = \varepsilon_{PET}$. Using this approximation, equation 7a becomes

$$\alpha = 2(h_{PET} + 4\varepsilon_{PET}\sigma T_0^3) - f_{NW} \left[(h_{PET} - h_{NW}) + 4\sigma T_0^3 (\varepsilon_{PET} - \varepsilon_{NW}) \right]$$
(7b)

We assume for simplicity that we can relate the transmittance to the amount of nanowires coating the substrate using the Lambert-Beer law: $f_{NW} \propto -\log T_R$ allowing us to write $\alpha = K_1 + K_2 \log T_R$. This functional form has been plotted on figure 2C and is consistent with the high T_R data so long as K₂>0.

While values of h_{NW} and ε_{NW} are not known, AgNW networks are known to exhibit thermal shielding behaviour,⁴⁹ leading to low values of ε_{NW} compared to other materials, suggesting ($\varepsilon_{PET} - \varepsilon_{NW}$) would be positive. Then, that K₂>0 implies that ($h_{PET} - h_{NW}$)>0, at least at low coverage. However, the behaviour described by equation 7b breaks down for T_R<80% (figure 2C). However, this is not surprising as figure 1E shows f_{NW} to be quite high at this transmittance, probably invalidating the assumptions leading to equation 7b for low T_R values.

As T_R falls below 80%, α begins to increase with decreasing T_R (*i.e.* with increasing nanowire coverage). However, in this regime, h_{NW} is probably controlled by network properties such as surface roughness⁵⁰ and high internal surface area. This may result in an increase in h_{NW} with increasing coverage, leading to the observed behaviour

Heating behaviour – steady state

The steady state temperature, T_{Sat} , can be found from equation 5 by setting dT/dt=0. This gives:

$$T_{sat} = T_0 + \frac{I^2 R}{\alpha A} = T_0 + \frac{I^2 R_s}{\alpha w^2}$$
(8)

where we have used $R = R_s l/w$ and A = lw. To test this, we applied a range of applied currents to a number of networks of various thicknesses. We measured the steady state temperature after ~10 minutes when it had clearly saturated. Shown in figure 3A is a graph of the steady state temperature increase, $\Delta T = T_{sat} - T_0$, plotted *versus* the applied current for a number of different networks. It is clear from this data that $\Delta T \propto I^2$ for all networks studied. We have plotted the temperature rise *versus* the areal power density (*i.e.* the power inputted by Joule heating per unit area) in figure 3B. To a first approximation it is clear that all samples fall roughly on the same mastercurve such that $\Delta T \propto I^2 R/A$. However, a closer look shows some deviation from a single mastercurve. These deviations are consistent with variations in α from sample to sample. From the fit curves shown on figure 3A we can calculate α using equation 8. We have plotted α *versus* the network transmittance, T_R, in figure 3C. We find behaviour very similar to that found by fitting the time dependent data (figure 2C). This shows that the model fits both time dependent and steady state data extremely well.

We note that equation 8 predicts ΔT to scale linearly with power (I^2R in our notation). Such behaviour has been reported by a number of other authors for nanostructured transparent heaters.^{21-24, 26, 28, 29} This allows us to use equation 8 to analyse previously reported data to extract α . The results are given in the SI and show values of α in the range 15-123 Wm⁻²K⁻¹. Averaging the results by material gave mean values of 21 Wm⁻²K⁻¹ (graphene), 70 Wm⁻²K⁻¹ (nanotubes) and 39 Wm⁻²K⁻¹ (metallic nanowires).

Relating temperature to transmittance

Once we know that the simple model outlined above describes the data reasonably well, we can extend it to describe the relationship between temperature increase and transmittance. To do this for bulk-like networks we simply rearrange equation 1 for R_s and substitute into equation 8 to give

$$\frac{\Delta T}{I^2} = \frac{Z_0}{2\alpha w^2} \frac{\sigma_{OP}}{\sigma_{DC,B}} \left(\frac{1}{\sqrt{T_R}} - 1\right)^{-1}$$
(9)

We can apply the same procedure to describe networks in the percolative regime, except this time using equation 2 instead of equation 1:

$$\frac{\Delta T}{I^2} = \frac{1}{\Pi^{n+1}} \frac{Z_0}{\alpha w^2} \left(\frac{1}{\sqrt{T_R}} - 1 \right)^{-(n+1)}$$
(10)

These equations imply that $\Delta T / I^2$ should scale with $T_R^{-1/2} - 1$ as a power law with an exponent that reflects whether the networks are in the bulk-like regime (*i.e.* thicker and exponent=-1) or the percolative regime (*i.e.* thinner and exponent=-(n+1)). To test this, we plotted $\Delta T / I^2$ versus $T_R^{-1/2} - 1$ for all samples (*i.e.* different thicknesses and currents) in figure 4A on a log-log plot. We do indeed find two separate regions described by different power laws. To demonstrate consistency with our model, we plot the curves described by equations 9 and 10 alongside the data in figure 4A using the parameters given above (*i.e.* w=2 cm, $\sigma_{DC,B} / \sigma_{Op} = 70$, $\Pi = 26$ and n=5.6). Because α is weakly thickness dependent, we use a representative value: $\alpha = 40$ Wm⁻²K⁻¹. The resultant curves overlay the data extremely well.

Figure 4A clearly illustrates the fact that the relationship between temperature and transmittance in AgNW networks differs between percolative and bulk regimes. Because Joule heating increases with the electrical resistance of the network, it is not surprising that $\Delta T/I^2$ values are higher for the less dense, more transparent networks. Therefore, at first glance the percolative regime appears most suitable for applications. This apparent supremacy of more resistive networks seems to be reinforced by comparison with literature data for transparent heaters fabricated from SWNTs and AgNWs (figure 4A). The AgNW data lies slightly below the data generated here while data for the more resistive SWNT networks give a larger temperature rise per unit current compared to AgNW networks. However, closer examination shows this to be very misleading: more resistive networks are not better transparent heaters. The reason is that more resistive networks require more power to drive a given current and so reach a given temperature increase. This makes them less efficient overall. We perform quantitative analysis to demonstrate this below.

Operating voltage and figures of merit

The analysis above shows that the model we have described fits real data extremely well. This allows us to consider what properties are required of a network to work in a real transparent heater application. It is likely that the details of the application will set ΔT and T_R

as well as the heater dimensions *i.e.* 1 and w. Then, ΔT will define the required power *via* equation 8 while T_R will define R_s *via* equation 1 or 2, depending on whether the network is bulk-like or percolative. Then the aim will be to achieve the required ΔT given the set value of T_R (and so R_s) for the lowest applied voltage, V. This will then minimise the power because $P = I^2 R = V^2 / R$. Then, using equation 8 and changing variable from current to voltage using $V = IR_s l / w$ gives an expression for the operating voltage:

$$V = \sqrt{\alpha \Delta T R_s l^2} \tag{11}$$

If, for example, the network is such that the required transmittance occurs in the bulk-like regime, then we can use equation 1 to replace R_s to give:

$$V = \sqrt{\frac{\alpha \Delta T Z_0 l^2}{2(T_R^{-1/2} - 1)\sigma_{DC,B} / \sigma_{Op}}}$$
(12)

This expression clearly shows that to minimise V, we need a network material with a low α and high $\sigma_{DC,B}/\sigma_{Op}$. Because this equation describes the bulk-like regime, it is only appropriate for networks with relatively low transmittances. The transmittance where a network changes from bulk-like to percolative depends on the specific nanomaterial being used. However, De el al. have shown that most networks are bulk-like for T_R=75%.⁴³ Thus, for descriptive purposes, we use equation 12 to find the voltage required to induce a temperature increase of 40 K in a network (T_R=75%) fabricated from the wires described here to be V=17 V (using l=0.1 m, α =40 Wm⁻²K⁻¹ and $\sigma_{DC,B}/\sigma_{Op}=70$).

For comparison purposes, we can calculate this operating voltage for transparent heaters fabricated from networks (T_R=75%) of a number of different nanostructured transparent conductors that have been described in the literature. To do this, we use the values of α extracted from the literature as described above. In addition, we make use of a recent review which has tabulated values of $\sigma_{DC,B} / \sigma_{Op}$ for solution processed networks of graphene, nanotubes and metallic nanowires⁴³. We use these tabulated values of $\sigma_{DC,B} / \sigma_{Op}$ coupled with the values of α reported above to predict the voltage, V, required to achieve a steady state temperature rise of 40 K for these previously reported networks of metallic nanowires, SWNTs and graphene (all solution-processed, and taking l=10 cm, figure 4B). The data in figure 4B shows that metallic nanowire networks require values of V<16 V

(median 10 V). However, solution processed SWNT and Graphene networks require voltages in the range 23-730 V (median 85 V) and 25-4500 V (median 172 V) respectively.

However, it is more likely⁴³ that the required transmittance will be considerably above 70% (usually $T_R \sim 85-95\%$) and so will occur in the percolative regime for solution processed networks.⁴² Then we can use equation 2 to replace R_s in equation 8 (and using $V = IR_s l / w$) to give:

$$V = \sqrt{\frac{\alpha \Delta T Z_0 l^2}{\left[(T_R^{-1/2} - 1)\Pi \right]^{(n+1)}}}$$
(13)

Applying this to our AgNW networks shows that V=283 V is required to raise the temperature by 40 K for a percolative network which is 95% transparent (l=10 cm, α =25 Wm⁻²K⁻¹, Π =26, n=5.6). This value is very high, clearly showing the problems associated with very thin nanostructured networks. Using literature data for Π and n tabulated by De *et al*,⁴³ and the values of α given above, we can calculate the voltage required for a 40 K rise for transparent heaters prepared from very thin networks of metallic nanowires, SWNTs and graphene (T_R=95%, l=10 cm, figure 4C). These are plotted as a function of Π in figure 4C which shows that metallic nanowire networks require values of V as low as 60 V. However, SWNT and Graphene networks require voltages which are >300 V and >500 V respectively. These results show that for solution processed networks, metallic nanowires are far superior to networks of nanotubes or graphene.

However, it is also very important to note the effect of percolation on nanowire performance. In general, increasing the transmittance of a heater from 75% to 95% involves moving from a network which is bulk-like to one which is percolative. According to the data in figure 4 B&C, this will result in an order of magnitude increase in operating voltage for networks of metallic nanowires. This will result in massive efficiency reduction and is obviously prohibitive. It is clear that networks which can retain bulk-like behaviour at low enough thickness such that T_R ~90-95% will be important for transparent heaters. Because the thickness (and so transmittance) defining the bulk to percolative transition depends on the diameter/thickness of the nanowires/nanosheets making up the network,⁴² thinner nanostructures (*i.e.* lower diameter nanowires/nanotubes or thinner nanosheets) will result in higher efficiency.

It is clear from equation 12 that, for bulk-like networks, the material parameters that control the performance of thermal heaters are α and $\sigma_{DC,B} / \sigma_{Op}$. For low operating voltages, low values of α coupled with high values of $\sigma_{DC,B} / \sigma_{Op}$ will be required. For percolative networks, the equivalent parameters would be α and Π^{n+1} . We note that with the exception of one paper²¹ none of the published work on transparent heaters give values of α , $\sigma_{DC,B} / \sigma_{Op}$, n or Π . In addition, none of these papers give enough information to ascertain whether they are bulk-like or percolative (there are two papers on CVD grown graphene which is clearly not a percolative material^{21, 25}). However, for simplicity we assume all are bulk-like and, where possible, we extracted values of α and $\sigma_{DC,B} / \sigma_{Op}$ from the reported data (see SI). We did this for three papers based on CNTs,^{24, 26, 29} two papers based on CVD graphene,^{21, 25} one paper on solution processed graphene (SP Gra),³⁴ two papers based on AgNW networks^{27, 30} and two papers based on hybrid structures.^{36, 37} We plot α *versus* $\sigma_{DC,B} / \sigma_{Op}$ in figure 4 D, including the results from this paper for comparison. Given that small α and large $\sigma_{DC,B} / \sigma_{Op}$ are required for effective performance, it is clear that CVD graphene and AgNW networks are far superior to CNT or solution processed graphene transparent heaters.

We can see this in another way by noting that inspection of equation 12 shows that $(\sigma_{DC,B}/\sigma_{Op})/\alpha$ can be used as a figure of merit for bulk-like transparent heaters (higher values give better performance). We have used the data in figure 4D to plot a bar chart of $(\sigma_{DC,B}/\sigma_{Op})/\alpha$ in figure 4E. This clearly shows that AgNW networks appear to have the most promise, followed closely by CVD graphene with CNT networks falling far behind. Similar considerations would suggest Π^{n+1}/α as a figure of merit for percolative transparent heaters. Because of the relationship between Π and $\sigma_{DC,B}/\sigma_{Op}$ (equation 3), we expect the materials ranking for percolative networks to be the same as for bulk-like networks.

CONCLUSION

In conclusion, we have prepared transparent conductors from networks of silver nanowires and shown them to work effectively as transparent heaters. By considering the balance of Joule heating and energy dissipation by both radiation and convection, we have developed a comprehensive model relating the heater temperature as a function of time to electrical and thermal parameters. This model described the temperature very well in both time-dependent and steady-state regimes. Like most nanostructured systems, these AgNW networks have electrical properties that are bulk-like for thick networks but percolative for thin networks. By combining the model described above with equations relating optical transmittance to network sheet resistance in both bulk-like and percolative regimes, it is possible to generate expressions relating the steady state temperature to transmittance and current. These expressions predict significantly different heating behaviour in the bulk-like and percolative regimes. This prediction is borne out by the data with theory and experiment matching extremely well.

A good transparent heater is one that achieves a given temperature rise at as low a voltage as possible. With this in mind, the models described above can be used to suggest figures of merit for both bulk-like and percolative networks: $(\sigma_{DC,B} / \sigma_{Op}) / \alpha$ and Π^{n+1} / α respectively. High values of these parameters will lead to low operating voltages. This work suggests AgNW networks to be most promising, followed by CVD graphene, followed by solution processed nanotube and graphene networks. This ranking should apply in both bulk-like and percolative regimes.

This work provides the first comprehensive, integrated description of the physics of nanostructured transparent heaters. It clearly demonstrates the parameters which are important for effective and efficient heater operation and allows the identification of materials which can fulfil the resultant criteria. We believe this information will be very useful to the development of nanostructured transparent heaters for real applications.

Methods

For this study, silver nanowires (AgNWs) were synthesized by Kechuang (http://www.ke-chuang.com/) and supplied as a suspension in isopropyl alcohol (IPA) (C_{AgNW} = 16 mg.ml⁻¹). These nanowires had mean length of ~5 µm and mean diameter of ~50 nm. A small volume of the dispersion was diluted to 1.5 mg.ml⁻¹ in IPA and subjected to 30 sec low power sonication in a sonic bath (Model Ney Ultrasonic) to eliminate bundles of nanowires. This solution was then further diluted to 0.15 mg.ml⁻¹ and sonicated another 30 sec immediately before being sprayed¹⁴ onto polyethylene terephthalate (PET) squares of 2×2 cm and thickness 135 µm on a hotplate at 120°C. The temperature was kept high during spraying in order to evaporate the IPA swiftly and remove polymer residue left over from synthesis.

Optical transmission spectra were recorded using a Cary Varian 6000i, with a sheet of PET used as the reference. Sheet resistance measurements were made using the four probe technique using a Keithley 2400 source meter. Scanning electron microscopy and helium ion microscopy images were taken using a Zeiss Ultra scanning electron microscope and Carl Zeiss Orion PLUS Helium Ion microscope respectively.

For temperature measurements, AgNW films on PET were used. To measure the temperature, a low mass thermistor was used. This was housed inside a hole in the side of a U-shaped copper clamp (secured using nail polish) and was held in place at the midpoint between the silver electrodes (at the edge of the sample) using a small screw. Electrical measurements were made using a Keithley source meter. For a given measurement, at time t = 0, a pre-defined current was driven through the AgNW film, resulting in an increase in the film temperature and so a change in thermistor resistance. The change in resistance was recorded using MATLAB and the temperature was extracted using the thermistor's calibration curve. The temperature was recorded as a function of time over the course of 15 minutes, significantly longer that the necessary time for the temperature to reach steady-state. Stabilising the experimental environment was vital since the lab temperature could fluctuate by up to 3 degrees centigrade. The system was sheltered from the ambient lab using a copper bell jar covered with commercially available insulating foam.

Supporting Information Available: Analysis of literature data for nanostructured transparent heaters. This material is available free of charge *via* the Internet at http://pubs.acs.org.

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Figures

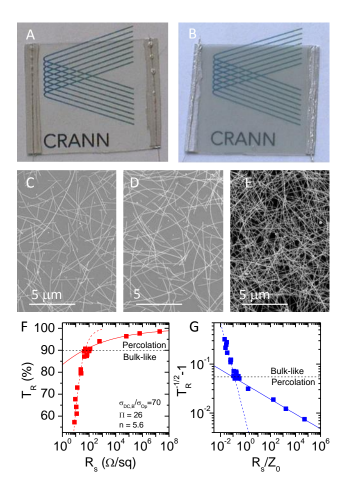


Figure 1: A-B) Photographs of spray coated AgNW networks with optical transmittance of A) 97% and B) 57%. C-D) SEM images of AgNW networks with transmittance of C) T_R =94% and D) T_R =90%. E) He ion micrograph of an AgNW network with T_R =80%. F) Optical transmittance (550 nm) plotted *versus* sheet resistance for all the networks prepared in this study. The red lines are fits to equation 1 (dashed) and equation 2 (solid). The fit constants are given in the panel. The black horizontal line indicates the boundary between bulk-like and percolative behaviour. G) The same data in F, plotted to illustrate linear behaviour.

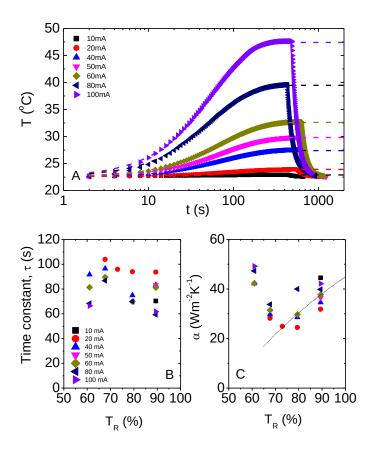


Figure 2: A) Time dependence of temperature rise for an AgNW network ($T_R=61\%$) with a number of different applied currents. The dashed lines represent fits to equation 6. B-C) Data derived from fitting time dependent data such as that in A. B) The time constant and C) heat transfer constant, α , for AgNW networks as a function of network transmittance. In each case, data is shown for a range of applied currents. In C the dashed line represents the behaviour suggested by equation 7b.

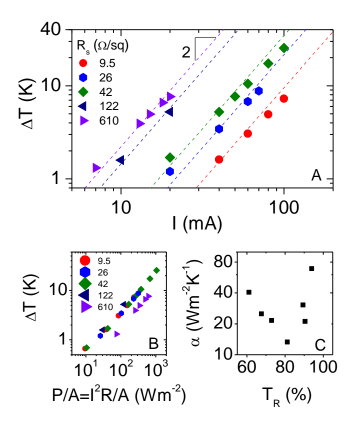


Figure 3: A) Steady state temperature, ΔT , rise plotted as a function of current, I, for networks with different sheet resistance, R_s (*i.e.* different thicknesses and so transmittances). The dashed lines represent $\Delta T \propto I^2$. B) Temperature rise plotted as a function of areal power density. The labels give the film sheet resistance. C) The heat transfer constant, α , calculated from the slope of the curves in A and plotted *versus* film transmittance. In all cases the interelectrode separation, 1=2 cm and the electrode width, w=2 cm. In A and B only some of the data sets are shown to avoid clutter.

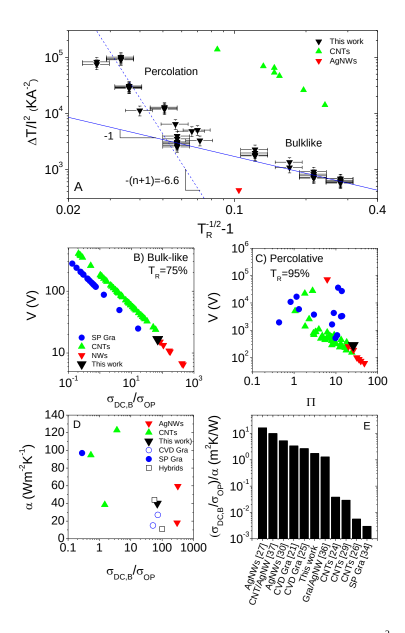


Figure 4: Temperature rise over current squared $(\Delta T/I^2)$ plotted *versus* $T_R^{-1/2} - 1$ for all networks at all applied currents. Note the latter parameter is proportional to network thickness. Measurements at all currents have collapsed onto the same master curve. The lines are plots of equation 9 (solid) and equation 10 (dashed), representing behaviour in the bulk-like and percolative regimes respectively. Also shown are data extracted from the literature for SWNT networks²⁹ (green) and an AgNW network²² (red). B) Predicted voltage required to reach a steady state temperature increase of 40 K as a function of $\sigma_{DC,B}/\sigma_{Op}$. The calculation assumes a bulk-like network with T_R=75%. The symbols represent reported values of $\sigma_{DC,B}/\sigma_{Op}$ for solution processed graphene (SP Gra), carbon nanotube (CNTs) and AgNW networks.⁴³ C) Plot of voltage required to reach a steady state temperature increase of 40 K as a function of percolative figure of merit, Π . The symbols represent data points

calculated using known values of Π and n for solution processed graphene, nanotube and AgNW networks.⁴³ The calculation assumes a percolative network with T_R=95%. Also included in B) and C) are values of V calculated for the networks studied in this work. In both B) and C), the calculations use the following values: l=10 cm, $\alpha/(Wm^{-2}K^{-1})=21$ (Graphene), 70 (SWNTs), 39 (NWs), 40 (This work). D) Comparison of performance of thermal heaters in literature^{21, 24-27, 29, 30, 34, 36, 37} with this work. Heat transfer constant, α , plotted *versus* $\sigma_{DC,B}/\sigma_{Op}$, for a number of solution processed (SP) Graphene, CVD Graphene, CNT, AgNW and hybrid (*i.e.* graphene AgNW and CNT/AgNW) transparent heaters. Both values were extracted by us from published data (see SI). E) Thermal heater Figure of Merit, $(\sigma_{DC,B}/\sigma_{Op})/\alpha$ calculated from data in D. The bracketed number denotes the reference.

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