Low-temperature electronic transport and the Coulomb blockade in oxidized films of bismuth

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We present low-temperature (0.2 < T < 10 K) transport measurements on semicontinuous films of bismuth which form a network of coupled Bi grains. By sequentially oxidizing such films we have observed a change in the character of transport that may be associated with the emergence of a finite charging energy and Coulomb blockade for single-electron tunneling between grains. This transition is manifested most dramatically as a divergence of the thermoelectric power with decreasing temperature. Magnetoresistance measurements indicate that the crossover in the thermopower occurs abruptly when the average oxide barrier resistance exceeds a fundamental value near $\hbar/e^2$.

Granular metal films have been the focus of considerable attention for the past decade in the study of the metal–insulator transition [1]. These systems generally consist of metallic grains separated by insulating barriers, the electronic coupling through which governs the character of the low-temperature resistance [2]. The limits of strong and weak coupling are well documented. In the former case the barriers have negligible activation energy and the low-temperature resistance exhibits small quantum corrections due to weak localization [3] and Coulomb interaction [4] effects. In the weak-coupling limit electrons are localized to individual grains and the resistance depends exponentially on temperature [5,6]. Not understood as well is the physics at the crossover between these regimes, where neither weak nor strong localization theories are appropriate and where transport properties other than the temperature dependence of resistance [7] have rarely been addressed.

In this Letter we report low-temperature measurements of resistance, magneto-resistance and thermoelectric power in semicontinuous bismuth films. The coupling between grains is controlled by sequentially oxidizing individual films. Our focus is on the crossover regime where we observe a new and striking behavior of the thermopower. We propose a possible explanation based on the Coulomb blockade in small-capacitance tunnel junctions. Magneto-resistance data suggest that a sharp charging energy onset occurs when the average intergrain tunnel resistance exceeds $\sim \hbar/e^2$.

Samples were prepared by vapor deposition of 99.9999% purity Bi onto room temperature glass substrates in an oxygen atmosphere of 50 mTorr. The films were deposited in a four-probe bridge pattern (0.8 x 10 mm²) with nominal thickness 55–70 Å as determined by a quartz crystal monitor. A 300 Å SiO₂ layer was then deposited over the conduction channel, allowing for slower, controllable oxidation. Films prepared in this fashion had post-deposition (dc) sheet resistances, $R_\square$, of < 1 kΩ. Transmission electron and scanning tunneling microscopies indicate [8,9] that the films are composed of closely packed crystalline Bi grains which form a multiply connected, filamentary structure. The grains are irregularly shaped with diameters ranging from 200 to 1000 Å. Exposing the films to air at room temperature promotes the growth of insulating Bi₂O₃ at the granular surfaces. The corresponding increases in $R_\square$ to values > 100 kΩ are associated with a reduced in-
tergrain coupling and the development of a percolative structure, as discussed in detail elsewhere [8]. Experiments were performed in a dilution refrigerator with a superconducting solenoid providing magnetic fields perpendicular to the substrates. Steady-state thermopower measurements employed two germanium resistance sensors thermally anchored to the back of the substrates. Seebeck probes were superconducting NbTi wire, soldered to the film with indium. Thermoelectric voltages were monitored on a nanovoltmeter with analog output to a chart recorder.

Fig. 1 shows the typical low-temperature resistance behavior for one of our films. For \( R_0 (4.2 \, \text{K}) < 8 \, \text{k}\Omega \) the resistance increases as \( \ln T \), with a slope that agrees with weak disorder theories for two-dimensional systems [3,4,10] and is consistent with previous studies of thin, homogeneous Bi films [11]. The temperature dependence for \( R_0 (4.2 \, \text{K}) > 8 \, \text{k}\Omega \) is faster than \( \ln T \) (approximately \( T^{1/2} \), but not yet exponential. For example, forcing a fit to the strong localization form [5,6], \( R \propto \exp \left( \frac{T_0}{T} \right)^{1/2} \), yields \( T_0 < 20 \, \text{mK} \). Such a low value for \( T_0 \) is not consistent with \( T \gg T_0 \) for which this expression applies.

The thermoelectric power, \( S \), for the same film from fig. 1 is shown in fig. 2. Similar thermopower data have been observed in three separate films, each oxidized in steps to span the range \( 1 < R_0 < 100 \, \text{k}\Omega \). For disordered thin films, phonon drag effects are rendered negligible and we interpret the data as reflecting the carrier diffusion thermopower. The negative sign indicates a dominant contribution from electrons. The thermopower of metals [12] is expected to decrease linearly with temperature, and this behavior is observed down to the lowest temperatures for \( R_0 (4.2 \, \text{K}) < 8 \, \text{k}\Omega \). For \( R_0 (4.2 \, \text{K}) > 8 \, \text{k}\Omega \) this \( T \)-linear dependence gives way to a dramatic transition where \( S \) diverges with decreasing temperature to values that are several orders of magnitude larger than those for samples having low \( R_0 \). Note that the temperature at which the thermopower diverges increases with increasing oxidation and the divergence becomes less rapid, tending toward an \( S \propto 1/T \) dependence. The latter behavior is characteristic of a thermally activated process, as in semiconductors [13] where the energy gap, \( E_g \), gives rise to a low-temperature \( (k_B T < E_g) \) thermopower, \( S \propto E_g / 2 k_B T \). Burns et al. [14] observed a similar crossover from metallic to activated \( (1/T) \) thermopower in thin, disordered Au and Au–Pd films. The surprising feature in our data is the more rapidly diverging thermopower that characterizes films with intermediate oxidation.

Developments [15,16] in the physics of small-capacitance tunnel junctions may offer insight into the
thermopower behavior of our films. Central to a semiclassical description [15] of the current and voltage response for a single junction of capacitance $C$, is the relationship between the charging energy or Coulomb blockade, $E_C = e^2/2C$, and the tunnel resistance, $R_t$. This is most clearly illustrated for a weakly coupled junction, characterized by $R_t \gg \hbar/e^2$ ($\approx 4.12 \text{ k}\Omega$). This resistance criterion [17] is equivalent to having localized electronic states on either side of the barrier. In this case the charge on the junction may be treated classically such that the capacitance, and hence $E_C$, have their usual geometric meaning. A fundamental issue is how this description should be modified when the weak coupling condition is relaxed. A theoretical treatment of this problem [18] for a single junction predicts that, as the temperature and capacitance tend to zero, an abrupt breakdown of the Coulomb barrier is possible when $R_t \sim \hbar/e^2$. This issue is addressed in a recent experimental study [19] of lithographically patterned single junctions and arrays, where experiments suggest that the relevant charging energy for transport may decrease below its geometric value for $R_t$ lower than about 10 kΩ.

For the arrangement of grains in our films we estimate an average geometric capacitance on the order of $C = 5 \times 10^{-16} \text{ F}$, which corresponds to a charging energy $E_C/k_B \approx 2 \text{ K}$. From the slope of the data for the highest level of oxidation in fig. 2 we find an activation energy of $\sim 1 \text{ K}$, which compares favorably with our estimate of $E_C$. This suggests that in the limit of high oxidation the charging energy is close to its geometric value with the thermopower taking on a simple activated form, $S \propto E_C/k_B T$. This $1/T$ behavior is expected for discontinuous metal films [20] when transport is by thermally activated tunneling between islands. We have recently calculated [21] the thermopower for a single tunnel junction, using a semiclassical approach [15]. In this model the charging energy is included as a shift in the chemical potential of the electrode into which an electron tunnels. Under certain conditions we find a thermopower that crosses over, with decreasing temperature, from a metallic ($T$-linear) to activated ($\propto E_C/k_B T$) form. A faster than $1/T$ divergence is not predicted by these calculations.

It is possible that the sharp upturn in the thermopower for intermediate oxidation is associated with the fact that the films form an array of coupled junctions rather than a single junction. During the time between tunneling events across an intergrain tunnel junction, there can be a redistribution of the charge onto neighboring grains. Such charge fluctuations will tend to increase the observed capacitance because the area probed by the electron is greater [22]. The net result will be a decrease in the charging energy, yielding an effective value, $E_C^{\text{eff}}$. This process will be determined by the mobility of electrons on the network and thus should be more pronounced when the intergrain tunnel resistance ($R_t$) is low. A similar picture might also be relevant for a single junction. As suggested in ref. [19], for low $R_t$ the tunneling rate may be sufficiently high for an electron to tunnel back and forth several times during the interval in which it contributes to transport. It is possible that the mobility which determines the charge redistribution in our films is temperature dependent. This would imply a $T$-dependent $E_C^{\text{eff}}$, which may be pertinent to the thermopower data. These ideas require further theoretical support. We now discuss magnetoresistance measurements which suggests that the average intergrain tunnel resistance in our films increases with oxidation and with decreasing temperature.

For $R_\square (4.2 \text{ K}) < 8 \text{ k}\Omega$ these films exhibit the same weak localization magnetoresistance (MR) that is predicted [23] and measured [11,9,24] for homogeneous Bi films. The phase breaking length at $T = 1 \text{ K}$ is $L_p \approx 1000 \text{ Å}$. For $H > H_c = \phi_0/4\pi L_p^2$ ($\phi_0$ is the flux quantum) and in the presence of strong spin-
orbit scattering (antilocalization) we have

$$\frac{\Delta R}{R} = \frac{R(H) - R(0)}{R(0)} = \frac{e^2}{4\pi^2\hbar} r_0 \ln H, \quad (1)$$

where $r_0$ is the sheet resistance of a homogeneous film. The MR for $R_0(4.2 \text{ K}) = 38.1 \text{ k}\Omega$, shown at several temperatures in fig. 3, has a dependence that is qualitatively similar to weak localization, behaving as $\ln H$ at high fields and $H^2$ at low fields. However, fitting to eq. (1) yields $r_0 \ll R_0$. This is to be expected since $R_0$ reflects percolation effects whereas $r_0$ is a measure of the sheet resistance on the scale of the Landau orbit radius, $L_H$. This is the characteristic diffusion length which cuts off the coherent backscattering responsible for weak localization in a field $H > H_\phi$. In the logarithmic field regime this length is smaller than the average grain diameter for our films. The following arguments lead us to conclude that $r_0$ measures the average intergrain tunnel resistance for highly oxidized films. Consider two points within a grain, one at the center and the other near the junction with a neighboring grain. The contribution at each such point to the $\ln H$ magnetoresistance is determined by summing the backscattering amplitudes associated with elastic scattering “trajectories” [25] (and their time-reversed contours), each of which begin and end at that point and have lengths less than $L_H$. For the point at the center of the grain all relevant trajectories are within the grain and hence the MR is the same as that for a homogeneous film with a sheet resistance equal to that of the weakly disordered grains (i.e. $<1 \text{ k}\Omega$). For the point near the junction we must consider trajectories which cross the junction and return. As long as these trajectories are statistically independent the contribution to the MR will have the same form as for the center of the grain, but will be weighted by the intergrain tunneling resistance $R_1$. When the latter is much larger than the intragrain sheet resistance, as in our highly oxidized films, the overall prefactor of the total MR is predominantly determined by $R_1$.

The inset in fig. 3 shows $r_0(T)$, as determined using eq. (1), for different levels of oxidation. A most remarkable observation is that, for each level of oxidation, $r_0$ crosses $\hbar/e^2$ at approximately the same temperature below which the thermopower data deviate from metallic ($T$-linear) behavior. This is consistent with our suggestion that $E_{\text{eff}}^c$ depends on temperature through $R_{\text{C}}(T)$. Extending this idea further, in fig. 4 we relate the magnetoresistance and thermopower data by plotting $E_C$ versus $r_0$. We first extract the dependence of $E_C$ on temperature by assuming that the thermopower is given by $S \propto E_{\text{eff}}^c / k_B T$. We draw $1/T$ curves through each of the $S(T)$ data points at temperatures below the crossover and $E_{\text{eff}}^c$ is then obtained from the slope. We next determine $E_C(r_0)$ by interpolating from the $r_0(T)$ data (fig. 3, inset). The universal sharp transition in $E_C$ versus $r_0$ for both different oxidation and temperatures may reflect a fundamental interplay between Coulomb interaction effects and localization in granular metals. This poses a stimulating theoretical problem for future investigation.

We conclude with some general comments. We expect $E_{\text{eff}}^c$ to saturate at its geometric value ($E_C$) when $R_i > 10 \text{ k}\Omega$, and presumably this occurs at temperatures somewhat lower than 0.2 K for the intermediate levels of oxidation. Thus in the $T \to 0$ limit we expect the thermopower for all levels of oxidation to behave as $S \propto E_C/k_B T$; the curves should not cross. The tendency of the curves to merge at the lowest temperatures suggests that the average geometric ca-

![Fig. 3. Magnetoresistance at several temperatures for the oxidized film with $R_0(4.2 \text{ K}) = 38.1 \text{ k}\Omega$. The inset shows $r_0$ versus $T$ for different oxidation levels as determined from the magnetoresistance data (see text). Values of $R_0(4.2 \text{ K})$ are, from bottom to top (in k\Omega): 1.47, 8.36, 18.0, 38.1, 62.6. The dashed line is $r_0 = \hbar e^2 = 4.12 \text{ k}\Omega$.](image)
Fig. 4. The charging energy $E_C$, as determined from the thermopower data, plotted versus $R_\Omega$ for three levels of oxidation (values of $R_\Omega$ (4.2 K)).

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References
