Dynamical Coulomb Blockade as a Local Probe for Quantum Transport

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Quantum fluctuations are imprinted with valuable information about transport processes. Experimental access to this information is possible, but challenging. We introduce the dynamical Coulomb blockade (DCB) as a local probe for fluctuations in a scanning tunneling microscope (STM) and show that it provides information about the conduction channels. In agreement with theoretical predictions, we find that the DCB disappears in a single-channel junction with increasing transmission following the Fano factor, analogous to what happens with shot noise. Furthermore we demonstrate local differences in the DCB expected from changes in the conduction channel configuration. Our experimental results are complemented by ab initio transport calculations that elucidate the microscopic nature of the conduction channels in our atomic-scale contacts. We conclude that probing the DCB by STM provides a technique complementary to shot noise measurements for locally resolving quantum transport characteristics.

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An important consequence of the downscaling of electronic circuits towards the atomic limit is the emergence of charge quantization effects [1–5]. The concomitant quantum fluctuations of charge and phase carry valuable information about transport processes [6], such as channel configuration, spin polarization, or effective charge [7–17]. Accessing them experimentally, however, for instance through shot-noise measurements [18] is quite challenging, but feasible [19–26]. Alternatively, the dynamical Coulomb blockade (DCB) is also a consequence of quantum fluctuations. It arises from the inelastic interaction of tunneling electrons with the local electromagnetic environment [27–32], in which the junction is embedded [see Fig. 1(a)]. It appears when the thermal energy $k_B T$ with the temperature $T$ and the Boltzmann constant $k_B$ is on the order of or smaller than the charging energy $E_C = e^2/2C_J$ associated with the capacitance $C_J$ of the tunnel junction ($e$ is the elementary charge). The DCB is directly observable in differential conductance data, where it manifests itself as a

![FIG. 1. (a) Schematic representation of an atomic tunnel junction in the DCB regime and corresponding energy diagram highlighting the environmental interaction. (b) Topography of a single Al adatom adsorbed on the Al(100) surface. The Al adatom is located in the lower half, in the upper part an intrinsic defect is visible. (c) Approach curve on the Al adatom with an Al tip (both in the normal conducting state) at a bias voltage well above the DCB dip. In (d) the dip in the normal conducting $dI/dV$ curve, prototypical for the DCB, is shown with a $P(E)$ fit in the low-conductance limit.](image-url)
dip in the voltage range on the order of $E_C/e$ around zero bias [33–36], as, for example, at very low temperatures ($\lesssim 1$ K) in small capacitance (few fF) mesoscopic circuits [29–38].

In this Letter, we exploit the DCB in ultralow temperature scanning tunneling spectroscopy (STS) as a tool to locally identify the quantum transport characteristics of atomic-scale junctions all the way from the tunnel to the contact regime. First, we use a junction formed between two single atoms featuring a single dominant transport channel [39]. The DCB is seen at low transmission, but disappears with increasing transmission following the Fano factor of a single-channel junction [40]. Extending the measurements to a junction between a single atom on one side and two atoms on the other side, we find a different signature in the DCB dip. This indicates a direct influence of the set of nonvanishing channel transmissions $\tau_i$, also referred to as the mesoscopic PIN code characteristic for the junction [41], on the DCB. We conclude that DCB measurements in STS below 1 K provide direct access to the mesoscopic PIN code as a technique complementary to shot noise measurements [14,40,42–45] as well as other techniques [46,47].

We first use the atomic manipulation capabilities of the scanning tunneling microscope (STM) to construct a junction between two single aluminum atoms [see Fig. 1(a)]. One Al atom is placed at the Al tip apex and one on the (100) surface of an Al crystal, as shown in the lower half of Fig. 1(b). By applying a magnetic field of 20 mT, the superconductivity in Al is quenched and we obtain a normal conducting junction at an experimental temperature $T = 15$ mK [48]. We can reproducibly and continuously tune the junction conductance up to the quantum of conductance $G_0 = 2e^2/h$ (with Planck’s constant $h$) by changing the tip-sample distance, as we illustrate in Fig. 1(c).

We start by studying the differential conductance $G(V)$ in the tunnel regime at bias voltage $V$, where the set point conductance $G_N = G_0 \sum \tau_i = G_0 \tau_i$ and $G_N \ll G_0$. As we show in Fig. 1(d) for $G_N = 0.027 G_0$, the conductance exhibits a dip at low bias voltage, which is the typical signature of DCB. To verify this observation we analyze our data using the $P(E)$ theory [29,37,49]. In the $P(E)$ model, the interaction of tunneling charged particles with the environment is taken into account by the environmental impedance $Z(\omega)$, as shown schematically in Fig. 1(a). The obtained fit is indicated in Fig. 1(d) as an orange line. We find for the junction capacitance $C_J = 21.7$ fF and for the effective temperature $T_{\text{eff}} = 84.9$ mK. The fit confirms that we operate in a low-impedance regime, where the zero frequency part of the environmental impedance is $R_{\text{env}} = 377 \, \Omega$ and much smaller than $1/G_0 = R_Q$ [40], resulting in a small reduction in conductance $\delta G(0) = G(0) - G_N$ at zero bias voltage of $\delta G(0)/G_N = -9\%$. The modeling is detailed in the Supplemental Material [50].

This establishes the DCB in the tunneling regime at low conductances. However, as we approach the tip to the adatom on the sample, the conductance increases, and we observe a clear reduction in the DCB. The experimental data is shown in Fig. 2(a) for different conductance values ranging from 0.03 $G_0$ close to 1 $G_0$. The spectra have been normalized to the set point conductance $G_N$ in the voltage range outside of the DCB dip. The reduction in conductance at zero bias voltage $\delta G(0)$ gradually decreases until it disappears at the highest conductance. This suppression of the DCB as the channel transmission approaches the ballistic limit of perfect transmission ($\tau_1 \rightarrow 1$) has been observed in other types of quantum point contacts [33–35,58]. It can be understood by considering the suppression of fluctuations in the number of electrons transmitted through the junction with increasing transmission, which is captured in the Fano factor $F = \sum \tau_i (1 - \tau_i)/\sum \tau_i$. The relative change in conductance $\delta G(V)/G_N$ for weak coupling to the environment $Z(\omega)$ and at zero temperature was derived for a single-channel system in Ref. [40] and for multiple channels in Ref. [59]:

$$\frac{\delta G(V)}{G_N} = -F \int_{V}^{\infty} \frac{d\omega}{\omega} \text{Re}Z(\omega).$$

FIG. 2. DCB dip as a function of junction conductance. In (a) we present $dI/dV$ data with junction conductances ranging from 0.03 $G_0$ up to 0.99 $G_0$. The data is normalized to the conductance values outside of the DCB dip. (b) The theoretical dependence based on Ref. [40]. Parameters were determined by the $P(E)$ fit in Fig. 1(d), the color code corresponds to (a). (c) The $dI/dV$ reduction at zero bias $\delta G(0)/G_N$ dependent on junction conductance, is plotted as blue circles for the single atom and as a yellow diamond for the dimer. We added a linear fit to the data assuming a single-channel junction $\tau_1 = \tau_i$, where the dip reduces in magnitude with increasing conductance as $(1 - \tau_1)$ from its value in the tunneling limit. For comparison, a dashed line $(1 - \tau_i/2)$ is shown, representing the behavior of a corresponding junction with two equal channels $\tau_1 = \tau_2 = \tau_i/2$. 

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The integral in Eq. (1) shows that for a generally small environmental impedance \(\text{Re}Z(\omega) \ll R_0\), as realized in the STM, the change in conductance will be comparatively small. In Fig. 2(b) we model the transmission-dependent DCB dip based on the theory in Ref. [40] for one transmission channel \(\tau_1\) (see also Supplemental Material [50]). We use the same parameters for the environmental interaction as before in the \(P(E)\) fit depicted in Fig. 1(d) and find good agreement with the data. The decrease of the conductance \(G_N\) is shown in Fig. 2(c) as blue circles. Here, we plot the reduction \(\delta G(0)/G_N\), which can be directly extracted from the experimental data [cf. Fig. 2(a)] and is independent of the \(P(E)\) fit. It follows a \((1 - \tau_1)\) dependence as expected from the Fano factor \(F\) [cf. Eq. (1)] and verified through the linear fit. This finding of pronounced single-channel characteristics in a junction between two Al atoms is consistent with previous experimental results obtained using the subgap structure of the current in the superconducting state [39].

In order to understand the observation of a single channel and to elucidate its origin, we have performed quantum transport calculations within the Landauer-Büttiker approach for coherent transport using a method that combines density functional theory (DFT) with nonequilibrium Green’s function (NEGF) techniques. In particular, this approach makes it possible to optimize the junction geometries, to compute their electronic structure and transport characteristics, including the transmission eigenchannels [60]. As in the experiment the Al sample is modeled as a (100) surface with an additional Al adatom. The structure of the tip oriented along a (100) direction, and the sample are displayed in Fig. 3(a). The channel transmissions \(\tau_i\) were extracted as a function of tip-sample distance, as is visible in Fig. 3(b). We can clearly see that the calculations reproduce the single-channel nature of the atomic Al contact. The transmissions of the second and third channel \(\tau_2\) and \(\tau_3\) are about 2 orders of magnitude smaller than those of the dominant channel \(\tau_1\) over the full range of \(z\) values considered, in contrast to the situation in break junction experiments [61–63]. Since higher order channels contribute even less, we focus on \(\tau_1, \tau_2, \tau_3\) in the following [64], corresponding to the valence states of Al [63,65]. Further insight can be obtained by calculating the complex-valued scattering-state wave functions of the transmission channels [66,67], as shown in Fig. 3(c). For an electron wave impinging on the contact from the substrate, we observe that the dominant first transport channel is of \(\sigma\) symmetry in the narrowest part of the junction. In comparison, the second and third channels exhibit a \(\pi\) shape when viewed along the transport direction. Thus, the theoretically calculated PIN code is \((0.575, 0.003, 0.001)\), which implies that the first channel provides 99.3% of the total transmission. Similar theoretical results were obtained for a junction geometry with an atomically sharp tip oriented along the (111) direction (see Supplemental Material [50]). From the experimental data at higher transmission, we estimate that channels beyond the first contribute no more than 3% to the total transmission at 0.99 \(G_0\), which agrees nicely with the theoretical results.

Exploiting the local atomic resolution and manipulation capabilities of the STM, we can build more complicated atomic structures on the surface such as a dimer of Al atoms. This is visualized in Fig. 4(a), where the dimer is marked in purple. We placed two Al atoms on face-centered cubic lattice sites parallel to the atomic rows, separated by one site. Approaching the tip over the bridge position of the dimer, we anticipate more than one significant transport channel in the junction. The DCB spectrum for the dimer is shown in Fig. 4(b) as a blue line together with a measurement on a monomer. Both of them are taken at a total conductance of 0.58 \(G_0\). The characteristic dip at zero bias voltage is clearly visible. Comparing the \(dI/dV\) curve on the dimer with the one on the monomer, we find that the DCB dip for the dimer is much more pronounced. From the experimental data on the dimer we extract a conductance reduction at zero voltage of \(\delta G(0)/G_N = −5.4\%\), whereas the reduction on the monomer at the same \(G_N\) value is \(\delta G(0)/G_N = −3.7\%\) [68]. Considering the identical total conductance, this is only possible if the number of transmissive channels has changed, such that the first channel has a lower transmission, which leads to a more pronounced DCB dip. Analyzing the dimer DCB dip, we...
consider two contributing channels and experimentally find a PIN code of (0.46,0.12), with an estimated uncertainty of ±0.05 for each channel.

Like for the monomer, we simulated the junction with the dimer to gain further insight into the microscopic origin of the transport channel configuration. The wave functions for the channels 1, 2 and 3 are displayed in Fig. 4(c) for $G_N = 0.58 G_0$. The simulations yield a PIN code of (0.543, 0.029, 0.003) for a (100)-oriented tip and (0.540, 0.034, 0.004) for a (111)-oriented tip, in acceptable agreement with the experimental findings (see details below). For these configurations, 93.6% and 93.1% of the total transmission is carried by the first channel, respectively. This is in contrast to the simulations of the monomer at the same conductance [(100)-tip orientation: (0.575, 0.003, 0.001); (111)-tip orientation: (0.576, 0.002, 0.002)], where in both configurations channel 1 contributes more than 99% to the total transmission (see Supplemental Material [50]). Hence, the transport channel configuration has clearly changed between the monomer and the dimer. Even if our calculations predict that the transport between the dimer and tip is dominated by the first channel, the transmission of the second channel is enhanced by one order of magnitude with respect to the monomer. For this reason we regard the dimer-tip system as a two-channel junction.

The experimentally observed more pronounced DCB dip on the dimer than on the monomer is in agreement with these predictions. Quantitative differences between theory and experiment for the dimer may arise from uncertainties in the precise atomic configuration of the tip, e.g., deviations from a perfect single-atom apex. Such deviations are visible as a small distortion of the dimer in Fig. 4(a). Considering that a change of the tip in the calculations, which is expected to be hardly visible in the topography, already yields a 14% change of $\tau_2$ demonstrates the sensitivity of our method.

To test the range of applicability of this technique, we measured the DCB also in the high-temperature limit. This data was taken on the crystal surface at 1.32 K and 0.13$G_0$, see Fig. 5(a). We model it with the same values of the parameters describing the electromagnetic environment in the $P(E)$ fit of the DCB in Fig. 1(d), only changing the temperature. While we find overall consistency between low- and high-temperature data and modeling, the dip at high temperature only reduces the conductance by about 1%, making it more challenging to detect changes. To reduce the error bar on these measurements, the strength of the DCB needs to be significantly increased. This can be achieved by changing the junction capacitance, since a smaller $C_J$ yields a more pronounced dip. To illustrate the effect, we model the DCB within an experimentally relevant range of $C_J$ between 1 and 60 fF and temperatures between 10 mK and 1.5 K based on the $P(E)$ model [36]. All other parameters are kept at the values used above. The obtained dependence is representative for the tunneling regime ($\tau_e \ll 1$) and is plotted in Fig. 5(b). Our calculation shows that even in the high-temperature limit, small-capacitance junctions should yield a reasonable $\delta G(0)/G_N$. The junction capacitance can be changed by adjusting the macroscopic tip geometry [36]. Therefore, we surmise that a number of experiments would profit by probing local PIN code variations using the DCB. The trade-off in energy resolution due to the reduced capacitance is likely not an issue at higher temperatures (of around 1 K) due to prevalent thermal broadening [36]. In this sense using the DCB to extract the transport
characteristics becomes a viable, complementary alternative to shot-noise measurements.

In summary, we have shown an alternative path to access transport properties on the atomic scale based on the DCB, applicable with standard measurement electronics. Apart from the Fano factor dependence of the DCB in the STM, we have demonstrated that it can be used in normal-conducting junctions to extract local changes of the mesoscopic PIN code, where Andreev reflections cannot be exploited [62,63]. As a perspective, the DCB measurements in the STM should be further extendable to other properties accessible by shot noise, including the spin polarization of tunneling particles and possibly also the determination of their effective charge.

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