TOPOCAL REVIEW

Charge transport through single molecules, quantum dots and quantum wires

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Abstract
We review recent progress in the theoretical description of correlation and quantum fluctuation phenomena in charge transport through single molecules, quantum dots and quantum wires. Various physical phenomena are addressed, relating to cotunneling, pair-tunneling, adiabatic quantum pumping, charge and spin fluctuations, and inhomogeneous Luttinger liquids. We review theoretical many-body methods to treat correlation effects, quantum fluctuations, non-equilibrium physics, and the time evolution into the stationary state of complex nanoelectronic systems.

List of symbols

In this section we provide a list of the most relevant symbols used:

| Symbol | Description |
|--------|-------------|
| $n_{a,k}$ | electron number |
| $k$ | label of the orbital state |
| $\rho_0$ | density of states |
| $\mu_0$ | electrochemical potential |
| $p(t)$ | dot density operator |
| $L_D$ | dot Liouville operator |
| $\Sigma(t,t')$ | transport kernel |
| $I_\alpha$ | tunnel current |
| $\epsilon_\sigma$ | level energy |
| $N$ | charge number |
| $\omega$ | frequency |
| $D_N$ | magnetic parameter |
| $\lambda$ | electron vibration coupling |
| $\Omega$ | dwell time of electrons in the system |
| $\delta_{i}$ | distance to resonances |
| $\Sigma(\omega)$ | self-energy |
| $G$ | conductance |
| $\tilde{p}$ | matrix $\tilde{p}$ in Laplace space |
| $h$ | magnetic field |
**1. Introduction**

In this review we consider different aspects of charge transport through nanoscale devices attached to electronic reservoirs, focusing on theoretical approaches dealing with interaction and quantum fluctuation effects. Transport experiments have shown convincingly that many of these systems—from carbon nanotubes down to single nanometer sized molecules—behave as quantum dots. A quantum dot is a confined system of electrons, which is so small that the discreteness of energy spectrum with a level spacing $\Delta \epsilon$ becomes important; it is therefore often referred to as an artificial atom. Due to the smallness of the quantum dot, Coulomb interaction between the electrons also plays an important role: the charging energy $U$ which can be of the same order as the level spacing. If the time evolution itself can be used as a tool to reveal various correlations and transport through inhomogeneities leads to complex excitations (e.g., due to spin–orbit interaction or mechanical vibrations), Luttinger liquid physics, non-equilibrium, and quantum fluctuation effects. Transport experiments have shown convincingly that many of these systems—from carbon nanotubes down to single nanometer sized molecules—behave as quantum dots. A quantum dot is a confined system of electrons, which is so small that the discreteness of energy spectrum with a level spacing $\Delta \epsilon$ becomes important; it is therefore often referred to as an artificial atom. Due to the smallness of the quantum dot, Coulomb interaction between the electrons also plays an important role: the charging energy $U$ which can be of the same order as the level spacing. If the quantum dot is attached to reservoirs by tunneling contacts, electrons can leave and enter the dot and the single-particle levels are therefore broadened due to the finite lifetime of the electrons. For sufficiently weak tunnel coupling $\Gamma$ and small temperature $T$, the level spacing $\Delta \epsilon$ and the charging energy $U$ can be resolved at standard cryogenic temperatures. This opens up the possibility of performing transport spectroscopy by measuring the differential conductance as a function of gate voltage $V_g$ and bias voltage $V$. The transport characteristics allow counting of the discrete charge states of such systems, and even more complex degrees of freedom of the quantum dot are revealed. As an example, the spin states can be identified with the help of the Zeeman effect, and also vibrational motion of the quantum dot system can be detected.

The transport behavior, however, depends strongly on the ratios of all the energy scales, and on the way transport is induced, e.g., by static or time-dependent applied voltages, or by measuring linear or nonlinear response in the transport voltage. In particular, for molecular scale quantum dots the excitation spectra may be quite complex and various splittings appear. This complicates matters further but also leads to a variety of transport effects. Besides stationary properties of quantum dots, the time evolution out of an initially prepared non-equilibrium state is another important issue of recent interest, providing the new energy scale $\hbar/\tau$. The time evolution itself can be used as a tool to reveal various many-body aspects. It is also of practical importance due to the progress in the field of solid-state quantum information processing, initiated by the suggestion to use spin states of quantum dots as basic elements of qubits [1]. In addition to the ‘bare’ energy scales mentioned so far, due to correlation effects new effective energy scales can be generated, which describe the physics of quantum fluctuations induced by the reservoir–dot coupling. These new energy scales are complex functions of the above ‘bare’ energies and we denote them in the following by $T_c$, where $c$ stands for ‘cutoff’.

Examples are the Kondo temperature describing the strong coupling scale for spin fluctuations (usually denoted by $T_K$) and renormalized tunneling couplings generated by charge fluctuations. These scales will dominate the measured transport in the strong coupling regime where $T_c$ is the largest energy scale. However, also in the weak coupling regime quantum fluctuations set in and are of particular interest close to resonances, where they can be analyzed by a perturbative treatment in the renormalized couplings. To understand the new energy scales physically and calculate the renormalized couplings, renormalization group (RG) ideas are required. RG transport theories reveal that interactions are of key importance for the generation of new effective energy scales by quantum fluctuations. Without interactions, quantum fluctuations are well understood, they just lead to an unrenormalized broadening of the energy conservation by the tunneling coupling. Although new effects from correlations are most pronounced in the regime of strong correlations $\Gamma \ll U$, interesting quantum fluctuation effects already start to become visible in the regime of moderate correlations $\Gamma \sim U$, where perturbative and mean-field theories are often not yet applicable. Another interesting regime is the case of a continuous level spectrum, i.e., where the quantum dot is replaced by a one-dimensional quantum wire. In this case, quantum fluctuations in the wire itself lead to renormalization effects and Luttinger liquid physics, with typical power law suppression of the bulk spectral density.

Similar to the case of quantum dots, the simultaneous presence of correlations and transport through inhomogeneities leads to very rich phenomena. A prominent example is the suppression of the spectral density at wire boundaries, which leads to vanishing conductance at zero temperature and bias voltage if a quantum wire is weakly coupled to two leads. This has to be contrasted to the Kondo effect in quantum dots, where conductance becomes maximal when a single spin is placed in an ‘insulating’ quantum dot.

In this review we will illustrate these key issues and concentrate on correlation effects in quantum dots and quantum wires. We will describe transport theories specifically designed to treat the case of moderate to strong Coulomb interactions and quantum fluctuation effects. With these methods it is possible to account for the complex interplay of various combinations of quantum mechanical effects (level quantization, interference, spin and angular momentum), complex excitations (e.g., due to spin–orbit interaction and/or mechanical vibrations), Luttinger liquid physics, non-equilibrium (due to static as well as time-dependent applied voltages), and the time evolution into the stationary state. An important aspect is that these methods apply to general models incorporating experimentally relevant microscopic details, and offer prospects for numerical implementation. Despite these

\[ J_{1/2} \] exchange couplings
\[ W_t \] rates for the process $s \rightarrow s$
\[ \chi \] susceptibility
\[ \Lambda \] flow parameter
\[ \Lambda_c \] cutoff
\[ K \] Luttinger liquid parameter
\[ v_{c/s} \] velocities of the charge/spin-density excitations
\[ k_F, v_F \] Fermi momentum, Fermi velocity
\[ U' \] nearest-neighbor interaction
\[ \alpha_{\text{bulk}} \] power law exponent of the wire spectral function
\[ \alpha_{\text{end}} \] power law exponent of an electron tunneling into the end of a LL.
common themes, the different sections of this review are meant to be almost self-contained and each section closes with a short outlook. We consistently use natural units $\hbar = k_B = e = 1$. The review is organized as follows.

In section 2, we will introduce a general approach to transport through quantum dots of arbitrary complexity, aiming at the description of three terminal single molecule junctions. The tunneling coupling is treated perturbatively up to second order in the coupling $\Gamma$, accounting for charge fluctuation effects, while not capturing spin fluctuations which start in higher order. Besides well-known level renormalization and cotunneling effects, electron pair-tunneling is shown to give rise to resonances, even for the most basic quantum dot model. It is shown how the complexity of, e.g., vibrational and magnetic excitations of molecular quantum dots needs to be accounted for in transport spectroscopy calculations.

Section 3 uses the same perturbation theory but extends it to transport through systems which are subject to time-dependent fields. Using the example of adiabatic charge pumping through a single-level quantum dot, it explains why time-dependent external fields in the adiabatic regime are of particular interest. It is shown how time-dependent fields can be used to reveal level renormalizations induced by charge fluctuations, an effect purely due to the Coulomb interaction. This offers the interesting perspective to use time-dependent fields as fingerprints for the combined influence of quantum fluctuations and correlations.

The regime of strong quantum fluctuations is addressed in section 4 in the linear response regime. It is described how a recently developed functional RG (fRG) method is a useful tool to treat generic quantum dot models. Within this method a systematic expansion in a renormalized Coulomb interaction is performed. However, the tunneling coupling is treated non-perturbatively, so that the strong coupling regime can be addressed for spin as well as for charge fluctuations, at least for moderate Coulomb interactions. Two prominent examples of strong quantum fluctuations are reviewed: the Kondo effect in moderate Coulomb interactions. Two prominent examples of strong quantum fluctuations are addressed for spin as well as for charge fluctuations, at least for non-perturbatively, so that the strong coupling regime can be accounted for in transport spectroscopy calculations.

Section 5 describes quantum fluctuations in nonlinear response, i.e. at finite bias voltage $V \gg T$, together with the time evolution into the stationary state. The basic physics of weak spin and strong charge fluctuations is illustrated on recent results for two elementary models: the anisotropic Kondo model at finite magnetic field and the interacting resonant level model. A real-time renormalization group (RTRG) method is introduced, where, complementary to the fRG scheme presented in section 4, an expansion in a renormalized tunneling coupling is performed whereas the Coulomb interaction is treated non-perturbatively. This allows the description of strong correlation effects $\Gamma \ll U$ for moderate tunneling.

Finally, in section 6 we describe transport properties of quantum wires. The status of Luttinger liquid physics is reviewed, in particular in connection with the presence of inhomogeneities, backscattering, interference effects, and non-equilibrium. As in section 4, the fRG scheme is shown to be a unique method, capable of describing the whole crossover from few-to multi-level dots up to the limit of continuous spectra in quantum wires. Its ability to simultaneously incorporate microscopic details is of particular importance for the description of experiments.

2. Transport through single molecule quantum dots

Molecular quantum dots present the ultimate miniaturization of quantum dot devices, which have been realized in semiconductor heterostructures and wires, metallic nanoparticles, and carbon nanotube molecular wires. Various methods have been developed to contact nanometer size single molecules. Transport measurements in three terminal junctions have most clearly demonstrated that single molecules also are quantum dots, although of a particularly complex nature, see for a review [2–4]. In view of this a central theoretical challenge is to describe the transport through a quantum dot of arbitrary complexity, covering the entire, broad class of systems mentioned above. We now first outline this general framework, which is relevant also for sections 3 and 5. For details see [5] and the references therein.

Starting from the limit of a completely opaque tunnel junction, we can specify the exact many-body level spectrum and states $|s\rangle$ of the quantum dot in this case,

$$H_D = \sum_s E_s |s\rangle \langle s|.$$  \hspace{1cm} (1)

Often, only a few of these states are actually accessible in an experiment. Subsequently, we incorporate the non-zero transparency of the junction in the coupling

$$H_T = \sum_{s,s',\alpha\kappa} T_{\alpha\kappa}(s'|s) |e_{\alpha\kappa}| + \text{h.c.}$$  \hspace{1cm} (2)

through the amplitudes $T_{\alpha\kappa}(s'|s)$ for an electron with spin $\sigma$ to tunnel on/off the dot from/to one of the electronic reservoirs labeled by $\alpha = L, R$. Generally, in equation (2) all states $s$ and $s'$ are coupled which differ by the addition of one electron. The reservoir $\alpha$ by itself is described by

$$H_\alpha = \sum_{\kappa\sigma} (\epsilon_{\kappa\sigma} + \mu_\alpha) n_{\alpha\kappa\sigma},$$  \hspace{1cm} (3)

where $n_{\alpha\kappa\sigma}$ denotes the electron number operator and $\kappa$ labels the orbital states. Each electrode is restricted to contain effectively non-interacting fermion quasi-particles with density of states $\rho_\alpha$. We make the further statistical assumption that the electrochemical potential $\mu_\alpha$ and temperature $T$ are well defined for each reservoir $\alpha$ separately. The central quantity is the reduced density operator $\rho$ of the dot, obtained by partial averaging of the full density operator over the reservoirs. From $\rho$ any non-equilibrium expectation value of a local observable can be calculated, and also (see below) the transport current. Starting from the above general model, one can derive the
kinetic equation from which the time evolution of the dot density operator can be calculated:

$$\frac{d}{dt} p(t) = -i L_D p(t) - i \int_0^t dt' \Sigma(t, t') p(t').$$

(4)

Here the Liouvillian superoperator $L_D$ acts on a density operator by commuting it with $H_D$, i.e., $L_D p(t) = [H_D, p(t)]$. This linear transformation generates the ‘free’ time evolution of the dot density matrix, i.e., in the case where it is not coupled to the electrodes. The challenge lies in the calculation of the transport kernel $\Sigma(t, t')$ which incorporates all effects due to the coupling to the electrodes which is adiabatically switched on at time $t_0$ ($t_0 < t$). The superoperator $\Sigma(t, t')$ is a non-trivial functional of three ‘objects’: (i) the dot Liouvillian $L_D$ (energy differences of initial, final and virtual intermediate states of processes), (ii) the dot part of the tunnel operator (tunnel vertices) and (iii) the product of the Fermi distribution functions $f((\epsilon_{\alpha s} - \mu_\alpha)/T)$ and density of states $\rho_\alpha$ of all electrodes $\alpha$. Importantly, compact diagrammatic rules for the calculation of this kernel in terms of these three objects are known exactly [5, 6]. In this section and section 3 we review the basic physics which emerges from the calculation of the kernel by perturbation theory in the tunnel operator $H_D^t \sim \Gamma$ to leading and next-to-leading order. This is applicable to the ‘high temperature’ regime, where higher order effects are present but still weak. In sections 4 and 5 the low temperature physics due to strong quantum fluctuations is illustrated. In particular, in section 5 the kernel is calculated non-perturbatively in a renormalization group framework. The technical details of these two approaches can be found in [6] and [5], respectively.

We now first review all possible basic types of transport resonances which can arise due to tunneling processes of the first and second order in $\Gamma$. For this purpose we consider a single energy level with Coulomb interaction, the so-called Anderson model, where we emphasize the need to consider a finite charging energy $U$ [7] such that double occupancy is possible (not only as a virtual intermediate state). This will set the stage for the subsequent discussion of examples of transport involving more complex spectra in section 2.2 and the time-dependent transport considered in section 3.

2.1. Transport spectroscopy: sequential, co- and pair-tunneling

Most types of transport resonances observed in quantum dot experiments can be illustrated by the Anderson model sketched in figure 1 where a single orbital level is coupled to electrodes $\alpha = L, R$ with tunnel amplitudes $T_\alpha$. The strength of the tunnel coupling is characterized by the spectral function, $\Gamma_\alpha = 2\pi \rho_\alpha |T_\alpha|^2$, whose energy dependence is neglected. Denoting by $N$ the electron number on the dot, the eigenstates $|s\rangle$ of the dot (when decoupled from the electrodes) are easily classified. For the empty dot ($N = 0$) there is one state $s = 0$ with energy $E_0 = 0$. For the singly occupied dot ($N = 1$) there are two states $s = \sigma = \uparrow, \downarrow$ with energy $E_\sigma = \epsilon_\sigma$. The $N = 1$ states are split by a magnetic field such that $s = \downarrow$ is the ground state and $s = \uparrow$ the excited state. Finally, for the doubly occupied dot ($N = 2$) the energy $E_d = \sum_\sigma \epsilon_\sigma + U$ of the state $s = d$ includes the charging energy. One can show that due to conservation of the electron number and the spin-projection along the magnetic field only the occupation probabilities enter into the description of the non-equilibrium state. We collect these into a vector $p = (p_0, p_\uparrow, p_\downarrow, p_d)^T$. Its time evolution is fully described by a kinetic equation with a matrix kernel $\Sigma(t, t')$,

$$\frac{d}{dt} p(t) = -i \int_0^t dt' \Sigma(t, t') p(t'),$$

(5)

where we have set $t_0 = -\infty$ (cf (4)). For time-independent parameters considered here, we have $\Sigma(t, t') = \Sigma(t - t')$ and the occupancies will reach a constant value in the stationary long-time limit: $p(t) \rightarrow p = \text{constant}$ for $t \rightarrow \infty$. With the subsidiary condition of probability conservation $\sum_i p_i = 1$, this state is determined uniquely by

$$0 = -i \Sigma p.$$

(6)

As expected, here the time integral over the kernel is needed, i.e., the zero-frequency Laplace transform

$$\lim_{\alpha \rightarrow 0} \int_{-\infty}^{\infty} dt e^{it\alpha} \Sigma(t, 0) = \Sigma.$$

This kernel has been calculated in closed form up to order $\Gamma^2$ for an arbitrary dot model Hamiltonian in [6], allowing arbitrarily complex transport spectra to be calculated. From a similar equation for the measurable tunnel current from electrode $\alpha$

$$I_\alpha = -i \int_{-\infty}^t dt' T_{D\alpha} \Sigma^\alpha(t, t') p(t') \rightarrow -i T_{D\alpha} \Sigma^\alpha p,$$

(7)

the stationary current can be found. This requires an additional zero-frequency kernel $\Sigma^\alpha$ which takes account of both the amount and the direction of the electrons transferred from electrode $\alpha$. Here $T_{D\alpha}$ denotes the trace over the dot degrees of freedom.

In figure 2 we show exemplary results for the Anderson model of the calculations outlined above. A complete overview of nonlinear transport resonances is obtained when plotting $dI/dV$ (quantifying changes in the current as new transport processes become energetically allowed) as a function of the static applied bias $V$ (which drives the current) and gate voltage $V_g$ (which uniformly shifts all energy levels $\epsilon_\sigma$ without changing the quantum states)$^4$. In the lower half of the panel

4 Here one assumes to a good first approximation that the bias and gate electric fields are spatially uniform. Corrections to this picture have been calculated in [8] for molecular junctions.
where sequential relaxation of the spin-excited state becomes possible. The Zeeman energy. The skewed green lines indicate where the energy of the spin-excited state becomes occupied when the bias exceeds the inelastic cotunneling threshold, which equals the Zeeman energy. The skewed green lines indicate where the sequential relaxation of the spin-excited state becomes possible (COSET).

![Figure 2. Occupation of excited spin-state, $p_t$ (upper panel) and differential conductance (lower panel) for the single-level Anderson model, plotted as a function of bias voltage $V$ and gate voltage $\beta V_g$, where $\beta$ is the gate coupling factor. The spin degeneracy is lifted by an applied magnetic field: $\epsilon_{\uparrow} - \epsilon_{\downarrow} = 50 T$, where $T$ is the electron temperature. The dot is symmetrically coupled to the left and right electrodes: $\Gamma_L = \Gamma_R = 10^{-4} T = 5 \times 10^{-4} U$. The horizontal green line indicates the inelastic cotunneling threshold, which equals the Zeeman energy. The skewed green lines indicate where the sequential relaxation of the spin-excited state becomes possible (COSET).]

we plot the differential conductance $dI/dV$, whereas in the upper half of the panel the occupation $p_t$ of the spin-excited state of the quantum dot is shown. This figure illustrates the insight provided by transport theory, linking the measured tunnel current to what goes on in the quantum dot. To emphasize this correspondence of the two panels, the bias axes in the lower panel are mirrored with respect to the upper panel.

To the far left and right of the plot the dot is in the $N = 0$ and $2$ state respectively around zero bias. Moving towards the center from there, $dI/dV$ peaks along the first encountered yellow skewed lines (marked ‘SET’) where the condition $\mu_{\alpha} = \epsilon_{\downarrow}$ and $\mu_{\beta} = \epsilon_{\uparrow} + U$ are met. Beyond this line a spin $\downarrow$ electron can now enter or leave the dot in a sequential or single-electron tunnel (SET) process which is of first order in $\Gamma$. In this way the $N = 1$ ground state $s = \downarrow$ can be reached starting from $N = 0$ and $N = 2$, respectively. Once beyond the next skew yellow line (marked ‘SET*’), defined by $\mu_{\alpha} = \epsilon_{\uparrow}$ and $\mu_{\beta} = \epsilon_{\downarrow} + U$, respectively, an electron with spin $\uparrow$ can tunnel onto the dot. This first order process makes the $N = 1$ excited state $s = \uparrow$ accessible, as the upper panel clearly shows.

Moving further towards the center, there is a large triangular region of the $dI/dV$ map where only the $N = 1$ ground state is accessible: sequential tunneling processes out of this state are suppressed to first order since $U \gg T$, which is referred to as Coulomb blockade. However, a second order process, referred to as inelastic cotunneling, allows the excited state to become occupied [9, 10] when the bias exceeds the excitation energy for a spin flip, $\mu_L - \mu_R = V > \epsilon_{\uparrow} - \epsilon_{\downarrow}$. This defines a gate voltage independent line (horizontal), reflecting the fact that the charge state is left unaltered by the spin fluctuation process which flips the spin. Below this onset voltage only elastic cotunneling is possible, giving a smaller conductance and current. In the small triangular region marked ‘ICOT’ in the upper panel, relaxation of the excited state is possible only by inelastic cotunneling. Above this, in the region marked ‘COSET’, the spin-excited state, populated by slow inelastic excitation, can relax by a much faster sequential relaxation process. It therefore is depleted again and the $dI/dV$ shows a small peak as a signature of this cotunneling-assisted sequential tunneling (COSET) [11].

The above processes are well known and have been experimentally observed in various types of quantum dots, see, e.g., [9, 12, 13] and the reviews [2–4, 14]. However, only recently, it was shown that for this elementary model yet another resonance exists in second order in $\Gamma$ [7] by taking into account all many-body states and all contributions to the transport kernel $\Sigma$. This resonance is related to electron pair-tunneling: it shows up along a skewed line in the $dI/dV$ map marked ‘PT’, the change in the contrast indicating a step in the conductance as a function of bias $V$. The resonance condition for this is $2 \mu_{\alpha} = \sum_{\sigma} \epsilon_{\sigma} + U$, indicating that the energy is paid (gained) by two electrons coming (going) from reservoir $\alpha$. This electron pair can be added to (extracted from) the dot in one coherent process, thereby changing the charge from $N = 0$ to $2$ (or vice versa). The electrochemical potential $\mu_{\alpha} = \frac{1}{2} (\sum_{\sigma} \epsilon_{\sigma} + U)$ at which this happens equals the average to the above mentioned SET resonances, i.e., due to quantum charge fluctuations the Coulomb energy paid (gained) per electron is reduced by $\frac{1}{2}$.

Finally, we note that the resonance positions of the sequential tunneling peaks are slightly shifted with respect to the above mentioned values $\mu_{\alpha} = \epsilon_{\downarrow}, \epsilon_{\uparrow} + U$ due to the second order energy level renormalization which is consistently accounted for. Although this may seem unimportant here, in section 3 it will be shown that in time-dependent transport this level renormalization effect may actually generate and dominate the full current.

2.2. Complex excitation spectra: vibrations and spin–orbit splitting

Molecular quantum dots are clearly more complex than the above considered model, for instance by their ability to mechanically vibrate. One important general aspect of this complexity is that the only conserved quantity in a tunnel process is the electron number$^5$. In contrast, the quantum numbers of, e.g., various vibrational modes of molecular quantum dot may change when an electron tunnels onto it. This implies that the transport theory has to account explicitly for the density matrix off-diagonal elements between different excited states with the same charge number $N$. As is well known, this is important already in first order in $\Gamma$ when there are degeneracies of the energy levels $E_r$ in the same charge ($N$) state on the scale set by the tunneling $[15–17]$. In the case where there are no such degeneracies only the probabilities matter to order $\Gamma$. However, it was shown only recently, that even in this simplest case this is no longer true.

$^5$ Conserved here refers to the system of reservoirs and dot, including their interaction.
In second (or higher) order in \( \Gamma \) and equation (6) can no longer be used. Instead, one can derive from (5) an effective stationary state master equation for the probabilities of the same form as (6), but with an effective kernel which contains important corrections from the off-diagonal elements [6, 18]. An example where these corrections are found to be very important is the Franck–Condon effect. This effect arises in its most elementary form in the Anderson–Holstein model, where the dot is described by an Anderson level (discussed in section 2.1) plus a single vibrational mode with frequency \( \omega \):

\[
H_D = \sum_\sigma \varepsilon_\sigma n_\sigma + U n_\uparrow n_\downarrow + \lambda \sqrt{2} Q \frac{\omega}{2} \sum_\sigma n_\sigma + \frac{\omega}{2} (P^2 + Q^2). \tag{8}
\]

Here \( Q = (b^\dagger b)/\sqrt{2} \) is the vibrational coordinate normalized to the zero-point motion and \( P = i(b^\dagger b)/\sqrt{2} \) the conjugate momentum. Due to the linear coupling term \( \lambda \), the vibrational mode is displaced when charging the molecule. For \( \lambda > 1 \) this causes a suppression of the Franck–Condon overlap of ground state vibrational wave functions for two subsequent molecular charges \( N \). As a result the tunnel rates calculated to lowest order in \( \Gamma \) are suppressed and the sequential tunneling current is blocked, even for gate voltages close to the resonance [19–21], an effect also called ‘Franck–Condon blockade’ [22]. This implies that second order processes become of crucial importance even at resonance [23]. The general approach developed in [6] consistently accounts for all second order effects, i.e., not just cotunneling rates as in [23]. In figure 3 we show the differential conductance calculated to second order where the vibrational mode is shifted by several multiples of its zero-point amplitude. Whereas for \( \lambda = 1 \) the standard Coulomb blockade picture is still identifiable, for \( \lambda = 4 \) it is drastically altered: inelastic cotunneling (threshold at \( \mu_L = \mu_R = \omega, 2\omega, \ldots \) ) and COSET [24] resonances proliferate in the transport spectrum. Experimentally, vibration assisted transport in three terminal molecular devices has been reported [25–27]. Significant Franck–Condon blockade has been observed in suspended carbon nanotubes and compared with theory in [28]. Other theoretical studies have studied the effect on current noise [22, 29], the Kondo effect [30, 31], the influence of mechanical dissipation (finite Q-factor) [32, 33] or the lack thereof [34], vibrational anharmonicity [35, 36] and interference effects [35, 37].

All above works rely on the Born–Oppenheimer separation of nuclear and electronic motion on the molecule. Transport effects signaling the breakdown of this separation, going under the generic name of (pseudo-) Jahn–Teller dynamics, have also been calculated [38–40]. A very recent STM experiment [41] has confirmed these transport signatures of the breakdown of the Born–Oppenheimer approximation. In this work SET resonances were measured as a function of a molecular parameter, as suggested in [40], in this case by measuring many individual molecular wires of various lengths.

Another class of examples of quantum dots with complex excitations are magnetic molecules. Here the spin–orbit interaction results in violation of spin-selection rules for electron tunneling and the same remarks as above apply. In particular, single molecule magnets, reviewed in [42], offer a basic example and have been of interest in view of quantum computing [43, 44]. In such molecules the ground state has a large spin value \( S > 1/2 \) due to strong exchange coupling of the spins of a few magnetic atoms. This multiplet is however split by strong spin–orbit perturbations. In many cases an accurate model for a given charge state \( N \) is

\[
H_D^{(N)} = -D_N S_N^2 + E_N (S_X^2 - S_Y^2), \tag{9}
\]

where the uni-axial (\( D_N \)) and transverse (\( E_N \)) anisotropy parameters describe the splitting and mixing of spin states, respectively. Recently, three terminal measurements on magnetic molecules have been reported, including comparison with theory [45–48], see for a review [49]. Calculations of the sequential [50–55] and cotunneling [56–60] regimes for models including the anisotropy have identified distinctive
transport signatures and possibly useful effects. An important goal is to gain control over the spin of a single molecule by electric means [61, 62]. Recently, the magnetic anisotropy splittings were measured in a three terminal junction, similar to two terminal STM measurements [63, 64]. Importantly, the magnetic field evolution of inelastic cotunneling [48] was measured in different charge states. It was found that the magnetic $D_N$ parameter depends strongly on the charge state $N$, which can be electrically controlled with the gate voltage.

Beyond the issue of control over single molecule magnetism, is the understanding of coupled molecular magnets or magnetic centers. Apart from spin–blockade effects, expected for large spin molecular ground states [65], the effect of spin–orbit interaction on the exchange coupling between two magnetic centers inside a magnetic double quantum dot is of interest. For instance, the spin–orbit induced Dzyaloshinskii–Moriya or antisymmetric exchange interaction was shown to give rise to a characteristic dependence of the transport on an externally applied magnetic field and the polarizations of magnetic electrodes [66].

Higher order tunneling effects for magnetically anisotropic molecular quantum dots described by (9) have attracted attention [67–69] after the prediction of a novel anisotropic pseudo spin Kondo effect [70]. The spin–orbit induced anisotropy in (9) not only raises a barrier which opposes spin fluctuations (due to $D_N$) (section 4), but it also generates intrinsic spin-tunneling (due to $E_N$). This interplay leads to interesting anisotropic effective exchange interaction with electrons transported through a molecular magnet, as will be discussed in section 5.

Besides their own interesting transport signatures, spin and vibrations can also interact or influence each other in transport. For instance, a mechanism of vibration-induced spin-blockade of transport was predicted for a mixed-valence dimer molecular transistor (‘double dot’) [71, 72].

Finally, we mention that level spectrum complexity may also deceive one when analyzing experimental data. A particularly clear example is offered by recent transport data on carbon nanotube ‘peapods’, i.e. fullerene molecules in a host nanotube [73]. Here weakly gate-dependent $dI/dV$ peaks were observed which are strongly reminiscent of inelastic cotunneling (second order in $\Gamma$, see section 2.1). However, detailed transport calculations have conclusively shown that the complex measured spectrum can be assigned to sequential tunneling processes only, i.e., of first order in $\Gamma$. This deceptive case occurs because the two coupled quantum dots, the host nanotube and the fullerene ‘impurities’ have a very different gate voltage dependence. Fortunately, the significant hybridization of the two dots gives a characteristic interference effect in the first order current, distinguishing it clearly from second order effects. In view of applications it is interesting that the analysis in [73] indicates that the charge state of the fullerene impurities is electrically tunable, independently of that of the host carbon nanotube.

2.3. Outlook

In summary, we have illustrated that general methods for spectroscopy calculations for molecular quantum dot models are essential for the experimental characterization and ultimately the design of nanoscale quantum dot devices. The complex spectra due to various strong many-body interactions play a crucial but complicating role, at the same time however, enabling interesting new possibilities. An exciting future direction important for further progress is to efficiently account for renormalization effects in complex models, in particular as outlined in section 5. Also, the approach outlined here can be extended to other transport quantities, such as spin and heat [74]. The molecule and junction models used here can be parametrized based on atomistic electronic structure calculations, as for instance, in [17, 75, 76]. Pursuing this further is another important step in understanding molecular quantum dots.

3. Adiabatic transport through a quantum dot due to time-dependent fields

Charge transport through quantum dots is not necessarily due to an applied static bias voltage: in the absence of a bias, a current can be obtained by the time-dependent modulation of fields, applied externally to a mesoscopic device. This is of interest for possible future applications in nanoelectronics, where the repeated and fast operation of a device is of interest. Indeed the mesoscopic analog of a capacitor has been realized [77, 78] serving as a fast coherent electron source and charge pumping [79–82] is a candidate for the realization of a quantum current standard, to name some examples. From a fundamental point of view the non-equilibrium created by time-dependent electric fields provides enhanced insight into the quantum properties of a nanoscale device, which are not accessible with equilibrium methods. If the variation of the fields is slow with respect to the internal timescales of the system itself, the modulation is called adiabatic. The system then remains in a quasi-equilibrium state, meaning that the system is not brought to an excited state by the time-dependent modulation.

A prominent example for transport in the absence of a bias voltage is adiabatic pumping: the cyclic variation of at least two of the system parameters, see figure 4, leads to a dc charge transfer through the quantum device. In this case the pumped charge does surprisingly not depend on the detailed time evolution of the pumping cycle; this can be shown by formulating it in terms of geometric phases [83–85]. Theoretically, adiabatic pumping has been extensively studied in the non-interacting regime [86–92]. In this case a scattering matrix approach is convenient [86, 93] and has been broadly used. While indeed Coulomb interactions are of minor importance in some setups [77, 78, 94], they play an important role in many quantum dot systems and lead to a variety of interesting results. Several approaches have been used to address the problem of adiabatic pumping through interacting quantum dots in different regimes, including weak Coulomb interaction [95, 96], pumping in Luttinger

6 The slow variation of the externally applied fields becomes experimentally particularly important for delicate systems such as molecules, see section 2, where problems related to heating, surface excitations (plasmons) and laser-induced chemical reactions (photo-bleaching) should be avoided.
liquids [97], and a quite general approach using non-equilibrium Green’s functions [98–100]. The latter works are particularly appropriate for the study of weak interaction or for the calculation of spin and charge pumping in the Kondo regime [99] and the role of elastic and inelastic scattering processes has been studied in detail [100]. The treatment of the tunnel coupling is non-perturbative in these works and they are in this sense complementary to the results presented in the following.

It turns out that the interplay of Coulomb interaction and coherent tunneling between a quantum dot and leads, leading to an effective renormalization of the energy level, can be at the origin of the pumping mechanism. The result is that this energy renormalization, which is a pure Coulomb interaction effect described above we use a perturbative expansion in the dependence of all parameters to an effective renormalization of the energy level, can be at smaller than \( T^* \), with the lead’s constant \( R \) with time-dependent parameters. This linear correction to the kernel is indicated by the superscript \( (i) \) for instantaneous. The subscript \( t \) to emphasize that the system parameters \( X(\tau) \rightarrow X(\hat{t}) \) are frozen at time \( \hat{t} \), i.e., the functional dependence on \( X(\tau) \) is replaced by \( X(\hat{t}) \rightarrow X(\hat{t}) \). The first order term is obtained by linearizing the time dependence of all parameters \( X(\tau) \) with respect to the final time \( \tau, \) i.e., \( X(\tau) \rightarrow X(\tau) + (\tau - \tau) \frac{d}{d\tau} X(\tau) |_{\tau=} \), and retaining only linear terms in time derivatives. This linear correction to the kernel is indicated by the superscript \( (a) \) for adiabatic. Finally, it is necessary to perform an adiabatic expansion for the occupation probabilities in the dot,

\[
p(t) \rightarrow p^{(i)}(t) + p^{(a)}(t)
\]

\[
\Sigma(t, t') \rightarrow \Sigma^{(i)}(t - t') + \Sigma^{(a)}(t - t').
\]

The instantaneous probabilities \( p^{(i)}(t) \) are the solution of the time-independent problem with all parameter values fixed at time \( \hat{t} \). They are obtained by solving the stationary master equation, equation (6), with all parameters replaced by time-dependent parameters evaluated at time \( \hat{t} \). Once the instantaneous probabilities \( p^{(i)}(t) \) are known, the adiabatic corrections \( p^{(a)}(t) \) are found from the master equation in first order in \( \Omega \) obtained from the expansion described above. Similarly an equation for the current can be developed, see equation (7), and adiabatically expanded, where a current of the decoupled system are given by \( s = \{0, \uparrow, \downarrow, d\} \). Tracing out the lead degrees of freedom the system is described by its reduced density matrix, where here only the occupation probabilities \( p = (p_0, p_\uparrow, p_\downarrow) \) are of interest. Their time evolution in the long-time limit, \( \hat{t}_0 \rightarrow -\infty \), is fully described by the generalized master equation, equation (5). The goal being the description of slowly varying system parameters, it is useful to perform an adiabatic expansion [105]: the frequency of the variation therefore has to be much smaller than the dwell time of electrons in the system, \( \Omega \ll \Gamma \). The effect of the system parameters varying slowly in time is that the density matrix of the system at time \( t \) is not only described by the instantaneous values of the parameters, but it lags a bit behind the parameter change. Therefore, an expansion around this time \( \hat{t} \) suggests itself. To obtain such an expansion as a first step a Taylor expansion of \( p(t') \) is performed around \( \hat{t} \) up to linear order in the integrand on the right-hand side of equation (5). This is related to the fact that memory effects of the kernel have to be taken into account. Furthermore, an adiabatic expansion of the kernel \( \Sigma(t, t') \) itself is performed. The zeroth order term, \( \Sigma^{(i)}(t - t') \), is indicated with the superscript \( (i) \) for instantaneous.

Figure 4. (a) Sketch of a single-level quantum dot with time-dependent parameters attached to two reservoirs. (b) Two of the system parameters are time-dependent as indicated in (a) and perform a closed cycle in parameter space.
kernel $\Sigma^L_\tau$ takes account for the amount and the direction of electron transfer.

On top of the adiabatic expansion a perturbative expansion in the tunnel coupling strength $\Gamma$ is performed up to second order, allowing the actual evaluation of the kernel using diagrammatic rules as elaborated in [101–105], see section 2. This perturbative expansion is valid in the regime of moderate couplings with respect to temperature, i.e. $\Gamma \lesssim T$.

3.2. Pumping current

We are interested in the time-resolved pumping current as well as the charge pumped through the dot per pumping cycle. The pumped charge, $Q = \int_0^T I_L(t)\,dt$, in the bilinear response regime, i.e. for modulations with small amplitudes, is proportional to the area spanned in parameter space, see figure 4. As a first step the time-resolved pumping current is calculated in lowest order in the tunnel coupling. It is straightforward to relate the pumped current to the dynamics of the average instantaneous charge of the dot $\langle n^{(i)}_t \rangle$ and one finds

$$I_L^{(0)}(t) = -\frac{\Gamma_L(t)}{\Gamma(t)} \frac{d}{dt} \langle n^{(i,0)}_t \rangle,$$

where $\langle n^{(i,0)}_t \rangle$ denotes the instantaneous occupation in lowest order in $\Gamma$. This suggests the following interpretation. As the dot occupation in lowest order, $\langle n^{(i)}_t \rangle$, is constant in time by varying the pumping parameters the charge moves in and out of the quantum dot generating a current from/into the leads. Note that one of the pumping parameters must be the level position since $\langle n^{(c,0)}_t \rangle$ is given by Boltzmann distributions and is therefore independent of the tunnel coupling strengths. This means that $\frac{d}{dt} \langle n^{(i,0)}_t \rangle = 0$ if the level position is constant. The charge moving in and out of the quantum dot is divided to the left and to the right, depending on the time-dependent relative tunnel couplings $\Gamma_L(t)/\Gamma(t)$.

The next step is to calculate the next-to-leading order correction to the lowest order result shown in equation (12). Also the next-to-leading order can be written in a compact way

$$I_L^{(1)}(t) = -\frac{1}{\Gamma(t)} \frac{d}{dt} \left\{ \langle n^{(i,\text{broad}L)}_t \rangle + \frac{\Gamma_L(t)}{\Gamma(t)} \frac{d}{dt} \langle n^{(i,\text{ren})}_t \rangle \right\}.$$  

The first term of equation (13) contains the contribution due to the correction of the average dot occupation induced by lifetime-broadening. It contains a total time derivative, and as parameters are periodically changing in time, it will not lead to a net pumped charge after the full pumping cycle. The second term has the same structure as the zeroth order contribution, equation (12). It can be understood as the correction term induced by the combined influence of charge fluctuations and correlation effects giving rise to a renormalization of the level position, $\varepsilon(t) \to \varepsilon(t) + \sigma(\varepsilon(t), \Gamma(t), U)$. The level renormalization, $\sigma(\varepsilon, \Gamma, U)$, is positive when the level is in the vicinity of the Fermi energy of the leads and negative for $\varepsilon + U$ being close to the Fermi energy. The sign of the renormalization thus indicates whether transitions take place between an empty and singly occupied dot or between a single and doubly occupied dot. The energy level renormalization may be time-dependent via time-dependent tunnel couplings or a time-dependent level. The result is that a finite charge can be pumped by means of level renormalization. This is a pure Coulomb interaction effect, it vanishes for $U \to 0$. We note that correction terms from the renormalization of the tunnel couplings do not contribute in first order in $\Gamma$. For the dc current, different contributions from higher order processes are present at the same time, which makes it challenging to identify them separately in an experiment. For the adiabatically pumped charge, where correction terms associated with a renormalization of the tunnel couplings and level-broadening effects vanish, the situation is distinctively different. Studying adiabatic pumping is, therefore, a convenient tool to access the energy level renormalization. This is most dramatic in the case when the zeroth order pumped current is zero, i.e., when pumping is done by changing both tunnel couplings. In this case, the dominant contribution to the pumped charge is due to time-dependent level renormalization. These results suggest that adiabatic pumping can be used to directly access the level renormalization in quantum dots.

The results for the pumped charge per pumping cycle are shown in figure 5, where $\eta$ is the enclosed surface in parameter space. Modulating the level position and one of the tunnel couplings the pumped charge has a maximum contribution at the resonances, see figure 5(a). Importantly the contribution from sequential tunneling is dominant here. Figure 5(b) shows the pumped charge obtained by modulating the two tunnel couplings. As described above, the pumped charge is due to level renormalization and therefore vanishes for vanishing Coulomb interaction. The sign change between the two contributions at the resonances reflects the opposite sign of the level renormalization for the two resonances.

The method described here is generally formulated and it is therefore extendable to a variety of systems in which Coulomb interaction is important and to which time-dependent fields are applied. Charge and spin pumping through an interacting quantum dot has been studied in the presence of ferromagnetic leads [106]. Pumping through one or two metallic interacting islands with a continuous density of states has been examined in [107]. In [108] the validity of the frequency regime has been extended to faster modulations and in [109] the influence of quantum interference was studied on the pumped charge through one or two quantum dots embedded in an Aharonov–Bohm geometry. Finally this approach has been used to study the capacitive and the relaxation properties of a driven quantum dot figuring as a mesoscopic capacitor in [110].

3.3. Outlook

In this section we have presented results on adiabatic pumping, where interesting effects due to quantum charge fluctuations and finite Coulomb interaction are revealed. The underlying method shown here uses an adiabatic expansion of the master equation based on a real-time diagrammatic technique; this method is applicable whenever a mesoscopic system is exposed to a certain number of adiabatically time-dependent fields. The intriguing results found for systems with a relatively
and tunneling coupling and, therefore, the Kondo effect in quantum dots is only visible for strongly coupled leads. In contrast, the importance of charge fluctuations is controlled by $\Gamma$ and signatures are already significant for $\Gamma \sim T$. Already in section 6, we showed how renormalized level positions can be identified in the adiabatically pumped charge. Experimentally, effects from charge fluctuations have been detected in transmission phase lapses through multi-level quantum dots in Aharonov–Bohm geometries [113, 114]. New light was shed on this long-standing puzzle by the insight that the combined influence of broadening and renormalization effects induced by charge fluctuations is responsible for this effect [115, 116].

The analysis of the signatures of correlation and quantum fluctuation effects requires appropriate methods for their theoretical description. The importance of the development of analytical as well as numerical techniques for their treatment constitutes therefore an important issue. In recent years functional renormalization group (fRG) methods [117] have been established as a new computational tool in the theory of interacting Fermi systems. These methods are particularly powerful in low dimensions. The low energy behavior usually described by an effective field theory can be computed for a concrete microscopic model by solving a differential flow with the energy scale as the flow parameter. Thereby also the non-universal behavior at intermediate scales is obtained. For applications to quantum dots and wires the fRG scheme turned out to be an efficient approach for the description of the single-particle spectral properties and the linear transport through generic models in the presence of moderate correlations. Arbitrary tunneling couplings can be considered, allowing access to the regime of strong coupling between the reservoirs and the quantum system. Due to the flexibility of the microscopic modeling the whole parameter range can be easily explored, and more complex geometries of multi-level quantum dots, as realized in recent experiments, can be treated. In the following sections we will discuss applications of the fRG to the description of strong spin and charge fluctuations in quantum dots, exemplified by the discussion of the Kondo effect and transmission phase lapses in multi-level quantum dots.

4. Quantum fluctuations in linear response

In this section we will discuss the physics of strong quantum fluctuations in combination with correlation effects in quantum dots. We will concentrate on the linear response and static regime, the dependence on finite bias voltage together with the time evolution will be discussed in section 5. One of the most prominent examples of the drastic effects of spin fluctuations in quantum dots is the experimental observation of the Kondo effect [111]. In bulk solids a small amount of magnetic impurities leads to an increased magnetic scattering of the electrons at low temperature, which results in an increased resistance. In quantum dots, the mesoscopic realization of a single spin coupled to two leads displays instead a zero bias peak in the conductance [112]. The particular advantage of quantum dots is that they allow for a very flexible tuning of the parameters and can easily be extended to study the effects of more complex impurities, where orbital Kondo, quantum critical and interference effects may arise. Moreover, additional environmental degrees of freedom in the presence of ferromagnetic or superconducting reservoirs coupled to the quantum dot as well as finite size effects affect the transport properties. For spin fluctuations, the characteristic energy scale $T_c$ is given by the Kondo temperature $T_K$, which, for the special case of a single-level dot with Coulomb energy $U$ and tunneling coupling $\Gamma$, is given by $T_K \sim \sqrt{U \Gamma e^{-\frac{U}{\Gamma}}}$. This scale is exponentially small for large Coulomb energy and, therefore, the Kondo effect in quantum dots is only visible for strongly coupled leads. In contrast, the importance of charge fluctuations is controlled by $\Gamma$ and signatures are already significant for $\Gamma \sim T$. Already in section 6, we showed how renormalized level positions can be identified in the adiabatically pumped charge. Experimentally, effects from charge fluctuations have been detected in transmission phase lapses through multi-level quantum dots in Aharonov–Bohm geometries [113, 114]. New light was shed on this long-standing puzzle by the insight that the combined influence of broadening and renormalization effects induced by charge fluctuations is responsible for this effect [115, 116].

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Figure 5. (a) Pumped charge due to a parameter variation of the dot level and of one of the barriers as a function of the average dot level. The dominant contribution to the pumped charge is due to first order tunneling processes. (b) Pumped charge due to the pure variation of the barrier strengths as a function of the average dot level. The dominant contribution to the pumped charge is due to second order tunneling processes only. In both plots the temperature equals $T = 2\Gamma$.
4.1. The Kondo effect in quantum dots

The appearance of an upturn followed by a low temperature saturation of the resistivity in metals containing diluted magnetic impurities was explained by Kondo in 1960 as an enhanced scattering due to the screening of the local impurity spins by the conduction electrons [118]. The Fermi liquid nature of the ground state [119] and the various theoretical treatments, including Wilson’s numerical renormalization group (NRG) [120], make the Kondo problem one of the best understood many-body phenomena in condensed matter physics, as well as a paradigm for correlated electron physics [121]. In the late 1990s, the prediction by Ng and Lee [122], as well as Glazman and Raikh [112] of the Kondo effect in quantum dots led to a revival of the subject, and has been observed in 2D GaAs/AlGaAs heterostructure quantum dots [111, 123, 124], silicon MOSFETs, as well as in carbon nanotubes [125] and contacted single molecules [126, 127]. Confinement and electrostatic gating enabled investigation of the Kondo effect in an artificial Anderson impurity through electrical transport measurements, characterized by a unitary conductance $2e^2/h$ at zero temperature [128]. Besides providing a paradigm for a variety of physical effects involving strong electronic correlations, the Kondo effect in nanostructures allows for the realization of spintronic device constituents as spin filters or quantum gates. The understanding and control of the transport of spin-polarized currents are of fundamental importance for the advancement of semiconductor spintronic device technology [129].

The coupling of a quantum dot with spin-degenerate levels and local Coulomb interaction to metallic leads gives rise to Kondo physics [121]. A small quantum dot with large level spacing is described by the single-impurity Anderson model (SIAM) in the regime in which only a single spin-degenerate level is relevant. At low temperatures and for sufficiently high barriers the local Coulomb repulsion $U$ leads to a broad resonance plateau in the linear conductance $G$ of such a setup as a function of a gate voltage $V_g$ which linearly shifts the level positions [112, 122, 130–132]. It replaces the Lorentzian of width $\Gamma$ found for non-interacting electrons. On resonance the dot is half-filled implying a local spin-$1/2$ degree of freedom responsible for the Kondo effect [121]. In the limit of large $U \gg \Gamma$ the charge degrees of freedom can be integrated out and the spin physics is described by the Kondo model, which will be discussed in section 5. For the SIAM the zero temperature conductance is proportional to the one-particle spectral weight of the dot at the chemical potential [133]. The appearance of the plateau of width $U$ in the conductance is due to the pinning of the Kondo resonance in the spectral function at the chemical potential for $-\frac{U}{2} < V_g < \frac{U}{2}$ (here $V_g \equiv \epsilon + \frac{U}{2} = 0$ corresponds to the half-filled dot case) [121, 132]. Kondo physics in transport through quantum dots was confirmed experimentally [111, 134], and theoretically using the Bethe ansatz [130, 132] and the NRG technique [120, 135]. However, both methods can hardly be used to study more complex setups. In particular, the extension of the NRG to more complex geometries beyond single-level quantum dot systems [131, 132, 136] is restricted by the increasing computational effort with the number of interacting degrees of freedom. Alternative methods which allow for a systematic investigation are therefore required. Here the fRG approach is proposed to study low temperature transport properties through mesoscopic systems with local Coulomb correlations.

A particular challenge in the description of quantum dots is their distinct behavior on different energy scales, and the appearance of collective phenomena at new energy scales not manifest in the underlying microscopic model. An example of this is the Kondo effect where the interplay of the localized electron spin on the dot and the spins of the lead electrons leads to an exponentially small (in $U/\Gamma$) scale $T_K$. This diversity of scales cannot be captured by straightforward perturbation theory. One tool to cope with such systems is the renormalization group: by treating different energy scales successively, one can often find an efficient description for each one. The fermionic fRG [137] is formulated in terms of an exact hierarchy of coupled flow equations for the vertex functions (effective multi-particle interactions) as the energy scale is lowered. The flow starts directly from the microscopic model, thus including non-universal effects on higher energy scales from the outset, in contrast to effective field theories capturing only the asymptotic behavior. As the cutoff scale is lowered, fluctuations at lower energy scales are successively included in the determination of the effective correlation functions. This allows control of infrared singularities and competing instabilities in an unbiased way. Truncations of the flow equation hierarchy and suitable parametrizations of the frequency and momentum dependence of the vertex functions lead to new approximation schemes, which are devised for moderate renormalized interactions. A comparison with exact results shows that the fRG is remarkably accurate even for sizable interactions.

To exemplify the effect of correlations, we first focus on a quantum dot with spin-degenerate levels. For simplicity only a single level with a local Coulomb repulsion $U$ described by the SIAM [121] is considered, see figure 1. The hybridization of the dot with the leads broadens the levels on the dot by $\Gamma_\alpha = 2\pi T_{\alpha 0}^2 \rho_\alpha$, where $\rho_\alpha$ is the local density of states at the end of lead $\alpha = L, R$ assumed to be independent of frequency here. The energy level is determined by the gate voltage $V_g$. Integrating out the lead degrees of freedom, the bare Green function of the dot [121] is

$$G_0(i\omega) = \frac{1}{i\omega - V_g + i\frac{\Gamma_0}{2}\text{sgn}(\omega)},$$

where $\Gamma = \Gamma_L + \Gamma_R$. By solving the interacting many-body problem a self-energy contribution $\Sigma(i\omega)$ is obtained dressing the bare propagator through the Dyson equation. The fRG is used to provide a self-energy describing the physical properties of the $T = 0$ linear conductance obtained by $G(V_g) = \frac{2}{\pi}\Gamma\rho(0)$ [133] in terms of the dot spectral function $\rho(\omega) = -\frac{1}{\pi} \text{Im} G(\omega + i0^+)$. In order to implement the fRG, an infrared cutoff in the bare propagator is introduced $G_0^\Lambda(i\omega) = G_0(i\omega)\Theta(|\omega| - \Lambda)$. As the cutoff scale $\Lambda$ is gradually lowered, more and more low energy degrees of freedom are included, until finally the original model is recovered for $\Lambda \rightarrow 0$. 

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Changing the cutoff scale leads to an infinite hierarchy of flow equations for the vertex functions. In the static approximation the flow equation for the effective level position $V^\Lambda = V_\Sigma + \Sigma^\Lambda$ reads

$$\partial_\Lambda V^\Lambda = \frac{U^\Lambda V^\Lambda / \pi}{(\Lambda + \frac{1}{2})^2 + (V^\Lambda)^2}$$

with the initial condition $V^\Lambda=\infty = V_\Sigma$ [138]. In first approximation the two-particle vertex $U^\Lambda \equiv U^\Lambda=\infty = U$ equals the bare Coulomb interaction. At the end of the flow, the renormalized potential $V = V^\Lambda=0$ determines the conductance $G(V_\Sigma) = 2e^2/h \arctan(V_\Sigma / \Lambda_1)$ [121, 132]. Results for different values of $U/\Gamma$ are shown in figure 6, together with the occupation of the dot.

For $\Gamma \ll U$ the resonance exhibits a plateau [132]. In this region the occupation is close to 1 while it sharply rises/drops to 2/0 to the left/right of the plateau. For asymmetric barriers the resonance height is reduced to $2e^2/h \arctan(2V_\Sigma / \Lambda_1)$ [121, 132]. Focusing on strong couplings $U \gg \Gamma$, the solution of the above flow equation $V \simeq V_\Sigma e^{-\pi \delta}$ describes the exponential pinning of the spectral weight at the chemical potential for small $|V_\Sigma|$ and the sharp crossover for a $V$ of order $U$. While already the first order in the flow equation hierarchy captures the correct physical behavior, the inclusion of the renormalization of the two-particle vertex improves the quantitative accuracy of the results. For details on the parametrization and extensions including dynamical properties we refer to [138–142]. In the presence of a magnetic field the Kondo resonance splits into two peaks with a dip in $G(V_\Sigma)$ at $V_\Sigma = 0$, providing a definition of the Kondo scale as the magnetic field required to suppress the total conductance to one half of the unitary limit. For the single dot at $T = 0$ the conductance and transmission phases are related by a generalized Friedel sum rule to the dot occupancy [121] by $G = 2e^2/h \sin^2(\alpha/2)(n)$ and $\alpha = \pi(n)$. For gate voltages within the conductance plateau the dot filling is 1 and the phase is $\pi/2$. The description of transmission phases for more complex setups will be discussed in section 4.2.

4.2. Mesoscopic to universal crossover of the transmission phase of multi-level quantum dots

The following application to a multi-level quantum dot illustrates the strength of the fRG approach, the flexibility and simple implementation, in the description of the intriguing phase lapse behavior observed in experiments by the group of Heiblum at the Weizman Institute [113, 114]. The transmission amplitude and phase $T = |T|e^{\alpha}$ of electrons passing through a quantum dot embedded in an Aharonov–Bohm geometry showed a series of peaks in $|T|$ as a function of a plunger gate voltage $V_\Sigma$ shifting the dot’s single-particle energy levels. Across these Coulomb blockade peaks $\alpha(V_\Sigma)$ continuously increased by $\pi$, as expected for Breit–Wigner resonances. In the valleys the transmission phase revealed “universal” jumps by $\pi$ for large dot occupation numbers. In contrast, for small dot fillings the appearance of a phase lapse depends on the dot parameters in the ‘mesoscopic’ regime. From the theoretical side, the observed behavior is captured by an fRG computation of the transport through a multi-level quantum dot with local Coulomb interactions [115, 116]. An essential aspect in the description of the generic gate voltage dependence of the transmission lies in the feasibility of a systematic analysis of the whole parameter space. The interaction is taken into account approximately, but a comparison to numerically exact NRG data for special parameters proves the fRG results to be reliable as long as the interaction parameter and the number of almost degenerate levels do not become too large simultaneously. The results are shown in figure 7, for spin-degenerate levels we refer to [116].

Assuming level spacings of the order of the level width or smaller for large dot fillings (similar to a hydrogen atom with
understanding of the charging of a narrow level coupled to a
and emptied via the narrow levels [144, 145]. The detailed
also in the dot level occupancies, the broad level being filled
figure 7). The mechanism leading to the phase lapses reflects
lapses appear in resonance phenomena (see the lower panels in
figure 7). For several almost degenerate levels in the ‘universal’
level with an interaction-dependent scaling function [147].

4.3. Outlook

The described fRG approach presents a reliable and promising
tool for the investigation of correlation and quantum fluctuation
effects in quantum dots, the computational effort being comparable
to a mean-field calculation. In contrast to the latter the fRG does not lead to unphysical artifacts. Besides
the Kondo effect and transmission phases in multi-level dots, as explained above, also the competition between the Kondo
effect and interference phenomena in more complex quantum
dot systems can be described [138, 148–151]. Furthermore, correlation effects for a large number of single-particle levels
can be discussed up to the limit of one-dimensional quantum
wires. These applications involve inhomogeneous Luttinger
liquids [152–155], which will be discussed in section 6, as
well as the influence of Luttinger liquid leads coupled to a
quantum dot [156]. Challenging extensions and theoretical
developments of the fRG involve the analysis of inelastic processes to describe dynamical properties, and the inclusion
of higher order contributions to access the physics at large
$U/\Gamma$. A fundamental question concerns the combination with the Keldysh formalism to describe non-equilibrium problems addressed in section 5.

5. Quantum fluctuations in non-equilibrium and time
evolution

In this section we discuss quantum fluctuations in quantum
dots or molecules in the presence of a finite bias voltage
together with the time evolution into the stationary state. As in
section 4 the quantum fluctuations are induced by the coupling
to the reservoirs. If the maximum of temperature $T$ and the
distance to resonances $\delta$ decreases, correlation effects from the
Coulomb interaction lead to an increased renormalization of the
couplings and the excitation energies $h_i$ of the quantum dot
(like e.g. magnetic field, level spacing, single-particle energy, etc). Renormalization group (RG) methods, which expand
systematically in the reservoir–system coupling (in contrast to the
fRG method described in the previous section 4, where an expansion in the Coulomb interaction is performed), show
that these renormalizations are typically logarithmic or power
laws. In non-equilibrium or for the time evolution, the voltage
$V$ and the inverse timescale $1/\Gamma$ are two new energy scales,
which can cut off the RG flow. New physical phenomena
emerge, which we will illustrate in this section by considering
very basic two-level quantum dots, like two spin states (Kondo
model) or two charge states (interacting resonant level model
(IRLM)). Conceptually, one has to distinguish between the two
different regimes of strong and weak quantum fluctuations. In
strong coupling, an expansion in the renormalized coupling is
no longer possible, and, except for the case of strong charge
fluctuations in the IRLM or for the case of moderate Coulomb
interactions (which can be treated with fRG methods), the non-
equilibrium case remains a fundamental yet unsolved problem.
Especially for the case of strong spin fluctuations, represented
by the non-equilibrium Kondo model or the non-equilibrium
single impurity Anderson model, an analytical or numerical
solution is still lacking. Even very basic questions, like e.g. the splitting of the Kondo peak in the spectral density by
a finite bias voltage, are not yet clarified. In weak coupling,
where a controlled expansion in the renormalized coupling is
possible, many results are already known. The simplest
ansatz is to take the bare perturbation theory, as explained
in section 2, use a Lorentzian broadening of the energy
conservation (induced by relaxation and decoherence rates),
and replace the bare couplings and excitation energies by the
renormalized ones obtained from standard equilibrium poor
man scaling (PMS) RG equations cut off by the maximum
$\Lambda_c = \max(T, h_i, V, 1/\Gamma)$ of all physical energy scales.
However, it turns out that this approach is not sufficient. In
contrast to temperature, the bias voltage is not an infrared
cutoff since it tunes the distance to resonances, where new
physical processes, like e.g. cotunneling (see section 2), can
occur. Therefore, the correct cutoff is not the voltage but the
distance $\delta_i$ to resonances, where RG enhanced contributions
occur. Furthermore, at resonance $\delta_i = 0$, these contributions are cut off by relaxation and decoherence rates $\Gamma_i$, i.e. one
should use the energy scale $|\delta_i + i\Gamma_i|$ as the cutoff parameter for the couplings. An RG approach has to be developed
to reveal how the various cutoff parameters influence the
couplings. It turns out that the rates $\Gamma_i$ are transport rates,
which have to be determined from a kinetic (or quantum
Boltzmann) equation, in contrast to rates describing the decay of a local wave function into a continuum. Therefore, the
RG has to be combined with kinetic equations. The transport
rates $\Gamma_i$ depend also on voltage and are important parameters
to prevent the system from approaching the strong coupling
regime for voltages larger than the strong coupling scale $T_c$. Furthermore, it turns out that terms can occur in the renormalized perturbation theory which are not present in the bare one, like e.g. the renormalization of the magnetic field in linear order in the coupling for the Kondo model (similar effects can also happen in higher orders for generic models). For the time evolution, these terms are of particular interest in the short-time limit, since in this regime they are cut off by the inverse timescale $1/t$ and lead to universal time evolution. At large times, non-Markovian parts of the dissipative kernel of the kinetic equation lead to many other interesting effects. Among them are unexpected oscillation frequencies involving the voltage, unexpected decay rates, power law behavior, and a different cutoff behavior at resonances compared to stationary quantities. In particular, for metallic reservoirs with a constant density of states, it can be shown that all local physical observables decay exponentially accompanied possibly by power law behavior. Since all of these effects occur already for very basic two-level systems, it has to be expected in the future that many more interesting effects will be found with respect to the physics of quantum fluctuations in non-equilibrium.

From a technical point of view the description of correlated quantum dots in the presence of quantum fluctuations in non-equilibrium is a very challenging problem and a playground for the development of new analytical and numerical methods. Concerning numerical methods, numerical renormalization group (NRG) with scattering waves [157], time-dependent NRG (TD-NRG) [68, 158–160], time-dependent density matrix renormalization group (TD-DMRG) [161, 161–165], quantum Monte Carlo (QMC) with complex chemical potentials [166], QMC in non-equilibrium [167–169], and iterative path-integral approaches (ISPI) [170] have been developed. However, the efficiency of these methods is often not satisfactory in the regime of either strong Coulomb interaction, large bias voltage, or long times. Exact analytic solutions exist for some special cases [162, 171–174] and scattering Bethe ansatz methods have been applied to the IRLM [175]. Other analytical methods are mainly based on RG approaches. Besides conventional PMS [176], improved frequency-dependent RG schemes [177–179], flow equations [180], and iRG methods [141, 142, 181–184] have been used. Whereas the latter expands systematically in the Coulomb interaction, we will introduce in this section a formally exact RG method, which expands systematically in the reservoir–dot coupling. It uses the perturbation theory described in section 2 and allows for a direct calculation of the kernel of the kinetic equation (4) in Liouville space, the so-called real-time RG method (RTRG) [185–187], which recently has been technically optimized and formulated in pure frequency space (RTRG-FS) [5]. The method has been successfully applied to calculate analytically all static and dynamical aspects of weak spin fluctuations for the anisotropic Kondo model at finite magnetic field [188–190] and strong charge fluctuations for the IRLM [191]. Many aspects of these models can also be derived with other methods, which will be mentioned in the following at the appropriate places.

5.1. Real-time RG in frequency space (RTRG-FS)

The aim of RTRG-FS is to combine the diagrammatic expansion in Liouville space, as described in section 2, with RG to resum systematically infrared divergences. We consider a time translational invariant system and use as an initial condition at $t_0 = 0$ that the dot and the reservoirs are decoupled. The kinetic equation (4) can be written in Laplace space as

$$\tilde{\rho}(z) = \frac{i}{z - L_{\text{eff}}^\text{D}(z)} p(t = 0),$$  \hspace{1cm} (16)

where $\tilde{\rho}(z) = \int_0^\infty dt e^{zt} p(t)$ is the reduced density matrix of the dot in Laplace space and $L_{\text{eff}}^\text{D}(z) = L_D + \tilde{\Sigma}(z)$, where $\tilde{\Sigma}(z)$ denotes the Laplace transform of the kernel $\Sigma(t - t') = \Sigma(t, t')$. The effective dot Liouville operator $L_{\text{eff}}^\text{D}(z)$ contains all reservoir degrees of freedom and is of dissipative nature. The qualitative dynamics can be analyzed from the analytic structure of $\tilde{\rho}(z)$, which is an analytic function in the upper half of the complex plane with poles and branch cuts only in the lower half. The single poles located at $z_{\text{eff}}^i = h_i - i\Gamma_i$ correspond to exponential decay with oscillation frequencies $h_i$ and decay rates $\Gamma_i$. From the pole at $z = 0$, the stationary state can be obtained via the solution of $L_{\text{eff}}^\text{D}(i\delta_0) p = 0$. Due to non-Markovian terms arising from the $z$-dependence of the Liouvillian, the weight of these poles is changed by $Z$-factors and branch cuts can occur. For a constant density of states in the leads (normal metallic case) it can be shown generically [190, 191] that the $Z$-factor leads to universal short-time behavior, whereas the branch cuts give rise to an exponential decay $\sim e^{-z_{\text{eff}}^i t}$ accompanied by power law behavior. $z_{\text{eff}}^i$ denotes the position of the branching point, which is shifted from some pole position by multiples of the electrochemical potentials of the reservoirs leading to oscillation frequencies involving the voltage. From the point of view of error correction schemes in quantum information processing it is quite important to understand these corrections to Markovian behavior [192–194].

It is unique to the RTRG-FS method that it provides direct access to the important quantity $L_{\text{eff}}^\text{D}(z)$. By systematically integrating out the energy scales of the reservoirs step by step, a formally exact RG equation can be derived for $L_{\text{eff}}^\text{D}(z)_{\Lambda}$, where all reservoir energy scales beyond $\Lambda$ are included. This RG equation is coupled to other RG equations for the couplings. Similar schemes can be developed for the calculation of the transport current [5] and correlation functions [189]. All RG equations involve resolvents similar to the one occurring in (16) where $z$ is replaced by $\Lambda$ together with other physical energy scales. As a consequence, it can be shown that, besides temperature, each term of the RG equation has a specific cutoff scale $\Lambda_i$, which is generically of the form

$$\Lambda_i = |E + \sum_j n_j \mu_{\alpha j} - h_i + i\Gamma_i| \equiv |\delta_i + i\Gamma_i|,$$  \hspace{1cm} (17)

Here, $E$ is the real part of the Laplace variable, $n_j$ are integer numbers, and $\mu_{\alpha j}$ denotes the electrochemical potential of reservoir $\alpha$. It shows that the cutoff scale is given by the distance $\delta$ to resonances. Furthermore, it provides the
generic proof that, at resonance $\delta_i = 0$, the cutoff scale is
given by the corresponding rate $\Gamma_i$. This issue was under
debate for some time because it was speculated that electrons
tunneling in and out via the same reservoir correspond to
low energy processes, which could possibly lead to a strong
coupling fixed point even in the presence of a finite bias
voltage [195]. However, it was argued that voltage-induced
decay rates prevent the system from approaching the strong
coupling regime [177, 196, 197]. The microscopic inclusion
of decay rates as cutoff scales into non-equilibrium RG
methods was achieved within RTRG [185–187], flow equation
methods [180], and RTRG-FS [5].

5.2. Applications

The two models used to illustrate the basic physics of spin and
charge fluctuations are sketched in figure 8. One model is the
Kondo model at finite magnetic field $h$ already discussed in
section 4, where a spin-$1/2$ couple via anisotropic exchange
couplings $J_{i/\perp}$ to the spins of two reservoirs. We have
assumed a symmetric coupling to the leads and note that during
the exchange it is also allowed that a particle is transferred
between the reservoirs. The model results from the Coulomb
blockade regime of a quantum dot with one level, where charge
fluctuations are frozen out and only the spin can fluctuate. This
leads to an effective band width of the reservoirs of the order of
the charging energy $U$. Anisotropic exchange couplings can
be realized for a molecular magnet, see section 2. The other
model is the IRLM, where the quantum dot consists of a
single spinless energy level at position $\epsilon$. The dot interacts
with the reservoirs via tunneling processes, which are characterized
by the tunneling rates $\Gamma_{a} = 2\pi\rho_{a}|T_{0a}|^{2}$, with $a = L, R$.
In addition, there is a Coulomb interaction $u_{\alpha}$ between the first
site of the reservoir leads and the quantum dot, which are
characterized by the dimensionless parameter $U_{\alpha} = \rho_{\alpha} u_{\alpha}$.
In the following we denote the bare parameters by a super-
index ($0$).

The two models have in common that due to spin/charge
conservation the effective Liouvillian has the same matrix
structure. There are three non-zero poles of the resolvent (16)
at $z_{p}^{1} = -i\Gamma_{1}$ and $z_{p}^{2} = \pm h - i\Gamma_{2}$. For the Kondo model, $\Gamma_{1,2}$
corresponds to the spin relaxation/decoherence rate and $h$ is the
renormalized magnetic field. For the IRLM, $\Gamma_{1}$ is the charge
relaxation rate, $\Gamma_{2}$ describes the broadening of the local level,
and $h = \epsilon$ is the renormalized level position. Denoting the two
eigenstates of the dot by $\pm \uparrow$, $\downarrow \equiv 1, 0$, the matrix elements
$L_{ii}^{\alpha\beta}(0\uparrow,\downarrow) = -i\epsilon_{\alpha} W^{-1}$ of the Liouvillian involve the rates
$W$ for the process $s \rightarrow s$. The stationary occupations
are given by $p_{s} = W^{s}/W$, which contain already most of
the interesting non-equilibrium physics. Similar rates can be
defined to calculate the current. We consider temperature $T$
from now on to reveal the physics of quantum fluctuations.

5.2.1. Kondo model. We consider first the stationary case
in the weak coupling regime $\Lambda_{c} = \text{max}[V, h] \gg T_{K}$, where
the Kondo temperature $T_{K} \equiv T_{c}$ is the strong coupling scale
for spin fluctuations. The renormalized couplings from PMS
cut off at the scale $\Lambda_{c}$ are denoted by $J_{i/\perp}$. For the isotropic
case, they are explicitly given by $J = (1/2) \ln(\Lambda_{c}/T_{K})$, with
$T_{K} \sim D^{1/2} e^{-1/(2D)}$ ($D \sim U$ denotes the effective band
width of the reservoirs).

In lowest order in $J_{i/\perp}$, one obtains a ‘golden rule’ like
expression for the rates [188]

$$W = \frac{\pi}{2} J_{z}^{2} [4h\delta_{--} + (V-h)\theta_{I_{z}}(V-h)],$$

where $h, V > 0$ and $\theta(\omega) = 1/2 + (1/\pi) \arctan(\omega/\Gamma)$ is
a step function broadened by $\Gamma$. The latter corresponds to
a Lorentzian broadening of the energy conservation law by
quantum fluctuations. The spin relaxation/decoherence rates
and the renormalized magnetic field are given by

$$\Gamma_{1} = \frac{\pi}{2} J_{z}^{2} [4h + (V-h)\theta_{I_{z}}(V-h)],$$

$$\Gamma_{2} = \frac{1}{2} \Gamma_{1} + \frac{\pi}{2} V J_{z}^{2},$$

where $\Gamma_{12} = \Gamma_{1} - \Gamma_{2}$ and

$$\begin{align*}
h &= (1 - J_{z} + J_{z}^{(0)}h^{(0)})/h^{(0)} - h J_{z}^{2} \ln \frac{\Lambda_{c}}{|h + i\Gamma_{12}|} \\
&\quad + \frac{1}{2} (V-h) J_{z}^{2} \ln \frac{\Lambda_{c}}{|V-h + i\Gamma_{12}|},
\end{align*}$$

where unimportant terms $\sim O(J^{2})$ without a logarithm have
been left out for $h$.

The expression for the rates $W$ up to $O(J^{2})$ can be
interpreted as follows. At high energies $\Lambda > \Lambda_{c}$, the voltage
and magnetic field are not relevant and the exchange couplings
are renormalized according to the PMS equations. This leads to
an effective band width $\Lambda_{c}$ for the reservoirs with renormalized
exchange couplings cut off at $\Lambda_{c}$. The precise value of $\Lambda_{c}$ is not important since the renormalization is logarithmic.
After this step one uses lowest order perturbation theory with
broadened energy conservation leading to the result (18) for

Figure 8. Two fundamental quantum dot models. (a) is the Kondo model, a spin-$1/2$ coupled via exchange couplings $J_{i/\perp}$ to two reservoirs. (b) is the IRLM, a spinless 1-level quantum dot coupled via tunneling rates $\Gamma_{L,R}$ and Coulomb couplings $U_{L,R}$ to two reservoirs. The electrochemical potentials are given by $\mu_{L,R} = \pm V/2$. 

the rates. However, this interpretation does not work for the rates in \( O(J^3) \) and it fails for the renormalized magnetic field \( h \) already in \( O(J) \) and \( O(J^2) \). We see that \( h \) contains a term linear in \( J \), which does not occur in perturbation theory, and was also discussed e.g. in \[198\]. It shows that it is generically not possible to calculate coefficients of certain orders in the renormalized couplings by comparing with bare perturbation theory. In addition, we find logarithmic corrections in \( O(J^2) \).

As was already discussed generically by equation (17), they occur at resonances \( h = 0 \) or \( V = h \). As was emphasized in \[176\], they are of \( O(J^2) \) at resonance and remain a perturbative correction for \( J \ll 1 \). They occur because the exchange couplings are not completely cut off by \( \Lambda_c \) but partially by smaller cutoff scales. The calculation of the prefactor of the logarithmic terms is quite subtle and follows from the structure of the RG equations, where each term has its own cutoff scale \( \Lambda_c \), see equation (17). Similar logarithmic corrections occur in \( O(J^3) \) for the rates and for \( \Gamma_{1/2} \). It turns out that the logarithmic terms for \( h \) and \( \Gamma_{1/2} \) contain an unexpected cutoff scale \( \Lambda_c = \Gamma_1 - \Gamma_2 \). This is generic for all quantities entering the time evolution. It occurs because the pole positions of the resolvent (16) have to be calculated self-consistently.

Logarithmic enhancements have been studied for the magnetic susceptibility and for the conductance also in \[178, 179, 199\], using slave particle and flow equation methods. The logarithmic terms in the magnetic field have first been calculated in \[188\] using RTRG-FS. They lead to a suppression of the renormalized g-factor as a function of \( h/V \) at \( h = 0 \) and \( h = V \), i.e. a non-equilibrium induced effect, see figure 9. Experimentally, it is proposed to be measured by using a three-lead setup, where the third lead is a weakly coupled probe lead \[188\]. A similar interesting non-equilibrium effect occurs for the logarithmic enhancement of the magnetic susceptibility \( \chi(h/V) \) at \( h = 0 \), as first proposed in \[178\]. All spin-spin correlation functions were calculated analytically in \[189\] using RTRG-FS. Here, the frequency variable of the correlation function enters equation (17) as well. At resonances, kink structures and logarithmic terms were found for the imaginary and real part of response functions, respectively.

In \[190\], the time evolution has been calculated from the inverse Laplace transform of equation (16). In the short-time limit \( t \ll \Lambda_c^{-1} \), the PMS equations are cut off by the energy scale \( 1/t \) leading to time-dependent exchange couplings \( J_{1/2} \).

In \( O(J) \) only the \( Z \)-factor from the linear \( z \)-dependence of \( L_{\text{eff}}(z) \) is important. In terms of the Pauli matrices \( \sigma \), the local density matrix can be written as \( \rho(t) = \frac{1}{2} + \langle S \rangle(t) \sigma \) with

\[
\langle S \rangle(t) = \frac{1}{2} \left( 1 - 2 \left( J_{1/2} - J_{1/2}^{(0)} \right) \right) \langle S \rangle(0).
\]

Inserting the solution of the PMS equations, one obtains universal logarithmic (power law) time evolution for the isotropic (anisotropic) case. A similar result has been found for the longitudinal spin dynamics in \[200, 201\] for the special case of the ferromagnetic Kondo model, which was also confirmed by the experimental results in \[190\] it was also shown that the short-time behavior of the conductance can be calculated from the golden rule expression and replacing the exchange couplings by \( J_{1/2} \).

In the long-time limit \( t \gg \Lambda_c^{-1} \), the cutoff scale for the PMS couplings is given by \( \Lambda_c \). In leading (Markovian) order one obtains the usual exponential behavior from the single poles of the resolvent (16), where the longitudinal/transverse spin decays with \( \Gamma_{1/2} \) and the transverse spin oscillates with \( h \). Interesting non-Markovian corrections occur in \( O(J^2) \), where logarithmic contributions \( \sim J^2 (z - z_0) \ln(\Lambda_c/(z - z_0)) \) in \( L_{\text{eff}}(z) \) lead to branch cuts. The branching point of the logarithm is generically given by \( z_b = z_0 + nV \), i.e. is shifted by multiples of the voltage from some pole position. As a consequence, an exponential behavior \( \sim J^2 e^{-t/\Gamma} e^{-nV} \) is obtained, accompanied by power law behavior \( \sim 1/t^2 \) from the branch cut integral. The result explains why the voltage occurs generically in the oscillation frequency, which is consistent with exact solutions at special Thouless points of two-level systems \[173\]. Furthermore, it shows that all terms decay exponentially but with unexpected decay rates and oscillation frequencies relative to the Markovian terms. This is illustrated in figure 10 by the time evolution of the transverse spin for the strongly anisotropic case \( \Gamma_2 \gg \Gamma_1 \), which is typical for molecular magnets. For short times, one obtains the expected result of oscillations with the magnetic field and decay with the spin decoherence rate \( \Gamma_2 \). These terms decay quickly for large decoherence rate. For longer times, a crossover is obtained to an oscillation with the voltage \( V \) and a decay with the opposite spin relaxation rate \( \Gamma_1 \). Finally, we mention that close to resonances, i.e. for \( |\delta| \ll \Lambda_c, 1/t \), with \( |\delta| = |V - h|, h \), logarithmic terms \( \sim J^2 |\delta| \ln(|\delta + i\Gamma_{1/2}|) \) occur for the time evolution of the transverse spin. Similar to the logarithmic terms in \( h \) and \( \Gamma_{1/2} \), they are cut off by \( \Gamma_{1/2} = \Gamma_1 - \Gamma_2 \).

5.2.2. Interacting resonant level model. For the IRLM the PMS equations lead to a renormalized tunneling rate, given by the power law \( \Gamma_\alpha = \Gamma_\alpha^{(0)}(D/A)^{\alpha} \), with exponent \( \alpha_d = 2U_\alpha - \sum_\beta U_\beta^{(0)} \). Cutting off this scale at \( T_c = \Gamma_1 + \Gamma_k \), one defines self-consistently the strong coupling scale \( T_c \) for the importance of charge fluctuations. The level position \( \epsilon \equiv \epsilon^{(0)} \) and the Coulomb interaction \( U_\alpha \equiv U_\alpha^{(0)} \) remain unrenormalized \[202\]. In lowest order in \( \Gamma_\alpha \), one obtains a golden rule like expression for the rates

\[
W^\alpha_s = \sum_u W^\alpha_{su}, \quad W^\alpha_{su} = \Gamma_\alpha \theta_{\text{T}_s}(s(\mu_u - \epsilon)), \quad (22)
\]
where \( \mu_{\alpha, R} = \pm V/2 \) are the electrochemical potentials of the leads. The energy broadening is given by \( \Gamma_2 = \Gamma_1/2 \) with the charge relaxation rate \( \Gamma_1 = \sum_a \Gamma_a \). Since a single tunneling process changes the charge, there is a unique cutoff parameter \( \Lambda_\alpha^c = |\mu_\alpha - \epsilon + i\Gamma_2| \) for the PMS tunneling rates. This cutoff scale will be used in the following to define \( \Gamma_\alpha \). For \( |\mu_\alpha - \epsilon| \sim T_c \), we are in the regime of strong charge fluctuations. We note the essential difference to the Kondo model, where several cutoff parameters are relevant. The current in lead \( \alpha \) can be calculated from the rate equation \( J_\alpha = \frac{\pi}{\hbar} 2(\mu_\alpha - \mu_\alpha - \epsilon) \). The dips have their origin in the oscillations of \( S(x) \).

The status of the field of quantum fluctuations in non-equilibrium for strongly correlated quantum dots is that powerful techniques have been developed to study the weak coupling limit in a controlled way. Elementary models of spin and charge fluctuations are well understood in this limit and the methods can now be applied to more complex quantum dot models, such as discussed in section 2.2. However, two fundamental issues are still open. First, the case of strong spin fluctuations, represented in its most elementary form by the isotropic Kondo model in the limit \( \max(T, h, V) \sim T_k \), is one of the most fundamental unsolved problems. The case of strong charge fluctuations at resonances has so far only been understood for the IRLM, where it seems that the broadening of the level together with a PMS renormalization of the tunneling couplings captures the essential physics. Whether this holds also for more complicated models has to be tested in the future. Second, most of the methods used to describe non-equilibrium properties of quantum dots are parametrized by the many-body eigenstates of the isolated quantum dot. Therefore, they can not be extended easily to larger systems like multi-level quantum dots or quantum wires. Two exceptions are the flow equation method [180] and non-equilibrium fRG methods [141, 142, 181–184]. In fRG, as already described in section 4, a perturbative expansion in the Coulomb interaction is used and the vertices are parametrized in terms of single-particle levels. Therefore, although fRG is restricted to the regime of moderate Coulomb interactions, the potential of the method lies particularly in the possibility to treat multi-level quantum dots and quantum wires. Preliminary studies have tested the fRG for the IRLM and the single-impurity
Anderson model (SIAM). For the IRLM, a static version was used, where the frequency dependence of the Coulomb vertex was neglected [184, 191]. An agreement was found with RTRG-FS and TD-DMRG [162]. Recently, a dynamic scheme with frequency-dependent two-particle vertices has been developed and was used to analyze the non-equilibrium SIAM in the regime of strong spin fluctuations [141, 142]. A good agreement with TD-DMRG, ISPI and QMC results was obtained for moderate Coulomb interactions [204]. As will be described in the next section, the static version of non-equilibrium IRG has also been applied to quantum wires and it is a challenge for future research to generalize the dynamic IRG scheme to quantum wires as well.

6. Correlation effects in quantum wires

6.1. An introduction to Luttinger liquid physics

A different type of correlation physics than discussed so far is found in (quasi) one-dimensional (1d) quantum wires at low temperatures. By this we mean electron systems confined in two spatial directions such that only the lowest 1d subband is occupied. Compared to the quantum dots considered above they contain many correlated degrees of freedom and the single-particle level spacing of the wire becomes the smallest energy scale (‘almost’ continuous spectrum; see below). The dominant quantum fluctuations are now driven by the interaction $U$ itself. In fact, below we will first discuss isolated, translationally invariant wires. Still, as discussed in the main part of this section, the physics becomes even more interesting if the coupling to leads (or the coupling between two wires) is included. Quantum wires are realized in single-walled carbon nanotubes [205, 206], specifically designed semiconductor heterostructures [207, 208], and in atom chains which form on certain surfaces [209]. A different class of quasi 1d systems are highly anisotropic (chain-like) bulk materials (for a recent review see e.g. [210]). Compared to the first type of systems it is less clear how a single wire of this class can be incorporated as an element in an electronic transport nanodevice and they will thus play a minor role in the present discussion.

Although being far from experimentally realizable, translationally invariant models of 1d electrons with sizable two-particle interaction (Coulomb repulsion) were already studied in the 1950s and 1960s. Exactly solvable models were constructed leading to a rather quick (compared to higher-dimensional correlated systems) gain in the understanding of correlation effects. Tomonaga showed that all excitations of a 1d correlated electron system at (asymptotically) low energies are collective in nature (‘plasmons’) rather than single-particle-like as in the three-dimensional counterpart (Landau quasi-particles) [211]. In particular, collective spin and charge density excitations travel with different speed. Quite often this difference in velocities of collective excitations is mistaken as being characteristic for 1d systems although it might also occur in higher-dimensional Fermi liquids. Only when adding, that no additional quasi-particle excitations are possible spin-charge separation becomes a unique feature of 1d correlated electrons. As a second (related) interaction effect certain correlation functions display power law behavior as a function of energy $\omega$ and momentum $k$ as first shown for the single-particle momentum distribution function $n(k)$ by Luttinger [212] and Mattis and Lieb [213]. Using the concepts of statistical physics one can view 1d chains of electrons as being (quantum) critical, although the exponents depend on the details of the underlying single-particle model (band structure, band filling) as well as on the interaction strength. Sólyom [214] and Haldane [215] showed using renormalization group (RG) arguments that the physics of the two exactly solvable models—being quite similar they are nowadays considered as a single model, the Tomonaga–Luttinger model—is generic for 1d interacting electrons as long as correlations do not drive the system out of the metallic phase (e.g. into a Mott insulator phase). This led to the term Luttinger liquid (LL) physics for the above correlation effects. In particular, Haldane argued that all of the exponents appearing in correlation functions of a spin-rotational invariant system with repulsive two-particle interaction can be expressed in terms of a single model parameter dependent number $K < 1$, with $K = 1$ for non-interacting electrons. Together with the velocities of the charge-and spin-density excitations $v_c$ and $v_s$, $K$ completely determines the low energy physics. For a given microscopic model the strategy to obtain the low energy physics is thus: one has to determine the three Luttinger liquid parameters as functions of the model parameters (ways to achieve this are e.g. discussed in [216]) and plug them into the analytic expressions for correlation functions computed within the Tomonaga–Luttinger model.

Despite the intense effort (see e.g. [205–210]) a commonly accepted experiment which shows LL behavior beyond any doubts is still pending. This has to be contrasted to the situation in quantum dots discussed earlier where the appearance of Kondo physics as a consequence of local correlations has convincingly been shown (see above). The lack of clear cut experiments is partly related to the status of the theory. LL theory only makes predictions for the asymptotic low energy behavior but does not provide the scales on which this sets in. In the construction of the Tomonaga–Luttinger model all microscopic scales as e.g. given by the band curvature [217] and the shape of the two-particle interaction [218] are disregarded. One can imagine that the upper energy scale $T_c$ beyond which LL physics can be found becomes so small that power laws are masked by other scales such as the single-particle level spacing set by the length of the wire. A profound comparison to experiments requires a knowledge of $T_c$ and possible lower bounds for LL physics while methods which allow extraction of these scales for microscopic models are rare. It is thus mandatory to develop new methods which capture LL physics and can directly be applied to microscopic models. Further down, we will return to this issue. We note in passing that the standard ab initio method—density functional theory—cannot be applied as the existing approximation schemes fail when it comes to LL physics.

When using a quantum wire as part of a low temperature nanodevice one has to be alert that correlation effects might alter the electronic properties. To understand this let us first imagine an electron tunneling from the tip of a scanning
tunneling microscope into the bulk of a 1d quantum wire. Using golden rule like arguments it becomes clear that the differential conductance $dI/dV$ computed from the tunneling current $I$ is determined by the product (more precisely the convolution) of the single-particle spectral function $\rho(\omega)$ (the ‘local density of states’) of the wire and the tip. As the latter is constant at small energies, $dI/dV$ displays the LL power law of the wire spectral function $\rho(\omega) \sim |\omega - \mu|^\alpha_{\text{bulk}}$, with the wire’s chemical potential $\mu$ and $\alpha_{\text{bulk}} = (K + K^{-1} - 2)/4$ [216]. The exponent changes to $\alpha_{\text{end}} = (K^{-1} - 1)/2$ if the electron tunnels into the end of a LL as is e.g. the case in end-contacted wires [216]. Evidently, tunneling in and out of either the bulk or the end of a wire plays a crucial role when using it in a nanodevice.

Second, we imagine a local inhomogeneity in the quantum wire. It might either appear (in a difficult to control way) in the process of producing the wire or might intentionally (in a controlled way) be introduced to design a functional device. As examples for the latter one can think of double barriers [219] defining a dot region with energy levels that can be tuned via a backgate (see the discussion on transport through quantum dots) or junctions of several quantum wires [220, 221]. One can easily imagine that because of the collective nature of the excitations in a LL any inhomogeneity with a (single-particle) $2k_F$-backscattering component, with $k_F$ being the Fermi momentum, strongly affects the low energy physics. A first step to substantiate this expectation is to compute the $2k_F$-component of the static density-density-response function $\chi(q)$ of the Tomonaga–Luttinger model [222]. For a non-interacting 1d electron system it diverges logarithmically when $q$ approaches $2k_F$ (Lindhard function). The divergence is enhanced and becomes power law like $\chi(q) \sim |q - 2k_F|^\alpha$ if the interaction is turned on. This shows that the inhomogeneity strongly couples to the system and linear response theory breaks down. This physics was further investigated applying different RG methods to different models [154, 223–226]. These studies show that even a single impurity at low energy scales effectively acts as if the chain is cut at the position of the impurity with open boundary conditions at the two end points. For example, for temperatures $T \to 0$ the linear conductance $G(T)$ across the impurity is suppressed in a power law fashion $G(T) \sim T^{2\alpha_{\text{end}}}$ as can again be understood using Fermi’s golden rule and the power law scaling of the spectral function at the end of a quantum wire (tunneling from end-to-end of two LLs). Remarkably, the exponent is independent of the bare scattering potential. Also a Breit–Wigner resonance of $G$ as a function of the level position (modified by $V_g$) showing up in the double-barrier geometry is strongly altered by the interaction. In the case of a ‘perfect’ resonance with peak conductance $e^2/h$ (per spin), that is for equal left and right barriers (which is experimentally difficult to realize), the resonance width becomes zero, while the resonance completely disappears, that is $G \to 0$ for all $V_g$, in all other cases.

The discussed physics of inhomogeneous LLs can be understood in terms of an effective single-particle problem. Using RG techniques one can show that the interplay of a local inhomogeneity and two-particle correlations leads to an effective oscillatory and slowly decaying ($\sim 1/x$, with the distance $x$ from the inhomogeneity) single-particle scattering potential of range $1/\delta$, where $\delta$ is the largest of the relevant energy scales (e.g. temperature). The wavelength of the potential is set by the (common) chemical potential $\mu$ of the leads. Scattering off this (so-called Wigner–von Neumann potential) leads to the discussed power laws in transport with the exponent given by the amplitude of the potential [154, 224–226].

In the remaining part of this section on coherent transport through quantum wires we describe four examples of recent attempts (by three of the present authors together with varying colleagues) to gain a detailed understanding of the interplay of correlations and local inhomogeneities in systems with different geometries. We believe that our insights should be kept in mind when designing transport setups to search for LL physics. They might also be of relevance when quantum wires (in the above defined sense) are used as elements in future nanoelectronic devices. In this case correlation effects might either be used to enhance the functionality or must be tuned away if they corrupt currents.

In all of the examples we use the functional renormalization group (fRG) method [137] (for a brief review see also [227]) to (approximately) treat the Coulomb interaction. It was already introduced in section 4. Compared to other methods it has the advantages that it (i) can directly be applied to microscopic models (continuum or lattice; flexible modeling), (ii) captures all energy scales (not restricted to the asymptotic low energy regime), and (iii) can be applied to systems with many correlated degrees of freedom. In the present context (iii) means chains of realistic length (in the micrometer range) coupled to effectively non-interacting (Fermi liquid) leads. The aspects (i) and (ii) together assure that $T_F$ and additional energy scales affecting LL physics can be investigated. As mentioned in section 4, the approximations inherent to the fRG approach are justified up to moderate renormalized two-particle interactions. In the implementation (static self-energy) used for quantum wires (i) the LL exponents come out correctly (only) to leading order in the two-particle interaction and (ii) inelastic processes generated by the two-particle interaction are neglected. We will return to the latter in the outlook.

6.2. The role of two-particle backscattering

In 1d systems at low energies and incommensurate fillings only forward (momentum transfer $q \approx 0$) and backward (momentum transfer $q \approx 2k_F$) two-particle scattering is active. All other processes are suppressed due to momentum conservation and phase space restrictions (Fermi surface consists of only two points). The relative importance of the two processes is determined by the (real space) range of the two-particle interaction, which in turn is set by the screening properties of the ‘environment’, e.g. the substrate on which the carbon nanotube is placed. If screening is strong, the two-particle interaction becomes short ranged in real space and backscattering is sizable and vice versa. Backscattering processes involving a spin up and a spin down electron are not part of the Tomonaga–Luttinger model. For a translationally invariant LL it was shown using an RG analysis [214] that...
these processes do not modify the asymptotic low energy physics (they are ‘RG irrelevant’), but that they become irrelevant only on a scale which is *exponentially small* in the bare backscattering strength (they flow to zero only logarithmically). From this one can expect that the LL power laws only occur on exponentially small scales $T_c$ also for inhomogeneous wires. For the *Hubbard model* with open boundaries it was shown that $T_c \sim \exp(-\pi V_F/U)$, where $U$ is the local Coulomb repulsion and $V_F$ the Fermi velocity [228]. In figure 11 this behavior is exemplified for the $T$-dependence of the linear conductance $G$ of a wire with a single impurity described by the *extended Hubbard model* [155]. This lattice model consists of a standard tight-binding chain with nearest-neighbor hopping $t$, a local two-particle interaction $U$, as well as a nearest-neighbor one $U'$. Throughout the rest of this section we use $t$ as the unit of energy (that is we set $t = 1$). For a fixed band filling the relative strength of the forward and backward scattering can be modified by varying the ratio $U'/U$. The considered size of $N = 10^7$ lattice sites corresponds to wires in the micrometer range, which is the typical size of quantum wires available for transport experiments. For $U' = 0$ (strong backscattering) due to logarithmic corrections the conductance increases as a function of decreasing $T$ down to the lowest temperatures in the plot, $T_c$ being smaller than the latter. For increasing nearest-neighbor interactions $U'$ the (relative) importance of backscattering decreases and a suppression of $G(T)$ at low $T$ becomes visible, but in all of the data obtained at quarter-filling $n = 1/2$ the suppression is much less pronounced than what one expects from the asymptotic power law with exponent $2\alpha_{\text{end}}$. By contrast, the suppression is much stronger and follows the expected power law more closely if parameters are chosen such that two-particle backscattering becomes negligible at low $T$, as can be seen from the conductance curve for $n = 3/4$ and $U' = 0.65$ in figure 11. The value of $K$ for these parameters almost coincides with the one for another parameter set in the plot, $n = 1/2$ and $U' = 0.75$, but the behavior of $G(T)$ is completely different. To avoid interference effects discussed next, in the present setup the two leads are coupled ‘adiabatically’ to the wire, such that in the absence of the impurity the conductance would be $2e^2/h$ for all temperatures smaller than the band width [154].

One can conclude that for wires in the micrometer range clear power laws can only be observed if screening is weak, leading to small two-particle backscattering [155, 224–226]. Note that at $T \sim \pi V_F/N$ finite size effects set in (the level structure becomes apparent), as can be seen at the low $T$ end of some of the curves in the figure.

6.3. Overhanging parts

In most transport experiments the wires are *not* end-contacted and the leads do *not* terminate at the contacts. This has to be contrasted to the modeling in which almost exclusively end-to-end contacted wires are considered. It is thus crucial to understand how overhanging parts of the wire and the leads alter the transport characteristics. Before tackling this problem we have to discuss transport through an interacting wire with two contacts—above we only mentioned the single contact case with power law scaling $G(T) \sim T^{\delta_{\text{aad}}}$. One might be tempted to argue, that two contacts can be understood as two resistors in series, which then have to be added leading to $G(T) \sim T^{\delta_{\text{aad}}}$. One might be tempted to argue, that two contacts can be understood as two resistors in series, which then have to be added leading to $G(T) \sim T^{\delta_{\text{aad}}}$. One might be tempted to argue, that two contacts can be understood as two resistors in series, which then have to be added leading to $G(T) \sim T^{\delta_{\text{aad}}}$ also for a (clean) wire with two contacts to semi-infinite leads. Although the result is correct, as can be shown using (coherent) scattering theory [154, 229], this simple argument ignores, that adding of the resistance (of resistors in series) only holds if the transport is *incoherent*, which is not the case at low temperatures as considered here.

We here exemplify the role of overhanging parts by considering overhanging leads. In figure 12 $G(T)$ is shown for a *spinless* tight-binding wire with nearest-neighbor interaction $U'$ coupled to two semi-infinite leads via tunneling hoppings $T_{L,R}$. For spinless models the low energy LL physics can be described by $K$ and the charge velocity $v_c$. Several correlation functions are again characterized by power law scaling. Compared to models with spin the analytic dependence of exponents on $K$ is modified (see e.g. [216]). Three setups with different overhanging parts are considered (see the left inset). As becomes clear each non-zero number of overhanging lattice sites $N_{\text{lead}}^{L,R}$ inflers a new energy scale $\pi v_F/N_{\text{lead}}^{L,R}$ and the simple power law (with exponent $\alpha_{\text{end}}$; dashed line) is divided in subsections separated by extensive crossover regimes. If clearly developed the power laws in the subsections all have exponent $\alpha_{\text{end}}$ (see the inset of figure 12). Overhanging parts of the wire lead to the same effect [230]. There is no reason to believe that adding the spin degree of freedom will change this. The same holds for the two effects discussed next and we thus stick to the spinless lattice model for the rest of this section.

One can conclude that in a generic experimental setup with overhanging parts the power law is much more difficult to observe than suggested by most theoretical studies considering end-to-end coupling.

![Figure 11. Temperature dependence of the linear conductance for the extended Hubbard model with 10⁷ sites and a single site impurity (‘δ-impurity’) of strength $V = 10$, for a Hubbard interaction $U = 1$ and various choices of $U'$; the density is $n = 1/2$, except for the lowest curve, which has been obtained for $n = 3/4$ and $U' = 0.65$ (leading to a very small backscattering interaction); the dashed line is a power law fit for the latter parameter set.](image-url)
6.4. A wire with two contacts and a single impurity

We now exemplify the very rich interference effects which occur when considering coherent linear transport through interacting wires with several inhomogeneities. A simple setup of this class is a wire with two end-to-end tunneling contacts to semi-infinite leads and an additional impurity in its bulk. As mentioned in the last subsection the concept of ‘adding resistances’ cannot be applied in the present context. Using averaging over the (two) scattering phases one can show that for sufficiently large $T$ (of the order $10^{-2}$ in figure 13) the total linear conductance is given by $G \sim G_1^\alpha G_2^\alpha G_3^\alpha$ with the conductances $G_i$ of the individual barriers. For the present setup we have $G_{1/3}(T) \sim T^{\alpha_{\text{end}}}$ and $G_2(T) \sim T^{2\alpha_{\text{end}}}$, leading to $G(T) \sim T^{2\alpha_{\text{end}}}$. As is shown in figure 13 the dashed–dotted curve obtained by phase averaging nicely follows the bare data (solid and dashed line) down to $T_p \approx O(10^{-2})$ and a power law with the expected exponent develops for $T_p \lesssim T \ll D$.

For $T < T_p$ details of the relative energy level structure of the two ‘dots’ defined by the three barriers matter and might even lead to resonances (see the solid line). The ‘dots’ energy level structure is set by the position of the bulk impurity and the solid and dashed line in figure 13 display data obtained for different positions. The example clearly shows that interference due to multiple inhomogeneities can set a lower bound to power law scaling of the conductance.

6.5. Transport at finite bias voltages

As our final example we study transport at finite bias voltages. In particular we consider the steady state current through a clean wire end-to-end coupled to two leads (reservoirs) which are held on different electrochemical potentials $\mu_L$ and $\mu_R$. The respective amplitudes are proportional to $T_{\mu_{LR}}^\alpha(T_{\mu_{LR}}^\alpha + T_{\mu_{LR}}^\beta) = \Gamma_{\mu_{LR}}^2/(\Gamma_L + \Gamma_R)$, with $\Gamma_\alpha \sim T_\alpha$ defined as in the preceding sections. Accordingly, the non-equilibrium local spectral function close to either of the two contacts displays power law suppressions at $\mu_L$ and $\mu_R$. The respective exponents are proportional to the respective couplings $\alpha_{\mu_{LR}} = \Gamma_{\mu_{LR}}^2/(\Gamma_L + \Gamma_R)$. In contrast to the linear response regime the exponents now depend on the strength of the inhomogeneities. This behavior is shown in figure 14. The spectral weight remains finite even at $\mu_{LR}$ as the power laws are cut off by the single-particle level spacing $\sim 1/N$ of the disconnected wire. A transport geometry in which the two power laws show up was proposed in [183].

6.6. Outlook

This short review on correlation effects in 1d quantum wires gives a flavor of the rich and partly surprising physics resulting from the interplay of the two-particle interaction and local (single-particle) inhomogeneities. An important ingredient missing in the current description using the (approximate) fRG for microscopic models are inelastic scattering processes resulting from the (Coulomb) interaction and leading to decoherence and dephasing at intermediate to large energies. We note that they are also only partly kept in the standard modeling which is based on the Tomonaga–Luttinger model. One can expect that inelastic processes set an upper energy
and nearest-neighbor interaction $U$. References FG 723, FG 912, SPP-1243, the NanoSci-ERA, the Ministry of present in this review. This work is supported by the DFG-

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Figure 14. Solid line: local spectral function (close to the left contact) in the steady state of an interacting tight-binding wire of length $N = 2 \times 10^4$ with tunnel couplings $T_L = 0.075$, $T_R = 0.15$, and nearest-neighbor interaction $U = 0.5$ as a function of $\omega$. The voltage is $V = 0.1$. Scale $T_c$ for power law scaling smaller than the maximal scale given by the band width. They will also play a prominent role when further considering non-equilibrium transport. First attempts to include such inelastic two-particle scattering for quantum dots in and out of equilibrium were mentioned in section 4 and are discussed in [139–141].
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