Sources of negative tunneling magnetoresistance in multilevel quantum dots with ferromagnetic contacts
Koller, Sonja; Grifoni, Milena; Paaske, Jens

Published in:
Physical Review B Condensed Matter

DOI:
10.1103/PhysRevB.85.045313

Publication date:
2012

Document Version
Publisher's PDF, also known as Version of record

Citation for published version (APA):
I. INTRODUCTION

Recent transport experiments on quantum dots coupled to ferromagnetic leads have demonstrated the existence of spin-dependent energy level shifts through the observation of negative tunnel magnetoresistance (TMR) effects in the single electron tunneling regime\(^1\) and spin splitting in the Kondo regime.\(^2\)\(^-\)\(^5\) So far qualitatively different approaches for explaining the origin of the underlying shifts coexist. For example, negative TMR data from carbon nanotube (CNT) measurements have been fitted with a model relying on spin-dependent interfacial phase shifts\(^6\) picked up by the wave function during multiple reflections at a spin-active interface.\(^7\) The concept is related to that of spin-mixing conductance\(^8\) and, because it only depends on the properties of the spin-active barrier region, is only weakly gate dependent and present in both interacting and noninteracting systems.\(^9\)

In contrast, experiments on CNTs\(^3\)\(^,\)\(^\,\)\(^5\) and InAs nanowires\(^4\) in the Kondo regime have demonstrated a combination of gate-dependent and gate-independent contributions to the energy level shifts. Both can be explained in terms of charge fluctuations,\(^10\) whereby electron-electron interactions are responsible for the logarithmic gate dependence,\(^11\) while a Stoner splitting of the energy bands of the magnetically polarized leads accounts for the almost gate-independent part.\(^12\)\(^,\)\(^13\)

While the effects of the energy level shifts in the Kondo regime are by now well understood, a thorough understanding of their influence on the TMR phenomenon in interacting quantum dots is still missing. The negative TMR data\(^1\) have been satisfactorily fitted in terms of a generalized Anderson model already including gate-dependent level shifts. In Ref. \(^14\) a reflection Hamiltonian was included to account for reflection processes at the interface in a second-order sequential tunneling theory.

In this work, we specifically address the TMR phenomenon within a diagrammatic approach to the reduced density matrix of the nanosystem\(^15\) and discuss different level-shift-induced mechanisms for negative TMR. Spin-dependent level shifts can originate to the lowest order in the tunneling Hamiltonian (sequential tunneling limit) if off-diagonal elements of the reduced density matrix participate in the dynamics. This is the case for noncollinearly polarized leads\(^14\)\(^,\)\(^16\)\(^,\)\(^17\) but also for collinearly polarized leads for some double-dot setups.\(^18\) Here we consider situations where coherences of the reduced density matrix do not play a crucial role in the dynamics. We rather identify charge fluctuation terms in all orders in the tunneling Hamiltonian which sum up to a Taylor series yielding intrinsic level shifts.

In particular we discuss two types of spin-dependent level shifts: strongly gate-dependent ones deriving from a net difference of the majority and minority density of states at the Fermi level, and largely gate-independent ones caused by a Stoner shift of the majority and minority bands in the leads. Our results are valid for generic multilevel quantum dots and reproduce the results of Ref. \(^12\) in the case of a simple Anderson model.

Finally, we analyze the TMR of a CNT quantum dot and show that, due to the multilevel spectrum, the intrinsic contributions give rise to a marked gate voltage dependence and a TMR which can indeed become negative. Moreover, we show that charge fluctuations to states including bosonic excitations, which are present in CNTs due to the linearity of the noninteracting spectrum, can largely influence the gate voltage dependence of the level shifts.

The paper is structured as follows: Sec. II contains a discussion of possible mechanisms for negative TMR. In Sec. III the model Hamiltonian is elucidated, while in Sec. IV the level shifts are calculated analytically. Results for the conductance and TMR of armchair CNTs are presented in Sec. V, while in Sec. VI conclusions are drawn.

II. MECHANISMS OF NEGATIVE TMR

The basic mechanism underlying a negative TMR is the presence of an effective generalized Zeeman field \(\hbar P/A(V_F)\) accounting for both extrinsic (stray fields, applied magnetic field) and intrinsic sources of spin-dependent level shifts. Here \((P/A)\) refers to contacts with parallel/antiparallel
magnetization (see Fig. 1). For spin-dependent transport experiments, typically two ferromagnetic contacts of different widths [compare Fig. 1(a)] are utilized to obtain a hysteretic polarization behavior when exposing the setup to a sweep of an external magnetic field. Due to remanent magnetization, the polarization of the contacts is inverted only at a reverted field of some hundreds of millitesla in strength, until the magnetically softer of the two contacts switches first to align with the external field. Thus, for some range of the field, the contacts stay in antiparallel configuration. Intuitively, one would expect that the conductance for contacts polarized in parallel \((G_P)\) is larger than the one for the antiparallel case \((G_A)\), yielding positive values of TMR \(= (G_P - G_A)/G_A = 1\). Nevertheless, there are at least two different mechanisms which can lead to a negative TMR, as sketched in Figs. 1(b) and 1(c), respectively. Here we consider quantum dots in the single electron tunneling regime and look at the linear conductance as a function of gate voltage \(V_g\). For sufficiently small couplings the temperature determines the width of the conductance peaks, while the peak position signals at which value of \(V_g\) the many-body ground state configurations with \(N\) and \(N+1\) electrons involved in the transition have the same electrochemical potential: \(\mu(N,V_g) = \mu(N+1,V_g)\).

Because the effective Zeeman field leads to corrections to the energy difference \(E_b - E_a := E_{ba}\) associated with the transition from a many-body state \(a\) to a many-body state \(b\), this in turn modifies the position of the conductance peaks.

First, and most obviously, there can be negative TMR if there is a noticeable offset in the conductance peak positions in parallel and antiparallel configuration [see Fig. 1(b)]. This requires \(|h^P - h^A|\) of the order of the width \(k_BT\) of the conductance peaks. Second, however, negative TMR can also arise for \(|h^P| = |h^A| \approx k_BT\) [see Fig. 1(c)]. This is because the effective magnetic field responsible for the effective Zeeman shift also removes the spin-degeneracy of the ground states by favoring the states with maximum total spin. This situation is illustrated in Fig. 1(d) for the case of a CNT quantum dot, for which we consider the CNT Hamiltonian, Eq. (2). The four-electron shells of the dot are filled sequentially upon sweeping the gate voltage and only a single shell needs to be considered at a time. The case of orbital \((\Delta e = 0)\) and spin degeneracy is illustrated for the \(n\)th shell in Table I. For the \(4n \leftrightarrow 4n+1\) and \(4n+1 \leftrightarrow 4n+2\) transitions and \(h^P/A \neq 0\) only spin-up electrons are required. Hence, due to the larger density of spin-up electrons, peak heights for parallel contact polarization will exceed those for the antiparallel case. On the other hand, for the \(4n+2 \leftrightarrow 4n+3\) and \(4n+3 \leftrightarrow 4n+4\) transitions spin-down electrons have to be transferred through the dot. In this latter case, a configuration with antiparallel contact polarization might be favored, leading to \(G_A > G_P\).

### III. THE MODEL HAMILTONIAN

For a quantitative description we consider the transport setup of Fig. 1. In the limit of weak coupling to the leads it can be described by the total Hamiltonian

\[
\hat{H}_{\text{tot}}^{P/A} = \hat{H} + \hat{H}_{\text{ext}}^{P/A} + \sum_{l=r,d} \hat{H}_l + \hat{H}_F, \tag{1}
\]

where \(\hat{H} = \hat{H}_0 - e\alpha V_g \hat{N}\) comprises the Hamiltonian \(\hat{H}_0\) of the isolated quantum dot and the effects of a gate voltage \(\alpha\) is a conversion factor of the order of one). In the case of an armchair nanotube quantum dot of medium-to-large radius far from half-filling it reads\(^19\)

\[
\hat{H}_0 = \frac{1}{2} E_c \hat{N}^2 + \frac{1}{2} \sum_{r,a} (\varepsilon_0 \hat{N}_{F_r} + \tilde{r} \Delta e) \hat{N}_{F_r} + \hat{H}_B, \tag{2}
\]

where \(\varepsilon_0 = \hbar v_F/\pi L\), with \(v_F\) being the Fermi velocity and \(L\) the CNT length, is the level spacing; \(\Delta e\) is the orbital mismatch; and \(E_c\) is the charging energy. The number of electrons in the orbital band \(F = \pm\) with spin \(\sigma\) is determined by the number operator \(\hat{N}_{F_r}\) and the total number is determined by \(\hat{N} = \sum_{F_r} \hat{N}_{F_r}\). Finally,

\[
\hat{H}_B = \sum_{n \neq 0} \sum_{j=u,s} \varepsilon_{j,n} \hat{a}_{j,n}^\dagger \hat{a}_{j,n} \tag{3}
\]
accounts for bosonic charge, $c$, and spin, $s$, excitations. The energies $\epsilon_{s,n} = n e a / g$ are for $n > 0$ depending on the Coulomb interaction parameter $g \simeq 0.2$, while the others scale only with the level spacing: $\epsilon_{s,\pm|n|} = \epsilon_{s,|n|} = |n|e/a$. None of the bosonic excitations influences the particle number in the single bands and hence one can classify the states of the CNT by a vector $|\mathcal{N}, \mathcal{m}\rangle$, where $\mathcal{N} = (N_{+1}, N_{+\uparrow}, N_{-1}, N_{-\downarrow})$ determines the fermionic configuration and $\mathcal{m}$ determines the bosonic excitations (see, e.g., Ref. 20).

The ground states $|\mathcal{N}, \mathcal{m}\rangle = |0\rangle$ of shell $n$ have $4n$, $4n + 1$, $4n + 2$, and $4n + 3$ electrons and can be characterized in terms of the excess spins in band $\hat{\tilde{\mathcal{E}}}_{l,s} = \epsilon \phi_{l}q$, with $\hat{\tilde{\mathcal{E}}}_{l,s}$ annihilating an electron of energy $\epsilon_{l}q$ and of spin $s$ in lead $l$. The density of states in lead $l$, $D_{l}(\omega)$, is assumed to be constant over an energy range set by a spin-dependent interaction parameter $\Delta$.

It determines the leads polarizations $P_l$ at the Fermi level according to $P_l = (D_l - D_{\uparrow l})/(D_{l\uparrow} + D_{l\downarrow})$.

On the other hand, in order to account for a Stoner splitting $\Delta$, the range of available energies in the leads is given by $-\omega \leq W + \sigma \Delta_{l}/2 \leq W$, where $\sigma$ is the bandwidth at zero Stoner splitting. Finally the perturbative contribution is $\hat{\tilde{\mathcal{H}}} = \hat{\tilde{\mathcal{H}}} + \hat{\tilde{\mathcal{H}}} + \hat{H}_{\text{c}}$, where $\hat{\tilde{\mathcal{H}}} + \hat{\tilde{\mathcal{H}}} + \hat{H}_{\text{c}}$ is the CNT bulk electron operator as given in Ref. 19, and $\hat{\tilde{\mathcal{H}}} + \hat{\tilde{\mathcal{H}}} + \hat{H}_{\text{c}}$ is the lead electron operator with $\phi_{l}(\mathcal{r})$ denoting the corresponding single-particle wave function.

**IV. TRANSPORT THEORY**

The current, as any other observable of the transport setup, can be calculated by a trace over the associated operator multiplied by the reduced density matrix $\hat{\rho}(t) = \hat{\tilde{\mathcal{H}}} + \hat{\tilde{\mathcal{H}}} + \hat{H}_{\text{c}}$. Being obtained from the density matrix $\rho_{\text{tot}}$ of the total setup upon tracing out the lead degrees of freedom, $\hat{\rho}(t)$ stores the full information about the state of the dot in the presence of the tunnel coupling to the leads. The time evolution of $\hat{\rho}(t)$ follows the Liouville equation, and it can for the stationary state ($\hat{\rho}(t) = 0$) be cast into the form (see, e.g., Ref. 21)

$$0 = -i \sum_{aa'} \delta_{ab} \delta_{a'\bar{b}} (E_a - E_{\bar{b}}) \rho_{a\bar{b}} + \sum_{aa'} K_{ab}^{a\bar{b}} \rho_{a\bar{b}},$$

taking matrix elements with respect to the many-body eigenstates of the CNT: $\rho_{aa'} := \langle a|\hat{\rho}|a'\rangle$ and $E_a := \langle a|\hat{H}|a\rangle$. Furthermore $K_{ab}^{a\bar{b}} := \langle b|\hat{H}|a'\rangle ||a\rangle$, with the kernel superoperator $K$ arising from the perturbation, i.e., the tunnel coupling between quantum dot and leads. Depending on up to which order $2n$ in the tunnel coupling $K$ is calculated, one takes into account effects from $2n$ correlated tunnel events. The most involved part of a perturbative transport calculation is the determination of the kernel $K$.

A fully diagrammatic representation of $K$ has first been proposed in Ref. 15. For example, one contribution to the second-order kernel $(K^{(2)})_{bb_{1}}^{a_{1}a_{2}}$ is shown in Fig. 2(a). It relates to the tunnel process $|a_1\rangle \rightarrow |b_1\rangle$, represented by the solid line connecting upper and lower parts of the diagram contour. In the following we identify specific terms in all orders of the perturbation series as charge fluctuation processes, which sum up to a Taylor series yielding the intrinsic level renormalization $E_{b} \rightarrow E_{b} + h_{\text{intrinsic}}^{bb_{1}}$ to the energy difference $E_{ba} = E_b - E_a$, where $h_{\text{intrinsic}}^{bb_{1}}$ is given in Eq. (7) below.

**A. Diagrammatic evaluation of the intrinsic level renormalization**

Recently, it has been recognized that certain fourth-order diagrams can be related to charge fluctuations in the initial or final state during a tunnel event.

Here we identify a specific class of these charge fluctuation diagrams which, summed up in all orders, yield a whole Taylor series and therewith an actual level shift. Figure 2(b) shows a diagram of order $2n + 2$, contributing to $(K^{(2n+2)})_{bb_{1}}^{a_{1}a_{2}}$, which dresses Fig. 2(a) by $k$ charge fluctuations in the final state (k “bubbles” on the upper part of the contour) and $n - k$ charge fluctuations in the initial state $a$ (n − k “bubbles” on the lower part of the contour). We thereby look at fluctuations isolated, in the sense of separated in time, from each other: each bubble must start and end at consecutive times $\tau_i$ and $\tau_{i+1}$ (where $1 \leq i \leq 2n - 1$). The electron transfer event is initialized at the earliest time $\tau = \tau_0$ and ends at the latest time $\tau_{2n+1} = \tau$. This gives $(\gamma)$ possibilities for the time ordering of the bubbles among the upper and lower parts of the contour. Summing all those plus their hermitian conjugates, the total contribution is, as obtained by applying diagrammatic rules,15
Here $c_j$ and $c'_j$ serve as placeholders for the possible intermediate states; $f^p(\omega) := f^p(\beta \omega + \beta eV_l)$, $p = \pm$, which we defined $f^+(\omega) = f(\omega) = 1/(1 + e^{\omega})$, denoting the Fermi function, while $f^-(\omega) = f(-\omega)$. The lead indices $l, l_i, l_j \in \{s, d\}$ are summed over, and the lead energies $\omega_i$, $\omega_l$, and $\omega_j$ are integrated over. The values of $p$ and $\sigma$ are set by fixing the initial and final states $a$ and $b$, respectively. Likewise the values of the spin indices $\sigma_j$ and $\sigma_i$ and of the Fermi function labels $p_j$ and $p_i$ depend on the intermediate states $c_j$ and $c'_j$ that can be reached from $b$ and $a$, respectively, and are summed over. Finally, there are tunnel matrix elements characterizing transitions on the dot, $T^{(+)}_{l \sigma}(b,a) \sim T_l(\tilde{r}_i)(b|\psi_i^{(l)}(\tilde{r}_i)\rangle a$ ($\tilde{r}_i$ characterizes the position of lead $i$). A multitude of terms emerges from the expression Eq. (6) when applying the decomposition
\[
\lim_{\eta \to 0} \int d\tilde{\omega} \frac{D_{l\sigma}(\tilde{\omega}) f^p(\tilde{\omega})}{\tilde{\omega} - \tilde{\mu} + i\eta} = \int' d\tilde{\omega} \frac{D_{l\sigma}(\tilde{\omega}) f^p(\tilde{\omega})}{\tilde{\omega} - \tilde{\mu}} - i\pi D_{l\sigma}(\tilde{\mu}) f^p(\tilde{\mu})
\]
for all fractions and expanding the product ($\int'$ denotes a principal part integration; $\tilde{\omega} \in [\omega_i, \omega_l, \omega_j]$), etc.). We want to focus on a certain contribution, namely the combination where the fraction containing merely $\omega$,
\[
\left(\frac{1}{p\omega - E_{ba} + i\eta}\right)^{n+1} \equiv \frac{1}{n!} \frac{d}{d^nE_{ba}} \left(\frac{1}{p\omega - E_{ba} + i\eta}\right),
\]
has been replaced by the $\delta$ function and all others by their principal parts.

In terms of $f^{(\alpha)}(\omega)$ it reads
\[
\frac{2\pi}{\hbar} n! f^{(\alpha)}(\beta E_{ba} + \beta p eV_l)D_{l\sigma} |T^{(\alpha)}_{l \sigma}(b,a)|^2 \beta^n \frac{n!}{k}
\]
\[
\times \text{Re} \left[ \frac{1}{p\omega - E_{cb} + p\beta eV_l} \int d\omega \int' d\omega' |T^{(+)}_{l \sigma}(b,a)|^2 \right]
\]
\[
\times \frac{n-k}{k} \int d\omega' \frac{f^{(\alpha)}(\omega')D_{l\sigma}|T^{(\alpha)}_{l \sigma}(c',a)|^2}{\omega' - \beta E_{ac} + p\beta eV_l}
\]
\[
\times \frac{n-k}{k} \int d\omega' \frac{f^{(\alpha)}(\omega')D_{l\sigma}|T^{(\alpha)}_{l \sigma}(c',a)|^2}{\omega' - \beta E_{ac} + p\beta eV_l}.
\]

Exhausting all possibilities of choosing the states $c_j$ and $c'_j$ as well as setting $k = 1, \ldots, n$, it is clear that one generates all kinds of terms appearing in an expansion of the power $n$ of a quantity $h_{ba}^{\ln}$ defined as
\[
h_{ba}^{\ln} = \sum \sum |T^{(\alpha)}_{l \sigma}(c,b)|^2 \int d\omega \frac{f^{(\alpha)}(\omega)D_{l\sigma}(\omega)}{\omega - \beta E_{ba} + p\beta eV_l} + \sum \sum |T^{(\alpha)}_{l \sigma}(c',a)|^2 \int d\omega \frac{f^{(\alpha)}(\omega)D_{l\sigma}(\omega)}{\omega - \beta E_{ca} + p\beta eV_l}.
\]
\[
\delta E_b = \sum \sum |T^{(\alpha)}_{l \sigma}(c,b)|^2 \int d\omega \frac{f^{(\alpha)}(\omega)D_{l\sigma}(\omega)}{p\omega - E_{cb}},
\]
where the values of $p$ and $\sigma$ are settled once the state $c$ is assigned. For the case of the single impurity Anderson model this expression reduces to the result for the level shift obtained in perturbation theory. Hence Eq. (8) provides a diagrammatic interpretation of these results. As temperature goes to zero, these diagrams diverge logarithmically at the charge-degeneracy points, which gives a parametric selection of these specific diagrams.

For a multilevel quantum dot, the summation over the intermediate virtual states $c$ implies that the amplitude of the level shifts is expected to vary from a resonance peak to the other, as also shown in Figs. 4, 5, and 6. Notice also that for the CNT used in our calculations, the summation over the virtual states also includes states with bosonic excitations as described by the bosonic Hamiltonian $\hat{H}_B$ in Eq. (3).

### B. Flat band with Stoner shift

In order to better analyze the contribution of majority and minority spins, we consider for the evaluation of Eq. (7) the case of a flat band including a Stoner splitting $\Delta_S$, i.e., $D_{l\alpha}(\omega)$ as given in Eq. (4). Notice that in order to retain holomorphic functions, one can make use of the Fermi function representation of the step functions, $\theta(\pm \epsilon) = \lim_{\beta \to \infty} f^{\pm}(\beta \epsilon)$. Using
then the residue calculus, one obtains
\[ h_{\text{int}}^{ba} = \sum_i \left[ \sum_c D_{1c} \left| T_{1c}^{\sigma}(c,b) \right|^2 \tilde{\Psi}_{\sigma,c}^{(0)}(\beta E_{cb} + p\beta eV_I) \right. \]
\[ - \sum_c D_{1c} \left| T_{1a}^{\sigma}(c,a) \right|^2 \tilde{\Psi}_{\sigma,c}^{(0)}(\beta E_{ac} - p\beta eV_I) \left. \right], \tag{9} \]

where the upper/lower boundary \( \pm W - \sigma \Delta_{Sl}/2 \) of the integration enters via
\[ \tilde{\Psi}_{\sigma,c}^{(0)}(x) = \text{Re} \left[ \Psi^{(0)} \left( 0.5 + i \frac{x}{2\pi} \right) \right] \]
\[ -\Psi^{(0)} \left( 0.5 + i \frac{x - \beta(\pm W - \sigma \Delta_{Sl}/2)}{2\pi} \right) \],

where \( \Psi^{(0)} \) is the digamma function. An approximation of this result is obtained elegantly when using Eq. (4) to split the integration range of the integrals in Eq. (7) as
\[ \int_{W - \Delta_{sl}/2}^{-W + \Delta_{sl}/2} d\omega = \int_{W - \Delta_{sl}/2}^{-W + \Delta_{sl}/2} d\omega + \int_{W + \Delta_{sl}/2}^{-W - \Delta_{sl}/2} d\omega, \]
\[ \int_{W - \Delta_{sl}/2}^{W + \Delta_{sl}/2} d\omega = \int_{W - \Delta_{sl}/2}^{-W + \Delta_{sl}/2} d\omega + \int_{W + \Delta_{sl}/2}^{-W - \Delta_{sl}/2} d\omega, \]
and noting that in the region at the top/bottom of the band is \( f(\omega) = 0/1 \). Here we get the result
\[ h_{\text{int}}^{ba} = \sum_i \left[ \sum_c D_{1c} \left| T_{1c}^{\sigma}(c,b) \right|^2 \tilde{\Psi}_{\sigma,c}^{(0)}(\beta E_{cb} + p\beta eV_I) \right. \]
\[ - \sum_c D_{1c} \left| T_{1a}^{\sigma}(c,a) \right|^2 \tilde{\Psi}_{\sigma,c}^{(0)}(\beta E_{ac} - p\beta eV_I) \left. \right] + \sum_{i,c} D_{1c} \left| T_{1c}^{\sigma}(c,b) \right|^2 \ln \left( \frac{W_I + E_{cb} + peV_I}{W_I + E_{cb} + peV_I} \right) \]
\[ + \sum_{i,c} D_{1c} \left| T_{1c}^{\sigma}(c,a) \right|^2 \ln \left( \frac{W_I - E_{ac} + peV_I}{W_I - E_{ac} + peV_I} \right), \tag{10} \]

where \( W_{\sigma} = W - \sigma \Delta_{Sl}/2 \) is the finite bandwidth for the integration over the lead energies. The result is an approximation of Eq. (9), as can be seen when applying \( \text{Re} \Psi^{(0)}(0.5 + iA) = \text{Re} \Psi^{(0)}(0.5 + iB) \) in Eq. (10).

The first two terms depend strongly on gate voltage, interaction, and polarization. They correspond to the contribution to the level shift first proposed in Ref. 11.

In contrast, obviously, the last two terms in Eq. (10) vanish if \( \Delta_{Sl} = 0 \). Moreover, for \( E_{cb}, E_{ac} \ll W_{\sigma} \) they are very weakly dependent on the gate voltage and polarization, but do depend on the saturation magnetization via the Stoner splitting. Following Ref. 5, with tunneling coupling \( \Gamma \approx 1 \text{ meV} \), this contribution can yield for PdNi contacts the gate-independent part of the level shift of the order of a few hundreds of \( \mu \text{eV} \). This is of the same order as the constant shift used to fit the Kondo data in Refs. 3 and 5 but also the TMR data in Ref. 6. A full account of the interface phase shifts might be necessary for large transparencies and goes beyond the scope of our tunneling Hamiltonian.

However, in the regime where the tunneling model is valid (Coulomb blockade and Kondo regime), tunneling-induced shifts should suffice to explain the experiments.

All in all, we can recast the expression for \( h_{\text{int}}^{ba} \) in the form
\[ h_{\text{int}}^{ba} = h_0^{ba} + \Delta h_{\text{int}}^{ba} (\Delta_{St}). \tag{11} \]

The first term depends logarithmically on the gate voltage and the interaction strength, while \( \Delta h_{\text{int}}^{ba} \) is the contribution due to \( \Delta_{St} \), which depends on the finite bandwidth \( W_{\sigma} \) of \( D_{1c}(\omega) \) and will be almost gate and interaction independent and thus contribute in the same way as the terms \( h_{\text{ext}}^{P/A} \) induced by \( \hat{H}_{\text{ext}}^{P/A} \).

Notice that if the states \( a, b, c \) and \( c' \) have electron-hole symmetry partners \( a_\ast, b_\ast, c_\ast \) and \( c'_\ast \), there exists the mirror transition \( b_\ast \rightarrow a_\ast \), with \( a_\ast \) now being the final state and \( b_\ast \) the initial state. Thus the contributions from the gate-dependent part of the charge fluctuations to \( c_\ast \) are negative, and those to \( c'_\ast \) are positive, such that \( h_{\text{int}}^{ba_\ast} = h_{\text{int}}^{ba} \). This preserves the mirror symmetry of all curves when \( \Delta_{St} = 0 \) [cf. Fig. 3(a)].

V. LINEAR CONDUCTANCE AND TMR RESULTS

We consider the linear bias regime for spin polarized transport across a CNT without band offset (cf. Table I).

Figure 3 shows the conductances for parallel (pink solid) and antiparallel (blue dashed) lead magnetizations under the influence of the distinct types of level shifts, along with the corresponding TMR curves. Throughout this work we assume equal lead polarizations \( P_L = P_R = P \) and the coupling asymmetry \( \gamma = \Gamma_L/\Gamma_R \sim |T_I|^2 / |T_0|^2 \), with the actual values of \( \gamma \) given at each plot.

In the absence of any Zeeman shift, Fig. 3(a) shows the typical conductance patterns expected for the degeneracies according to Table I, for both a second-order (sequential tunneling) and a fourth-order (cotunneling, pair tunneling, single charge fluctuations, etc.) truncation in the calculation of the kernel \( K \). Second-order theory predicts a constant TMR value, but this changes upon inclusion of higher-order effects. Within fourth-order perturbation theory, the TMR exhibits an oscillatory gate voltage dependence, albeit the variation is small. Notice that the increase at the edges of the plot is nonphysical, stemming from a limitation of the numerical calculation to six \( 4n - 1 \leq N \leq 4n + 4 \) charge states.

To visualize the TMR mechanism from Figs. 1(c) and 1(d), we include in Fig. 3(b) merely a constant shift from an external source, \( |h_{\text{ext}}^{P/A}| = |h_{\text{ext}}^{A}| > k_B T \) (this can arise from the external field which controls the direction of contact polarization in TMR measurements, when for the measurement of \( G_P \) the field is swept back to the point where \( G_A \) has been measured before).

As explained above, among the formerly degenerate lowest-lying states, the ones with maximum spin projection \( S \), are selected as ground states, as highlighted in Table I. At the first two peaks transport is thus mediated by \( \uparrow \) electrons, and at the last two peaks by \( \downarrow \) electrons. This fact breaks the symmetry in the conductance curve for a parallel configuration which favors transport via the \( \uparrow \) electron channel. For an antiparallel magnetization of the leads, both spin species are minority charge carriers in one of the two leads and thus provide
FIG. 3. (Color online) Parallel ($G_P$) and antiparallel ($G_A$) conductance along with the resulting tunneling magneto-resistance (TMR) for a CNT of 500 nm in length ($\varepsilon_0 = 3.35$ meV) and charging energy $E_c = 6.7$ meV. The thermal energy was set to $k_B T = 0.3$ meV, the lead polarization to $P = 0.4$, the tunnel broadening to $\hbar \gamma_1 = 3$ $\mu$eV, and the bandwidth to $W = 3$ eV. (a) For a full fourth-order calculation a mirror-symmetric TMR slightly oscillating around a value of 20% is obtained. (b) An equal splitting $h_{ext}^P = h_{ext}^A = 0.4$ meV causes gate asymmetry and negative values of the TMR by the mechanism shown in Fig. 1(c). (c) For $h_{ext}^P = 2h_{ext}^A = 0.8$ meV, the regularity of the curve is broken and the negative TMR mechanism shown in Fig. 1(b) comes into play. (d) Same as panel (d) but for $\Delta St = 0.2W$, which breaks electron-hole symmetry. Including the influence of the virtual excited states negative TMR can be reached. (f) Combining intrinsic shifts with external splitting. An asymmetric coupling to the leads can enlarge ($\gamma < 1$) or diminish ($\gamma > 1$) the TMR effect.

equally weak transport channels, such that the symmetry in the conductance curve is preserved. In Fig. 3(c) we still neglect intrinsic shifts, but assume $|h_{ext}^P| > |h_{ext}^A|$ ($> k_B T$, e.g., due to stray fields) to bring also the first negative TMR mechanism, Fig. 1(b), into play. It overlays the effects observed in Fig. 3(b). The shift in the peak positions of parallel and antiparallel conductance peaks enforces a change of the TMR from positive to negative value from the first peak on.

Because there is no significant change in the curves of Figs. 3(b) and 3(c) upon inclusion of the fourth-order effects and the numerical implementation of the gate-dependent contribution $h_{ext}^P$ implies a summation over a large number of states for a CNT quantum dot, in the remaining figures we only show second-order curves with all level shifts fully included.

FIG. 4. (Color online) Gate-voltage dependence of the intrinsic level shift $h_{ext}^P$, Eq. (7), for exemplary electron-hole symmetric transitions, see state labels in panel (a). The electron-hole symmetry is nicely reflected in the curves for $\Delta st = 0$. Moreover, it becomes obvious that $\Delta St$ acts opposite to an external Zeeman splitting, i.e., increases (decreases) the energy difference for $\uparrow$ ($\downarrow$) transitions. Raising the energy cutoff for the inclusion of excited states, more and more transitions to virtual states will be contributing to the charge fluctuations for the exemplary transitions. (a) Only fluctuations to ground states. (b) Fluctuations to all states within the range $\varepsilon_0$. (c) All states within the range $2\varepsilon_0$. 

045313-6
can be identified. Note that going beyond a cutoff of 2\(\Delta_\text{SI} = 0\) for parallel configuration for the two states \(b = |↑, \downarrow\rangle\) and \(b' = |↓, \downarrow\rangle\) \((N = 4n + 1, \text{upper panel})\) and \(b = |↑, \downarrow\rangle\) and \(b' = |↑, \downarrow\rangle\) \((N = 4n + 3, \text{lower panel})\). Notice the pronounced influence of virtual states including bosonic excitations. For the legend, see Fig. 5.

are confirmed by Figs. 4(a)–4(c) which show the shifts \(\delta h_{\text{eff}}^\text{int}\) (Eq. (7), the quantity determining the shift of position of the conductance peaks) for selected electron-hole symmetric transitions \(|b\rangle \rightarrow |a\rangle\). Due to the orbital degeneracy of the CNT, the shifts are seen to differ from peak to peak. The trend to convergence is obvious, in particular, around the resonances the respective transitions belong to (curves thickened in the respective gate voltage region).

Finally, Fig. 3(f) combines the impact of the intrinsic shifts with an equal extrinsic splitting \(|h_{\text{ext}}^\text{int}| = |h_{\text{ext}}^\text{int}|\). The intrinsic effects suffice to change the TMR curve, Fig. 3(b), to a shape observed in experiments,\(^3\) though due to our limitation to the weak coupling regime, quantitative agreement cannot be expected. An asymmetric coupling is found to affect the curve quantitatively, while the qualitative shape is retained. Thereby, \(\gamma < 1\) \((\gamma > 1)\) relates to an increased (decreased) coupling to the drain contact—the one in which the parallel and antiparallel configurations differ from each other [see Fig. 1(a)]—and enhances (suppresses) the TMR effect. Even for the very small values of \(\Gamma_\text{D}\) used here to justify lowest-order perturbation theory, the difference between Figs. 3(b) and 3(f) reveals a marked influence of the intrinsic, tunneling induced, level shifts.

The dramatic influence of the excited states, and among these, in particular, the bosonic excitations, is further demonstrated in Figs. 5 and 6.

In Fig. 5 the difference of energy shifts \(\delta E^b_\text{SI} - \delta E^A_\text{SI}\) [see Eq. (8)] is plotted versus gate voltage for the same states as those used in Fig. 4. Notice that this quantity vanishes for states with total spin 0. For nonzero spin, the shifts exhibit a characteristic gate dependence, which proves to be most sensitive to the presence of bosonic excitations: omitting them (Fig. 5, magenta dotted + cyan dash-dotted) influences the curve not only quantitatively but also qualitatively. Compared to that effect, it plays only a minor role whether the cutoff is chosen to be \(\epsilon_0\) (blue solid) or \(2\epsilon_0\) (red dashed), and the curves for \(\Delta_\text{SI} = 0\) and \(\Delta_\text{SI} \neq 0\) practically fall upon one another. In perfect similarity to an externally induced splitting, there is a

FIG. 5. (Color online) Gate-voltage dependence of the difference of energy shifts \(\delta E^b_\text{SI} - \delta E^A_\text{SI}\) [see Eq. (8)] for states with spin 0 (a), spin \(\pm 1/2\) (b), and spin \(\pm 1\) (c). Notice the pronounced influence of virtual states including bosonic excitations.

FIG. 6. (Color online) Gate-voltage dependence of the difference of energy shifts \(\delta E^b_\text{SI} - \delta E^A_\text{SI}\) for parallel configuration for the two states \(b = |↑, \downarrow\rangle\) and \(b' = |↓, \downarrow\rangle\) \((N = 4n + 1, \text{upper panel})\) and \(b = |↑, \downarrow\rangle\) and \(b' = |↑, \downarrow\rangle\) \((N = 4n + 3, \text{lower panel})\). Notice the pronounced influence of virtual states including bosonic excitations. For the legend, see Fig. 5.
factor of 2 between the differences of the energy shifts for the spin-\(\frac{1}{2}\) and the spin-1 states.

In Fig. 6 the quantity \(\delta E_b - \delta E_p\) is shown for the case of parallel configuration, for the two states \(b = |\uparrow, \cdot\rangle\) and \(b' = |\downarrow, \cdot\rangle\) (upper plot) and \(b = |\uparrow, \downarrow\rangle\) and \(b' = |\downarrow, \uparrow\rangle\) (lower plot). These shifts for states with 1 and 3 excess electrons are those which have been measured, e.g., in the Kondo experiments of Refs. 3 and 5. For \(\Delta S_b = 0\) the result is already converged, as the curves for cutoff energies \(\varepsilon_0\) and \(2\varepsilon_0\) are mostly identical. For \(\Delta S_b = 0.2W\) the energy difference acquires a negative offset.

VI. CONCLUSIONS

We analyzed the impact of different kinds of effective Zeeman shifts in magnetically coupled multilevel quantum dots, obtaining a characteristic gate dependence and the possibility of negative TMR. In particular, we have provided a systematic way of including the important effects of tunneling-induced level shifts in a transport calculation by identifying a subclass of diagrams, to all orders in the tunneling Hamiltonian, describing charge fluctuations. This infinite class of diagrams is seen to correspond to a perturbative renormalization of the dot spectrum, an effect which is indeed expected on physical grounds. In conventional Feynman diagrammatics, these terms arise from summing up the second-order dot electron self-energy in the Dyson equation. Importantly, this concept generalizes to higher orders: virtual charge fluctuations during inelastic cotunneling (appearing first in sixth order) give a renormalization of the inelastic cotunneling threshold, as already experimentally observed in Ref. 26. In general, a TMR signal will be influenced by many parameters relevant to the given device. Nevertheless, following the lines of the analysis given here for a CNT, it should be possible to disentangle the importance of the various contributions.

ACKNOWLEDGMENTS

We acknowledge fruitful discussions with C. Strunk, S. Pfaller, and J. Hauptmann as well as the support of the DFG under Program SFB 689.

21The difference \(\Delta S_b\) is zero unless the electronic configurations of \(b\) and \(c\) differ by one electron of spin \(\sigma\). Then, \(p = \text{sgn}(N_r - N_b)\), with \(N_b := |a|\langle b|\).
22Equivalently to \(\int d\omega f^\ast(\omega)D_{\mu}(\omega)/(\omega - \mu)\) one can consider \(\text{Im} \int_\mu^\infty d\omega D_{\mu}(\omega) f^{\ast}(\beta \omega) e^{i\omega\tau_\sigma}/\omega\).
23Notice that \(\Gamma_i \sim D_i \tau_i^2\) and \(\Gamma = \sum_i \Gamma_i\).