1. NONLINEAR CONDUCTANCE TRACES FOR A TWO-IMPURITY KONDO SYSTEM: MODELING ASYMMETRIC DOUBLE-PEAK ZBAs

This section summarizes the approach that we used for calculating the nonlinear differential-conductance traces of Fig. 4 in the main text. The experimental results show nonlinear conductance traces with highly asymmetric double peak ZBAs: the peak height and width of the two peaks can differ significantly, and the minimum between the two peaks can appear at positions that differ significantly from $V_{sd} = 0$. The purpose of the theoretical modeling that is presented in this Section is to investigate whether such asymmetric double-peak ZBAs are consistent with the physics of the two-impurity Kondo system. The theoretical traces of Fig. 4 are calculated for such a two-impurity Kondo system, and show behavior that is for a large part consistent with the behavior of the experimentally observed asymmetric double-peak ZBAs. In addition, several devices showed rather symmetric single-peak ZBAs that had a position that differed significantly from $V_{sd} = 0$. This behavior also comes forward from the two-impurity Kondo modeling for parameters where the two impurity spins have unequal coupling to their reservoir and for a spin-spin coupling that is weaker than temperature (traces not shown here).

The main challenge for this theoretical modeling is to extend the two-impurity Anderson Hamiltonian to calculations of the differential conductance for transport through two series-coupled impurities at nonzero bias voltage and nonzero temperatures. The approach that we use here directly builds on refs. 1,2. The model parameters that were used for calculating the traces in Fig. 4 of the main text are summarized at the end of this Section.

Theoretical model

The system is modeled as a two-impurity Anderson Hamiltonian with an extra term accounting for inter-impurity coupling. Each impurity is connected to a different Fermi sea with chemical potential $\mu_L = \frac{eV_{sd}}{2}$ and $\mu_R = -\frac{eV_{sd}}{2}$, respectively. The full Hamiltonian reads

$$H = \sum_{k_\alpha \in \{L,R\}, \sigma} \epsilon_{k_\alpha} c_{k_\alpha,\sigma}^\dagger c_{k_\alpha,\sigma} + \sum_{\alpha \in \{L,R\}, \sigma} \epsilon_{\alpha\sigma} d_{\alpha\sigma}^\dagger d_{\alpha\sigma} + V_0 \sum_{k_\alpha \in \{L,R\}, \sigma} (c_{k_\alpha,\sigma}^\dagger d_{\alpha\sigma}^\dagger c_{k_\alpha,\sigma} + d_{\alpha\sigma}^\dagger c_{k_\alpha,\sigma})$$

$$+ V_I \sum_{\sigma} (d_{L\sigma}^\dagger d_{R\sigma} + d_{R\sigma}^\dagger d_{L\sigma}) + U_L n_{L,\uparrow} n_{L,\downarrow} + U_R n_{R,\uparrow} n_{R,\downarrow}. \tag{1}$$

The first two terms in Eq. (1) represent the electrons in the leads and in the impurities, re-
respectively. In these terms, \( c_{kL/R,\sigma}^\dagger (c_{kL/R,\sigma}) \) creates (annihilates) an electron with momentum \( k_{L/R} \) and spin \( \sigma \) in the left/right lead, and \( d_{L/R,\sigma}^\dagger (d_{L/R,\sigma}) \) creates (annihilates) an electron with spin \( \sigma \) in the left/right impurity. \( \epsilon_{kL/R} = \epsilon_k + \mu_{L/R} = \epsilon_k \pm \frac{\epsilon V_{sd}}{2} \) are the energies in the leads, while \( \epsilon_{\alpha\sigma} \) are the bare energies at each impurity. The third term describes the coupling between each impurity and its corresponding lead, and determines the coupling strength \( \Delta_{L,R}(\epsilon) = \pi V_0^2 \sum_{k_{\alpha \in \{L,R\}}} \delta(\epsilon - \epsilon_{k_{\alpha}}) \) (we neglect the \( k \) dependency of the tunneling matrix element for simplicity). Each lead is described by a parabolic density of states (energy bandwidth \( W = 2D \)) centered at the chemical potential, such that we can define the function

\[
\Delta_{\alpha}(\epsilon) = \pi V_0^2 \sum_{k_{\alpha}} \delta(\epsilon - \epsilon_{k_{\alpha}}) = \begin{cases} 
\Delta_0 \left[ 1 - \left( \frac{\epsilon - \mu_{\alpha}}{D} \right)^2 \right] & \text{if } -D \leq \epsilon - \mu_{\alpha} \leq D, \\
0 & \text{otherwise.}
\end{cases}
\]

The fourth term describes inter-impurity coupling. In the absence of such coupling, this Hamiltonian describes two independent Anderson impurities each of them coupled to different Fermi seas. In the limit of \( U_L, U_R \to \infty \) we can write the Hamiltonian (1) in terms of auxiliary pseudo-fermions and slave boson (SB) operators plus constraints:

\[
H = \sum_{k_{\alpha \in \{L,R\}},\sigma} \epsilon_{k_{\alpha}} c_{k_{\alpha},\sigma}^\dagger c_{k_{\alpha},\sigma} + \sum_{\alpha \in \{L,R\},\sigma} \epsilon_{\alpha\sigma} f_{\alpha\sigma}^\dagger f_{\alpha\sigma} + \frac{t_I}{N} \sum_{\sigma} (f_{L\alpha}^\dagger b_{L\alpha}^\dagger f_{R\sigma} + f_{R\alpha}^\dagger b_{R\alpha}^\dagger f_{L\sigma}) + 
\frac{V_{sd}}{\sqrt{N}} \sum_{k_{\alpha \in \{L,R\}},\sigma} (c_{k_{\alpha},\sigma}^\dagger b_{\alpha\sigma}^\dagger f_{\alpha\sigma} + f_{\alpha\sigma}^\dagger b_{\alpha}^\dagger c_{k_{\alpha},\sigma}). \tag{2}
\]

In the slave boson representation, the annihilation operator for electrons at the impurity sites, \( d_{\alpha\sigma} \) is decomposed into the SB operator \( b_{\alpha}^\dagger \) which creates an empty state and a pseudo fermion operator \( f_{\alpha\sigma} \) which annihilates the singly occupied state with spin \( \sigma \) in the impurity \( \alpha \): \( d_{\alpha\sigma} \to b_{\alpha}^\dagger f_{\alpha\sigma} \) \( (d_{\alpha\sigma}^\dagger \to f_{\alpha\sigma}^\dagger b_{\alpha}) \). Note that we have scaled the hopping parameters \( V_0 = \frac{V_{sd}}{\sqrt{N}} \) and \( V_I = \frac{t_I}{N}, \) \( N \) being the degeneracy of the level on each impurity. This scaling is done in such a way that the parameters \( \Gamma = N \Delta_0 \) and \( \Delta_0/V_I = N \Delta_0/t_I = \Gamma/t_I \) appearing in the expression of the Kondo temperature have a well defined \( N \to \infty \) limit, namely there is a well defined \( 1/N \) expansion of the physical quantities. At the end of the calculation, the physical limit \( N = 2 \) is, of course, taken. Finally, the physical constraint is that we must work in a subspace of the Hilbert space where the number of auxiliary particles (on each
impurity) is one, namely:

\[
\hat{Q}_L = \sum_\sigma f_{L\sigma}^\dagger f_{L\sigma} + b_L^\dagger b_L = 1,
\]

\[
\hat{Q}_R = \sum_\sigma f_{R\sigma}^\dagger f_{R\sigma} + b_R^\dagger b_R = 1.
\]

These two constraints come from the physical condition that each impurity has to be in one of the three states \(|0\rangle, |\uparrow\rangle\) or \(|\downarrow\rangle\).

At this point, we have reduced the original problem described by the Hamiltonian in Eq. (1) to a problem of fermions and bosons interacting through tunneling terms and subject to the constraints in Eq. (3). Properties of the physical electrons can be built up from the Green’s functions of the pseudo-fermions and slave bosons. These Green’s functions for the auxiliary fermions and bosons constitute the basic building blocks of the theory. Furthermore, our aim is to study the out-of-equilibrium properties of the system; we need, then, a fully non-equilibrium description of the dynamics of the Green’s functions of these auxiliary particles. The appropriate starting point is to derive equations-of-motion for the time-ordered double-time Green’s function of the auxiliary fermion and boson fields on a complex contour. In order to do this we employ the so-called non-crossing approximation (NCA)\(^3\)\(^-\)\(^6\) generalized to a two-impurity Anderson Hamiltonian\(^2\). Without entering into much detail of the theory, we just mention that the boson fields are treated as fluctuating operators such that both thermal and charge fluctuations are included in a self-consistent manner to order \(O(1/N)\). In particular, one has to derive self-consistent equations-of-motion for the time-ordered double-time Green’s function (sub-indexes are omitted here):

\[
iG(t,t') \equiv \langle T_{c} f(t) f^\dagger(t') \rangle,
\]

\[
iB(t,t') \equiv \langle T_{c} b(t) b^\dagger(t') \rangle,
\]

or in terms of their analytic pieces:

\[
iG(t,t') = G^>(t,t')\theta(t - t') - G^<(t,t')\theta(t' - t),
\]

\[
iB(t,t') = B^>(t,t')\theta(t - t') + B^<(t,t')\theta(t' - t).
\]

A rigorous and well established way to derive these equations-of-motion was first introduced by Kadanoff and Baym\(^7\), and has been related to other non-equilibrium methods (like
the Keldysh method) by Langreth, see ref. 8 for a review. In the paper, we just show numerical results of the coupled set of integral NCA equations for our problem and refer the interested reader to refs. 2–6 for details. In particular, the density of states is given by

$$\rho(\omega) = \frac{-1}{\pi} \sum_\sigma \text{Im}[A^\sigma_\omega(\omega)],$$

where $A^\sigma_\omega(\epsilon)$ is the Fourier transform of the retarded Green’s function $A^\sigma_\omega(t) = G^\sigma_\omega(t)B^\omega(-t) - G^\omega_\omega(t)B^\sigma(-t)$. Note that this decoupling neglects vertex corrections and, as a result, the NCA fails in describing the low-energy Fermi-liquid regime. Nevertheless, the NCA has proven to give reliable results even at temperatures well below the Kondo temperature (of the order of $T = 10^{-2} T_K$). Following Meir and Wingreen in ref. 10, the current is given by

$$I_{\alpha \in \{L, R\}} = -\frac{2e}{h} \sum_\sigma \int d\epsilon \Gamma_\alpha(\epsilon) \left[ 2\text{Im} A^\sigma_\omega(\epsilon)f_\alpha(\epsilon) + A^\omega_\omega(\epsilon) \right],$$

with $A^\omega_\omega(\epsilon)$ the Fourier transform of $A^\omega_\omega(t) = iG^\omega_\omega(t)[B^\epsilon(-t) - B^\omega(-t)]$ and $f_\alpha(\epsilon) = \frac{1}{1 + e^{(\epsilon - \mu_\alpha)/kT}}$ the Fermi-Dirac function at each reservoir held at a chemical potential $\mu_\alpha$ such that the applied bias voltage is defined as $eV_{sd} = \mu_R - \mu_L$.

In practice, we self-consistently solve the NCA integral equations for each isolated Anderson impurity until good numerical convergence is reached. In a second self-consistent step, we obtain the self-energies coming from inter-impurity coupling.

**Parameters used for the theoretical traces in Fig. 4**

The theoretical traces in Fig. 4 of the main text are calculated as follows and for the following parameters. The inset of Fig. 4a presents traces of $dI/dV_{sd}$ calculated with the theory above here. The numerics are performed by discretizing the Fourier space in a finite mesh of size $N_\omega = 2^{18}$ with cutoff $D = 20$. The asymmetry primarily arises from taking unequal coupling strengths $\Gamma_L$ and $\Gamma_R$ between the impurities and their respective reservoirs. Further reference to a value for $\Gamma$ assumes the relation $\Gamma = \frac{\Gamma_L + \Gamma_R}{2}$. The traces in the inset of Fig. 4a are for $\Gamma_L/\Gamma_R = 1.5$, and inter-impurity coupling $t_i = 1.2 \Gamma$. For the presentation in the main text all energy scales are with respect to the highest Kondo temperature $T_K$ of the two-impurities.

We define it by using the following estimate for $T_K$ of each impurity,

$$T_K = D \sqrt{\frac{1}{\pi |\epsilon_i|}} \exp \left( \frac{-\pi |\epsilon_i|}{\Gamma_i} \right),$$
where $\varepsilon_i$ is the energy of the Anderson impurity level $i = L, R$, and $\Gamma_i$ should be filled in as $\Gamma_L$ or $\Gamma_R$ for the interaction of impurity $L, R$ with its respective reservoir. (Note that for the single-impurity case with coupling to two reservoirs on two sides, one must use $\Gamma_i = \Gamma_L + \Gamma_R$, which yields a Kondo temperature that is significantly higher than the values for the two-impurity case.) The bias voltage $V_{sd}$ is thus expressed in units of $k_B T_K/e$ ($k_B$ is Boltzmann’s constant). The traces are for increasing temperature from $T = 0.76 T_K$ (black trace) to $T = 1.33 T_K$ (red trace), see also the colored dots in the inset of Fig. 4b. The inset Fig. 4b is derived from these traces in the same manner as for the experimental traces, simply by extracting the conductance between the two peaks. We used here for all traces the conductance level at $V_{sd} = -0.88 k_B T_K/e$.

For calculations of magnetic field dependence, as presented in the inset of Fig. 4c, we simply added a Zeeman energy term for each of the localized spins, namely $\pm \Delta Z/2$. The traces are for (top to bottom) Zeeman energies of $\Delta Z = 0$ to $\Delta Z = 11 k_B T_K$. It is well known that NCA (since it is a high-$N$ method) shows unphysical zero bias features at finite magnetic fields at low temperatures. Indeed, running the NCA for the single-impurity case reveals such a spurious anomaly. However, as one increases the temperature this feature goes away much faster than the physical features coming from Zeeman-split Kondo peaks. In order to avoid such spurious features, while obtaining good numerical convergence, we had to work with a better mesh resolution ($N_\omega = 2^{20}$) and more stringent convergence parameters (as a criterium for good convergence, the iteration stops when the relative error between successive occupations, as calculated from lesser Green’s functions, in the iteration loop is less than $10^{-6}$). Also, we use slightly different parameters (a bit larger inter-impurity coupling $t_i = 1.3 \Gamma$ and a bit smaller asymmetry $\Gamma_L/\Gamma_R = 1/0.9$) in order to resolve the Zeeman-split peaks without the need of going to lower temperatures that might be on the verge of reliability. This explains the slightly higher conductance values in Fig. 4c as compared to Fig. 4a (and it is known that this method can yield $dI/dV$ values that are slightly too high on the scale of $2e^2/h^4$). Hence, these plots at finite magnetic fields are meant to show qualitative agreement. Quantitative agreement in such $dI/dV$ calculations at finite magnetic field and temperature is beyond this technique (or any other technique that we know of).
2. SDFT SIMULATIONS

Summary of approach and main findings
To support the picture for the many-body effects that was presented in the main text we present numerical SDFT results for QPCs, generalizing earlier calculations\textsuperscript{11,12}, to allow for QPCs of changing length. The length of the channel was determined by a gate of variable length. Opening the QPC is controlled with a single gate-voltage parameter $V_g$. We studied whether the SDFT yields localised states with about one electron of charge as the state with lowest energy. For the true ground state, these states should have spin-singlet character, but this cannot be addressed with SDFT in the present approximation. SDFT can still be used for checking whether a spin-polarised solution (in an arbitrary direction) has a lower energy than an unpolarised solution. Since the polarisation direction is arbitrary, the SDFT then admits two degenerate ground states and the true ground state is a superposition of these two cases with spins in a singlet state. The electron density, though, is the same for the two degenerate SDFT ground states, and thus also for their linear superposition. To allow for a spin-polarised solution, we started the iterative procedure with a finite magnetic field in arbitrary direction that breaks the spin-symmetry, which was turned off in subsequent iterations. For most of the values of gate voltage below the first plateau, the polarised solution had lower energy than the unpolarised solution (Fig. S4). An example of such SDFT results is presented in the Fig. S1 (but similar results are found for a wide range of parameters). Figure S1a displays the electron density for one of the ground states, for both spins, with increasing QPC length, demonstrating an increasing number of ELSs, denoted by circles (the number of ELSs was determined from the total density in the QPC, by fitting the density to a sum of two-dimensional Gaussians, each of total density $e$). Figure S1b depicts the number of ELSs in the gate voltage-length plane, demonstrating the changing number of ELSs with these two parameters. This supports the picture we have presented in the main text: The number of ELSs increases by one each time when the QPC length increases over a range that allows for one additional period of the Friedel oscillation in the QPC channel.
FIG. S1: **Results of Spin-Density-Functional-Theory calculations (SDFT).** a, Spin-polarised electron density in QPC channels (top view on 2DEG plane) for spin-up (right panels) and spin-down (left panels) at constant gate voltage $V_g = -6$ (arb. units, giving a conductance just below $1 \cdot (2e^2/h)$) for three different lengths (540 nm, 680 nm and 830 nm). The number of ELSs (marked with white dotted circles) inside the QPC channel is 2, 3 and 4 respectively. The number of ELSs increases by one each time when the QPC length increases over a range that allows for one additional period of the Friedel oscillation in the QPC channel (the Fermi wavelength in the plane 2DEG area was for these calculations 150 nm). The color scale extends from 0 (black) to $2 \cdot 10^{14}$ m$^{-2}$ (yellow). b, The number of ELSs inside the QPC as a function of gate voltage $V_g$ (parameter for opening the QPC channel) and QPC channel length.
SDFT approach and methods

The SDFT simulations were carried out in a rectangular box (representing a piece of 2DEG), shown in Fig. S2, with along the x-axis (horizontal in Fig. S2) periodic boundary conditions. The external potential is composed of a harmonic part $\frac{1}{2}\omega_y^2 y^2$ that represents the wire and the QPC potential which is calculated by placing two negatively charged gate electrodes at height $z_0 = 100$ nm above the 2DEG in the middle, we are using the Yukawa potential

$$v(x_0, y_0) = \int \rho_g \frac{e^{-|r-r_0|/\gamma}}{|r-r_0|} dr$$

(7)

where $r_0 = \sqrt{x_0^2 + y_0^2 + z_0^2}$, $\gamma = 110$ nm and the integration is over the two-dimensional electrodes. The charge density of the electrodes $\rho_g$, is a function of the gate voltage on the electrodes $V_g$. In our simulation we set $\rho_g$, nevertheless, in order to present the results as in the experimental results, we regard it as gate voltage $V_g$ with arbitrary units.

For the Hartree term we use the appropriate two-dimensional system with one-dimensional periodic boundary conditions\(^{13}\). In addition, we add a positive image charge plane at height 100 nm above the 2DEG as the contribution from the donor layer\(^{11}\).

For the exchange and correlation functionals we use the local-density approximation, for the exchange we use slater exchange\(^{14}\) and the correlation functional is taken from quantum Monte-Carlo simulations of uniform electron gas\(^{15}\). The total number of electrons is $N = 108$ and the temperature is 300 mK. Though we have repeated some of the calculations with temperatures down to 60 mK, with very little change in the results. We used the Octopus code\(^{16}\) for solving the equations. For all the simulations the electron effective mass $m_e^* = 0.067m_e$ and the dielectric constant $\kappa = 12.9$. The actual 2DEG electron density is taken slightly lower than in the experiments to keep the computational time of a simulation at a reasonable level, but we work in a regime where we capture the relevant physics. As a result, the relevant length scales (which are relative to the Fermi wavelength) are for the simulations also slightly longer than the experimental values.

The simulation steps are as follows:

- Set an external potential for a given QPC gate length (parameterized by giving the gate electrodes a length $L$ along the channel) and given gate voltage of the electrodes. In this simulation, opening the QPC is thus controlled with a single gate-voltage parameter $V_g$ (in arbitrary units).
• Find the unpolarized ground state of the system by solving self-consistently the Kohn-Sham equations.

• Then we polarize the solution by applying a magnetic field perpendicular to the sample (only the Zeeman term) for a few iterations, turning it off and letting the system flow to its ground state again (in this procedure we are basically giving an educated initial guess for the density). In principle the field can be applied in any direction as it yields degenerate solutions.

• Repeat these steps for different QPC lengths and gate voltages.

Figure S3 shows two examples of the resulting densities, the left column is spin-down density while the right column in spin-up density. The top row presents the unpolarized solution, the middle row the spatially-symmetric polarized solution, and the bottom row the spatially-anti-symmetric polarized solution. The spatial symmetry of the magnetic field determines the symmetry of the solution.

Figure S4 presents the Free energy difference $\Delta E$ between the polarized solutions relative to the unpolarized solution. If there exists a polarized solution it has a lower energy than the unpolarized solution. Moreover, the spatially-symmetric solution has a region $V_g = -[6.5 - 5.5]$ where it is the ground state of the system. In the following we will concentrate on this region which is below the first plateau.

To determine the number of emergent localized states (ELSs) that are in the effective QPC channel we cannot use that they appear in the region that is determined by the lithographic gate length $L$. Instead, the effective QPC channel appears as a saddle-point potential that has an effective length that is typically shorter than $L$. We used the following approach to define whether localized maxima in the charge density can be interpreted as an ELS within the effective QPC channel. We study the unpolarized solution, and define that the effective QPC channel is located between the two points at a density of 80% of the maximum density, along a QPC cross section as depicted in Fig. S5.

As was discussed in the main text, the ELSs inside the QPC originate from Friedel oscillations and as the QPC gets longer the Friedel oscillations have more periods in the QPC channel. We use two complimentary procedures in order to determine the number of ELSs inside the QPC. First we look at the cross section of the density across the middle of
the sample (this is done separately for spin-up density and spin-down density). The criteria for the definition of a peaked feature that represents an ELS is given by

$$n_\sigma(r_{max}) * P \geq n_\sigma(r_{min}),$$

(8)

$$n_\sigma$$ is the density of spin $$\sigma$$, and $$r_{min}$$ is the closest minimum toward the outside of the QPC. This procedure is shown in Fig. S6 for $$P = 0.32$$. As can be seen in Fig. S6d, this procedure may be problematic when the solution is ferromagnetic, and the peaks overlap significantly. In such cases we use a second procedure: An example is given in Fig. S7. The total two-dimensional density for the up spin (panel b) can be fit to a sum of five Gaussians (only three of them inside the actual QPC channel), each of the total unit weight ($$e$$), even though the cross section reveals only two peaks inside the QPC.
FIG. S2: a, Schematic of the simulation box, the blue rectangles represent the gate electrodes above the 2DEG. b, Example of the external saddle-point and wire potential for a QPC (blue is lowest, red is highest potential).

FIG. S3: Spin-down (left) and spin-up (right) densities. Top row: unpolarized solution. Middle row: spatially-symmetric polarized solution. Bottom row: spatially-anti-symmetric polarized solution. The color scale extends from zero (black) to $2 \cdot 10^{14}$ m$^{-2}$ (yellow).
FIG. S4: Free energies of the symmetric and anti-symmetric polarized solutions relative to the unpolarized solution as a function of gate voltage, for a QPC length of 680 nm.

FIG. S5: Determination of the effective length of the channel in the QPC saddle-point potential (typically shorter than the lithographic gate length $L$). The blue line is a cross section of the total density across the middle of the QPC channel. Above is the total electron density zero (black) to $3.5 \times 10^{14} \text{ m}^{-2}$ (yellow). The red dots mark the density at 80% of the maximum density, and the distance between these points is used as the effective QPC channel length.
FIG. S6: Example for counting ELSs - first method. a, b, Electron density and its cross section for spin up (a) and spin down (b) at $V_g = -6$ and for $L = 540$ nm. The red crosses mark the Friedel oscillations that are counted as ELSs. Here we have a total of two ELSs in the QPC, each one has a density very close to one electron density. c, d, Cross section of spin-up (c) and spin-down (d) densities at $V_g = -6$ and for $L = 830$ nm. In this case there are four ELSs, as can be seen from the cross section.
FIG. S7: Example for counting ELSs - second method. a, b, Electron density and its cross section for spin up (a) and spin down (b) at $V_g = -6$ and for $L = 680$ nm. Here the integration of the density inside the QPC gives a total of three electrons, though there are only two peaks in the cross section. The two dimensional density can be fitted to a sum of 5 unit weight ($e$) Gaussians (all 5 Gaussians have weight $e$ when accounting for their transverse dimension). Three of these Gaussian are within the effective QPC channel (Fig. S5). c, We show how the five Gaussians give the corresponding SDFT density. From top to bottom (and with reference to traces in panel b): Center Gaussian (corresponds to green dashed line in the cross section), two nearest Gaussians (red dashed line), two next nearest Gaussians (cyan dashed line), all five Gaussians together (purple line), SDFT density (blue line). Hence we conclude that there are three localized ELSs inside the QPC.
3. EARLIER REPORTS SHOWING MODULATED 0.7 ANOMALIES AND DOUBLE-PEAK ZBAs

A remarkable aspect of our observations of double-peak ZBAs in many different QPC devices is that such observations were to our knowledge never explicitly reported before, despite about 20 years of research into many-body effects in QPCs (with a few hundred publications reporting on the topic). Similarly, our observations of a periodic modulation of the 0.7 anomaly as a function of QPC channel length is a phenomenon that -to our knowledge- has not been reported before. It is therefore justified to ask why this is the case, and to question whether our -seemingly unique- observations result from an irregularity that is unique to our devices. The main text already reported that we observed double-peak ZBAs in many conventional QPC\textsubscript{2F} devices from two different wafer materials and in 8 different QPC\textsubscript{6F} devices (and for both cases for multiple cool-downs and for different gate settings during cool-down), which all together provides convincing evidence that the new phenomena are not due to a particular device imperfection.

We noticed in our studies on many QPC\textsubscript{2F} that a strong expression of the many-body effects that we report here appears as quantized conductance plateaus that are not flat, and a 0.7 anomaly that appears as a peaked resonance instead of only a shoulder on the step from $G = 0$ to $1 \cdot (2e^2/h)$ (see for example also Fig. S9a near $V_{g1} = -0.35$ V). These two signatures can then also show replicas at and just below the second and third quantized conductance plateau. In such cases, the ZBA often has double or triple-peak character with strong asymmetries and smaller side peaks. Such QPC results were in fact observed long before by our group, but such results (and further studies of the devices) were discarded because a device imperfection was suspected. We know that most groups in the field had the same practice (see for example ref. 17). Also, such imperfections on quantized conductance traces were often removed by applying a small (typically 25 mT) perpendicular magnetic field during the QPC studies, or by measuring at about 600 mK instead of the lowest available temperature (see also ref. 17, not applied in any of our studies). Only our recent study on a large number of QPC\textsubscript{2F} showed us that such strong deviations on quantized conductance traces fall in fact on a regular trend from weak to strong expression of the 0.7 anomaly and the ZBA. Nevertheless, it remains difficult to investigate this systematically since the appearance is very diverse, and for results from one particular QPC\textsubscript{2F} it remains a challenge.
to rule out a (partial) role for an actual material or device defect.

In the remainder of this Section we report that several signatures of double-peak ZBAs and periodic modulation of the 0.7 anomalies can in fact be recognized in the experimental data of earlier publications (but they were never discussed or systematically investigated). Below here we review the literature on this, first for modulation of the 0.7 anomaly and subsequently for double-peak ZBAs. For both phenomena it is also illustrative to inspect all experimental data in the extensive review ref. 18.

**Reports showing modulated 0.7 anomalies**
Several publications report a very regular modulation of the 0.7 anomaly over about 1 period as a function of back(top)-gate voltage in experiments that have a large-area back(top) gate in addition to the split-gate structure of the QPC. A deviation from our experiment is that such a back-gate modulates at the same time the QPC saddle-point potential and the electron density of the reservoirs. A regular modulation of the 0.7 anomaly over about 1 period with a fixed density for the reservoirs was also reported before, in experiments with split-gate QPCs that have an additional side gate or narrow top gate. Examples are ref. 19 (Fig. 2), ref. 17 (Fig. 6-17, left panel on QPC2 on p. 105), ref. 20 (Figs. 2, 3a and 3b), ref. 21 (Fig. 2), and ref. 22 (Fig. 1). The latter three references show in fact modulation over about 1.5 period, with an anomaly at a conductance level of about $0.9 \cdot (2e^2/h)$ that disappears while a new one appears at about $0.6 \cdot (2e^2/h)$ (as in our data). Similar modulation is observed in ref. 23 (Fig. 1).

**Reports showing double-peak ZBAs**
We also found several publications that report double-peak ZBAs for data taken at zero magnetic field. An example that looks much like our data is in ref. 17, Fig. 6-19a on p. 107. Other examples can be found in ref. 24 (Fig. 4) and ref. 25 (Figs. 1a, 1c, 3b and 3d). A very recent publication by Zhang et al. (ref. 23) presents a few examples of double- and triple-peak ZBAs (Fig. 3).
4. REMOTE IMPERFECTIONS AND REDUCING CHARGE NOISE IN QPC DEVICES

Very weak static fluctuations on the QPC saddle-point potentials are to be expected. They can, for example, result from the small device-to-device variation that is inherent to the nanofabrication process. In addition, our devices have a doping layer at about 40 nm distance from the QPC channel. In this doping layer ionized doping centers are at random positions with an average inter-dopant distance of about 10 nm.

Such weak static fluctuations on the QPC saddle-point potentials can have a strong influence on the expression of many-body effects. For example, the parameter \( \Gamma \) that was introduced in the main text for the coupling strength between a localized electron and a neighboring reservoir depends exponentially on weak potential fluctuations that are present between the localized state and the reservoir.

For most of our measurements, we stabilized the dynamical character of the (non)ionized doping centers by cooling down with a small positive voltage on the gates \(^{26}\) (see also Methods).
5. DEFINING AND CALIBRATING $L_{\text{eff}}$ FOR LENGTH-TUNABLE QPCs

As introduced in the main text, operating the QPC$_{6F}$ devices induces a smooth saddle-point potential (as in Fig. 2b) for which the length along the transport direction is controlled by operating at a particular ratio $V_{g2}/V_{g1}$. For such a smooth saddle-point potential it is not obvious what the value is of the channel length. We characterize this channel length with the parameter $L_{\text{eff}}$, which corresponds to the value of the lithographic length of a QPC$_{2F}$ type device (the length $L$ in Fig. 1a) that gives effectively the same saddle-point potential.

The results of calibrating the relation between $L_{\text{eff}}$ and $V_{g2}/V_{g1}$ are presented in Fig. S8. A detailed account of this calibration can be found in a separate publication by our team$^{27}$. This work also investigated whether there is significant structure on the saddle point potential from the narrow gaps (44 nm wide) between the three gate fingers on each side of the QPC channel. The results show that we operate the QPC$_{6F}$ under conditions far away from the point where such structure would become significant. The electron flow is 200 nm away from the gate electrodes such that effective electrostatic potential only reflects the gate geometry in a highly rounded manner.

Opening or closing the transport channel for QPC$_{6F}$ at a particular value for $L_{\text{eff}}$ requires co-sweeping of $V_{g1}$ and $V_{g2}$ at a fixed ratio $V_{g2}/V_{g1}$. Notably, in the case of cooling down with about +0.3 V bias on the gates (see Methods and ref. 26), co-sweeping of $V_{g1}$ and $V_{g2}$ was carried out with respect to $V_{g1} = V_{g2} = +0.3$ V instead of $V_{g1} = V_{g2} = 0$ V (for details see ref. 27).

![Figure S8](image-url)

FIG. S8: Result of calibrating the dependence of $L_{\text{eff}}$ on $V_{g2}/V_{g1}$. 
6. FRIEDEL OSCILLATIONS AND FERMI WAVELENGTH IN THE QPC

The approximate form of a Friedel oscillation for screening of a charged scatter center at position $x = 0$ (valid for positions not too close to $x = 0$) is

$$\rho_{\text{devi}} \propto \frac{\sin(2k_F x + \phi)}{x^d},$$

where $\rho_{\text{devi}}$ is the deviation in electron charge density, $d$ is the dimensionality of the Fermi liquid, $k_F$ is the Fermi wavenumber, and $\phi$ is a phase parameter that depends on the details of the scattering$^{28}$. The factor 2 before $k_F$ shows that the wavelength of Friedel oscillations is half the Fermi wavelength.

The electron reservoirs in our experiment had a density $n_{2D} = 1.6 \cdot 10^{15}$ m$^{-2}$, which corresponds to a Fermi wavelength $\lambda_F = 2\pi/k_F = 62$ nm. In the QPC channel the electron kinetic energy is reduced. For an estimate we assume that the kinetic energy ($E_F = 5.7$ meV for the reservoir) gets reduced in the QPC entries to a value of about 1 meV, which is estimated by taking half the 1D subband energy spacing of our type of QPCs (value taken from our analysis in ref. 29). This yields that $\lambda_F$ increases here to about 150 nm, which corresponds to about 75 nm for the Friedel oscillation wavelength in the QPC channel. When increasing the length of the QPC, the number of Friedel oscillations in the channel increases at the same time (more or less symmetrically) in both entries of the QPC. Thus, a length dependence that relies on the number of Friedel oscillations in the channel should show a modulation with a periodicity of about 150 nm, or a bit smaller ($\sim 100$ nm) if the effective Fermi wavelength in the QPC entries is still a bit closer to the value for the reservoirs. Figure S12 shows that this is well in the range of the observed periodicity. The fact that the modulation occurs faster around $L_{\text{eff}} = 250$ nm than around $L_{\text{eff}} = 500$ nm is consistent with the fact that the latter case corresponds to a longer shallow channel (less abrupt saddle-point potential) where the Fermi wavelength is extended over a longer range.
7. TEMPERATURE DEPENDENCE OF LINEAR CONDUCTANCE

Figure S9 presents for the same device as used for Fig. 2, 3 and 4 in the main text results for the temperature dependence of the 0.7 anomaly in linear conductance traces. At the highest temperature (4000 mK), the linear conductance traces no longer show quantized conductance plateaus (Fig. S9a) and the only remaining feature is the 0.7 anomaly, which no longer shows a modulation as a function of $L_{\text{eff}}$. For the following discussion we focus on gate settings that give $G \approx 0.7 \cdot (2e^2/h)$ at 4000 mK. At these points, the linear conductance increases from 0.7 towards $1 \cdot (2e^2/h)$ when lowering the temperature. Notably, the increase in conductance is for all $L_{\text{eff}}$ for the largest part due to a growing height of the ZBA (observed in the corresponding nonlinear conductance results). Also, subtracting 4000 mK traces from 80 mK traces (defining the traces $\Delta G_T$ in Fig. S9b) shows that the conductance increase is largest around these points. However, some $\Delta G_T$ traces show a suppression (for $V_{g2}/V_{g1} = 0.3, 0.6$ and 1.0 in Fig. S9b) and these points coincide with a strong 0.7 anomaly at the lowest temperatures and pronounced double-peak character for the ZBA. That is, the $\Delta G_T$ curves show that the enhancement of $G$ due to many-body effects is strongest where the linear conductance is about $0.7 \cdot (2e^2/h)$ at high temperatures, but that there is a range within each $L_{\text{eff}}$ period where exactly at this point the strongest expression of a new effect causes in fact a dip in $\Delta G_T$. Further analysis shows that this coincides with the points where the double-peak ZBA shows behavior that is characteristic for the two-impurity Kondo model, and that the energy scale for the coupling between the two spins in this model appears maximum at this point.
FIG. S9: Temperature dependence of the linear conductance. a, Evolution of the linear conductance traces of Fig. 2c as a function of temperature. Traces are displayed for gate ratios $V_{g2}/V_{g1} = 0.0, 0.1, 0.2 \ldots 1.0$, as labeled in b. b, The difference in linear conductance $\Delta G_T$ between the 80 mK and 4000 mK traces of panel a. These traces directly reflect the enhancement of the linear conductance around the 0.7 anomaly with decreasing temperature.
8. MAGNETIC-FIELD DEPENDENCE OF DOUBLE-PEAK ZBAs

Three additional examples for the magnetic-field dependence of double-peak ZBAs are presented in Fig. S10. The dependence on field shows diverse behavior that includes the merging of the two peaks into a single broad peak, after which in some cases a revival of the double-peak character can be observed at higher fields (panels a, b). Panel c presents an example where one of the peaks of the double-peak ZBA at zero field develops a splitting before the full ZBA evolves into a single broad peak.

FIG. S10: Magnetic-field dependence of nonlinear conductance with double-peak ZBAs. a, Evolution of the nonlinear conductance as a function of applied in-plane magnetic field $B_{\text{ext}}$, for the QPC$_{6F}$ of Fig. 3c and Fig. 4, operated at fixed $V_{g1} = -0.646$ V and $V_{g2}/V_{g1} = 0.1$ (giving $L_{\text{eff}} \approx 220$ nm). Subsequent traces (offset $-0.01 \cdot (2e^2/h)$) are for increasing $B_{\text{ext}}$ from 0 to 2.2 T in steps of 0.1 T, with additional traces for an increase in $B_{\text{ext}}$ in larger steps as labeled. b, As panel a, for the device operated at fixed $V_{g1} = -0.403$ V and $V_{g2}/V_{g1} = 0.6$ (giving $L_{\text{eff}} \approx 377$ nm). c, As panel a, for the device operated at fixed $V_{g1} = -0.398$ V and $V_{g2}/V_{g1} = 0.6$ (giving $L_{\text{eff}} \approx 377$ nm).
9. ANALYSIS OF ZBA PEAK POSITIONS AT G = 0.4, 0.6, 0.7 AND 0.85 · (2e²/h)

A detailed analysis of the number of ZBA peaks in nonlinear conductance traces that also quantifies the positions, widths and amplitudes of these peaks is presented in Fig. S12. This analysis was carried out on the data that underlies Fig. 3c. Figure S12 presents results for 4 different conductance levels, as labeled. The symbol size in Fig. S12 is proportional to peak area, which was obtained from fitting Gaussian peak shapes to the ZBA peaks, see Fig. S11 (a phenomenological ansatz suited for extracting values for peak position, width and height). The largest symbols correspond to a peak area of 50 µV · (2e²/h). We mostly observe that the peak width roughly correlates with peak amplitude, such that similar plots with the symbol size proportional to peak width or peak amplitude roughly give the same picture. For the Full-Width at Half-Maximum (FWHM) of ZBA peaks we mainly find values in the range 0.1 to 0.4 mV (or meV for energy scale) for the conductance levels between 0.4 and 0.85 · (2e²/h).

![FIG. S11: Analysis of ZBA peaks. An example of results from fitting two Gaussian peak shapes on a double-peak ZBA. Fits are carried out on peak traces ∆G<sub>P</sub>, that are obtained by subtracting a background conductance level (linear, or parabolic where needed).](image-url)
FIG. S12: Analysis of ZBA peak positions at $G = 0.4$, 0.6, 0.7 and 0.85 · $(2\epsilon^2/h)$ (as labeled). The graphs present results of fitting ZBA peaks, displayed as peak positions (in $V_{sd}$ units) as a function of $L_{eff}$. The area of the symbols is proportional to the peak area (obtained as the product of peak height and FWHM from peak fitting).
10. LINEAR CONDUCTANCE RESULTS FOR DIFFERENT QPC\textsubscript{6F} DEVICES

Figure S13 presents linear conductance traces for a range of $L_{\text{eff}}$ values from 4 different devices (similar results for again another device were presented in Fig. 2c of the main text). These results illustrate that the periodic modulation of the 0.7 anomaly as a function of $L_{\text{eff}}$ was observed in all QPC\textsubscript{6F} that we measured. The results in Fig. S13a-c were obtained on devices where a gate voltage of +0.3 V was applied during cool down. The results in Fig. S13d are from a device that had 0 V on the gates during cool down.

FIG. S13: Linear conductance $G$ traces as a function of $V_{g1}$ for $L_{\text{eff}}$ tuned from a short to a long QPC. The results in a-d are for 4 different devices.
11. FULL DATA SET FOR A QPC$_{6F}$ DEVICE

A full data set of linear and nonlinear conductance for a QPC$_{6F}$ device as a function of $L_{\text{eff}}$ is presented in Fig. S14, as a sequence of 51 paired graphs. Here $L_{\text{eff}}$ is tuned by adjusting $V_{g2}/V_{g1}$ from 0 to 1 in steps of 0.02. This data set is the basis for the (zero-field, 80 mK traces in the) results that are presented in Figs. 2, 3, 4, S9, S10 and S12.
FIG. S14: (figure over next 13 pages) Linear and nonlinear conductance data for a QPC$_{6F}$ device, for $L_{eff}$ tuned from short to long by adjusting $V_{g2}/V_{g1}$ from 0 to 1 in steps of 0.02.
Nonlinear Conductance

Linear Conductance

$V_{sd}$ (mV)

$G$ ($2e^2/h$)

$V_{g1}$ (V)

$G$ ($2e^2/h$)

$V_{g2}/V_{g1}$

Values:

0.480

0.500

0.520

35
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<th>Vsd (mV)</th>
<th>G (2e^2/h)</th>
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Nonlinear Conductance

Linear Conductance

Vg1 (V)

Vg2/Vg1 0.640

Vg2/Vg1 0.660

Vg2/Vg1 0.680

Vg2/Vg1 0.700
Nonlinear Conductance

Linear Conductance

Vsd (mV)

G (2e^2/h)

Vg1 (V)

G (2e^2/h)

Vg2/Vg1

0.72

0.74

0.76

0.78


15. Attaccalite, C., Moroni, S., Gori-Giorgi P. & Bachelet, G. B. Correlation energy and spin


