Emerging magnetic order in platinum atomic contacts and chains

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The development of atomic-scale structures revealing novel transport phenomena is a major goal of nanotechnology. Examples include chains of atoms that form while stretching a transition metal contact or the predicted formation of magnetic order in these chains, the existence of which is still debated. Here we report an experimental study of the magneto-conductance (MC) and anisotropic MC with atomic-size contacts and mono-atomic chains of the nonmagnetic metal platinum. We find a pronounced and diverse MC behaviour, the amplitude and functional dependence change when stretching the contact by subatomic distances. These findings can be interpreted as a signature of local magnetic order in the chain, which may be of particular importance for the application of atomic-sized contacts in spintronic devices of the smallest possible size.

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Bulk platinum shows an enhanced magnetic susceptibility, indicating proximity to ferromagnetism according to the Stoner criterion. Upon reduction of the size down to the nanometre scale, this proximity is enhanced and eventually overcomes the Stoner criterion, rendering Pt ferromagnetic at the nanoscale. First experimental support for this emerging magnetism at reduced dimensions arose from work on Pt clusters, where large magnetic moments were identified that increase inversely proportional to cluster size. A characteristic property of Pt is the formation of mono-atomic chains, when pulling apart a thin Pt wire, as indicated by the appearance of long plateaus in the stretching curves at a conductance corresponding to the single-atom contact. These mono-atomic chains have been thoroughly investigated theoretically as truly one-dimensional model system. It has been predicted that the ground state of an atomic Pt chain is ferromagnetic. Experimentally, however, the existence of local magnetic order in atomic contacts is difficult to measure directly, because the sole occurrence of conductance values at half-integers of the conductance quantum makes a direct measurement of magnetic order in atomic contacts impossible. The same limitation holds for the observation of an enhanced monotonic and hysteretic orbital magneto-conductance (MC) in contacts with cross-sections of several nanometers of the nonmagnetic transition metal Pd.

In recent experiments, the shot noise of Pt atomic chains was studied and no indications of a magnetic ground state were found. A detailed theoretical analysis revealed that this does not exclude the existence of local magnetic order, but the electronic states carrying the magnetic information do not contribute to the shot noise.

For atomic-size contacts of 3d ferromagnets, very large MC and anisotropic MC (AMC) were observed and attributed to spin-dependent scattering. However, different origins of the large MC in magnetic nanocapillaries have been proposed, such as instabilities caused by magnetostriction or quantum interference effects. Nevertheless, a pronounced non-monotonic AMC always requires the existence of a spontaneous magnetization and MC effects caused purely by orbital motion of electrons in non-magnetic metals are much smaller and always monotonic. Atomic Pt chains represent a unique system, where locally ordered, magnetic moments in the chain are coupled directly to paramagnetic leads. This enables a direct study of the magnetic and transport properties of an atomic-size magnet, the properties of which are not induced or dominated by the magnetism in bulky leads.

Here we investigate the magnetotransport of atomic Pt contacts and chains at low temperature. We observe pronounced features in the MC and AMC curves that are subject to subtle changes of the atomic configurations. The analysis reveals the formation of local magnetic moments in the constriction, and therefore indicates a local origin of the observed magnetotransport effects. Our findings thus support so far unconfirmed theoretical predictions. Furthermore, they show a route to create novel atomic-scale electronic devices based on the direct vicinity of magnetic and nonmagnetic subsystems.

Results
Fabrication of single-atom contacts and atomic chains. Here, we investigate atomic contacts by stretching free-standing bridges of Pt fabricated via the mechanically controlled break-junction (MCBJ) technique. A schematic view of the experimental setup and an electron micrograph of a suspended Pt nano-bridge are shown in Fig. 1. Details on the sample preparation and measurement procedures are described in the Methods section. The conductance of a pristine Pt nano-bridge before breaking is typically around 200 $G_0$, where $G_0$ denotes the conductance quantum $G_0 = 2e^2/h$ with the elementary charge $e$ and Planck’s constant $h$. We adjust the cross-section of the junction by carefully stretching the Pt bridge while measuring its conductance. A single atomic contact is created, when the conductance drops below 2.5 $G_0$; further stretching of the contact results in mono-atomic Pt chains. Using this technique, Pt chains of up to seven atoms in length can be created repeatedly. The MC of these contacts and chains is measured at a constant length, sweeping the magnetic field parallel (x direction) or perpendicular (z direction) to the direction of the electrical current. We observe five characteristic features that indicate magnetic order, which are discussed below in separate paragraphs and compared with reference measurements on gold break junctions, where no magnetic ordering occurs in reduced dimensions.

Emergence of manifold shapes in MC of few-atom contacts. First, we record MC traces in magnetic fields up to $|B_z| < 8T$ perpendicular to the sample at 4.2 K. We concentrate on Pt single-atom contacts or chains with $G_0 < 0.9$ to 2.5 $G_0$. For large contacts with $G_0 > 30 G_0$, the MC shows a weak, negative field dependence, as expected for bulk Pt and as shown in Supplementary Fig. 1. For small contacts with cross-sections of few atoms ($G_0 < 5 G_0$), we observe MC curves similar to those found in atomic-size contacts of 3d band ferromagnets. Typical examples of such MC traces are collected in Fig. 2a. The occurrence of MC curves with different shapes and magnitudes serves as a first indication for local magnetic order in Pt chains. To quantify the MC effects, we define the MC ratio (MCR) as the conductance variation relative to the conductance close to zero field: $MCR(B) = (G(B) - G_{ext}(B ≈ 0))/G_{ext}(B ≈ 0)$, where $G_{ext}(B ≈ 0)$ is the extreme value of conductance at low field. Owing to hysteresis effects, which are discussed below, the reference value $G_{ext}(B ≈ 0)$ is not necessarily at zero field. The MCR values indicated in Fig. 2a are defined as the largest deviation in the MC trace from $G_{ext}(B ≈ 0)$ normalized to $G_{ext}(B ≈ 0)$. In Fig. 2a, the stretching length at which the curves have been recorded as well as the zero-field conductance are indicated for each curve. All curves show MC changes occurring...
on a field scale of several Tesla, all are non-monotonic and hysteretic. In the curves I–III and VI additionally the amplitudes of the finite-field extremes depend on the sweep-direction, which is typically observed in magnetic tunnel junctions (MTJs) and is attributed to a spin-polarization of the electrodes\(^1\). However, the field scales are typically much smaller for MTJs of band magnets. Figure 2b,c shows examples of the two most frequently observed MC traces (see Supplementary Table 1 for details about the relative abundance of the shapes). In one case (Fig. 2c), we observe a positive MC with a minimum conductance near zero field, whereas in the other case (Fig. 2b), the MC is negative with a maximum conductance close to zero. The insets depict a simple model we develop herein, which indicates how an external magnetic field may influence the local magnetic moments of a contact, leading to the observed MC. This model is described in detail in the Discussion section.

Sensitivity of MC to atomic configuration. Next we follow the development of the MC traces while stretching a mono-atomic chain, see Fig. 3. When arriving at the single-atom contact regime\((G<2.5 \ G_0)\), we interrupt the stretching at intervals of about 0.11 nm, which corresponds to 38% of an atomic diameter, to record series of MC curves at each fixed chain length. Figure 3 shows the evolution of the conductance in zero field (Fig. 3a) as well as the amplitude \(\text{MCR}_{\text{max}}\) (Fig. 3b) and hysteresis \(\Delta\) (Fig. 3c) of the MC curves; the stretching intervals are highlighted as alternating shaded regions. The multiple symbols for \(\text{MCR}_{\text{max}}\) and \(\Delta\) at a fixed chain length in Fig. 3b and c correspond to subsequent magnetic field sweeps, in which we observe only a small variation of the curve shape, hysteresis and \(\text{MCR}_{\text{max}}\). The MC-shapes corresponding to the \(\text{MCR}_{\text{max}}\)-Values labelled with roman numerals (I–VIII) in Fig. 3b are shown in Fig. 3d. The MC traces VII and VIII in Fig. 3d were recorded at the same nominal chain length of 3.8 atoms but both are drastically different. We observed such a large difference in subsequent MC measurements in particular for ultimately stretched Pt chains and attribute this behaviour to a lower stability and a larger sensitivity of these chains to magnetic field-induced variations of the contact geometry. Owing to this reduced stability the contact breaks while extending the nominal chain length above 4.1 atoms.

While extending the length of mono-atomic chains starting from the single-atom contact, the sign, amplitude \(\text{MCR}_{\text{max}}\) and hysteresis \(\Delta\) as well as the shape of the MC traces (Fig. 3d) change within subatomic variations of the chain length, whereas the overall conductance varies only slightly. Even a small variation of the electrode separation can induce changes in the atomic contact configuration. However, in single-atom contacts and chains, the conductance is rather independent of length. Hence, when an additional atom is extracted from the electrodes to join the mono-atomic chain, the bond-length between the chain atoms and the electrodes change, whereas the absolute conductance
remains fairly constant. However, the magnitude, anisotropy direction and coupling of magnetic moments in the chain are very sensitive to the bond length $8,26,27$. The observed variation of sign and shape of the MC curves upon stretching the contact by fractions of an atomic diameter hallmarks this high sensitivity. In the example shown in Fig. 3b, the MCR max values vary between 2 and $\pm 20\%$. For other chains, similar results and MCR max up to $\pm 40\%$ were found (see Supplementary Fig. 2). Both the large amplitude of the MC and its sensitivity to sub-atomic length changes point towards a local origin of the MC rather than a bulk effect and serve as a second indication for magnetically ordered Pt chains.

Hysteresis of magnetoconductance curves. Most MC traces recorded for Pt single-atom contacts and chains reveal a hysteretic behaviour of the conductance with a local extreme $G_{\text{ext}}(B) \approx 0$ located symmetrically around zero field, as indicated by the red and blue curves in Fig. 4. Here we define the hysteresis $\Delta$ as the deviation of position of this local extreme from zero field, $\pm 5\%$ to $\pm 20\%$ in all panels, the dashed line indicates zero magnetoresistance amplitude (MCR).

Figure 3 | Development of the magneto-conductance during evolution of a Pt chain. (a) Stretching curve of a Pt bridge interrupted at regular intervals to measure the magneto-conductance (MC). The sample was opened continuously until the single-atom contact regime at conductance $G < 2.5 G_0$ was reached. Then several MC traces in perpendicular ($B_z$) fields up to $\pm 8$ T were recorded at fixed chain length, while stepwise opening the contact at intervals of approximately 2/5 of a Pt atomic diameter in a chain as indicated by the shaded regions. The insets sketch the atomic chain (cyan spheres) coupled to the electrodes (blue) via apex atoms (purple) as a number of Pt atoms equivalent to the electrode assuming a rigid chain with a bond-length of 2.33 Å. $b$ shows the maximum MC ratio ($\text{MCR}_{\text{max}}$), and the hysteresis ($\Delta$) is shown in $c$; these values are extracted from individual MC traces exemplified in $d$. The symbols in $\text{MCR}_{\text{max}}$ and $\Delta$ represent consecutive measurements at fixed opening positions. $\text{MCR}_{\text{max}}$ varies between values close to zero and up to 20% as the chain is created. Moreover, the MC continuously changes its sign depending on the atomic configuration of the contact. The hysteresis also changes non-monotonically with the chain length between 20 and 300 mT. (d) Out-of plane MC traces for different contacts recorded along the stretching curve at positions I–VIII used to extract the $\text{MCR}_{\text{max}}$ and $\Delta$ values shown in $a$. The horizontal axis spans the range from $\pm 5\%$ to $\pm 20\%$ in all panels, the dashed line indicates zero magnetoresistance amplitude (MCR).

Figure 4 | Emerging hysteresis in an atomic Pt contact. After adjusting the contact the MC shows an extreme at zero while ramping the magnetic field the first time (cyan curve). After application of a high field, this minimum is shifted by about 350 mT in sweep direction when performing a magnetic field sweep. Please note that the curves have been shifted vertically for a better visibility. The cyan curve has been measured right after adjusting the contact when sweeping the field from 0 to $\pm 8$ T, then the blue curves when sweeping from $\pm 8$ T to $\pm 8$ T and finally the red curve when sweeping from $\pm 8$ T up to $\pm 8$ T.
as indicated by the vertical markers in Fig. 4. The numerical values of the hysteresis are calculated by fitting a quadratic polynomial to the area around the central extreme $G_{\text{ext}}(B \approx 0)$, the vertex of the polynomial marks the hysteresis, whereas the standard deviation of the best-fit result represents the error bars as indicated by the vertical markers in Fig. 4. The general shape of the AMC remains unaffected but its sign changes. This behaviour is incompatible with the bulk theory of AMC\textsuperscript{16}, but can be explained by spatial asymmetries and by spin polarization of the conduction channels\textsuperscript{24}. This model put forward for atomic contacts of band ferromagnets predicts a manifold AMC behaviour, including sign changes due to atomic reconfiguration and large amplitudes that are similar to our experimental findings. Therefore, the large AMC and the observed sign changes strongly imply that the conduction channels in Pt contacts are indeed spin-polarized by the emergence of a local magnetically ordered state.

**Dependence of MC on magnetic history.** The fifth indication for a local magnetic order is the observation of history- and time-dependent in-plane MC (x-direction). We observe a symmetric, parabolic MC (green and black traces in Fig. 5b), if only a moderate $|B| < 2.5$ T field is applied before measuring the in-plane MC, where the magnetic field is applied parallel to the current. However, if a strong, perpendicular field of $|B_z| > 7$ T was applied before recording the in-plane MC, a highly asymmetric and strongly hysteretic behaviour is observed (blue and red traces in Fig. 5b). The extreme values of the up- and down-ward sweeps occur at fields, which are in the same order of magnitude as the hysteresis observed in the out-of-plane MC traces, suggesting that the in-plane MC is governed by the magnetic properties of the electrodes. The MC relaxes, at the low temperature of our measurement over time scales of several hours, pointing again towards a local magnetic order\textsuperscript{16} rather than a bulk effect. The relaxation of the MC over such long time scales is interpreted as a relaxation of the perturbed magnetization state after the application of a large out-of-plane field to its ground state, where the magnetic moments are preferably aligned in-plane.

**Reference measurements on gold atomic contacts.** The general MC behaviour of Au differs drastically from those of Pt, although...
both metals form mono-atomic chains when stretching a single-atom contact. A collection of MC traces for Au single-atom contacts and chains is shown in Fig. 6a. Most traces reveal small MCR_{max} < 1% and changes of MC on much smaller field scales, which are reminiscent of conductance fluctuations in atomic contacts. The amplitude is not expected to show the universal value because of the non-homogenous voltage drop in atomic contacts. Similar to Pt, we find a large diversity of MC shapes and amplitudes, which occasionally change sign. Overall, there is a clear tendency for negative MCR_{max}. Nevertheless, similar shapes than in Pt chains can be identified, but the amplitude of the MC effect is approximately one order of magnitude smaller. For example, the shape of trace III in Fig. 6a shows similarities to trace III in Fig. 2a. Despite the occurrence of similar shapes of MC traces in Pt and Au atomic contacts and chains, in the case of Au we did not find a distinct curve shape that is more prominent than others. This is in contrast to Pt, where the two traces shown in Fig. 2b,c represent the most abundant form of MC traces observed in these contacts. In Au contacts and chains with a total conductance near 1 G_0 the MCR_{max} values hardly exceed 1%. This changes when the zero-field conductance differs markedly from multiples of G_0, that is, when partially opened conduction channels contribute, or when the conductance drops below 1 G_0. For a contact of 0.6 G_0 we found a MCR_{max} value of more than 1.85%, see Fig. 6a trace VIII. These findings are in agreement with earlier reports of conductance fluctuations of Au atomic contacts and changes studied upon sweeping the bias voltage.

As for Pt, we study the evolution of the MC and its hysteresis while extending a Au chain, starting from the single-atom contact. Again, the chain length is adjusted and then kept constant, while a series of MC traces is recorded before stretching the chain further. Figure 6 shows an example of the development of the zero-field conductance (Fig. 6b), the MCR_{max} (Fig. 6c) and the hysteresis (Fig. 6d) of a Au chain. Like for Pt chains, occasionally we observe an instantaneous switching behaviour on Au chains as well, which can be interpreted as arising from atomic rearrangements, but leading to a comparably small MC of 2.6% compared with 30% on Pt, see trace IX in Fig. 2a. To determine the hysteresis, the centre parts of the sweeps are evaluated in the same manner as for Pt. However, generally the MC traces of Au chains are only weakly or non-hysteretic. The exact evaluation shows Δ in the range of 20 mT, which occasionally increases up to 80 mT, however, the error bars are much larger than in the case of Pt. In summary, the MCR in Au is extremely small and non-hysteretic, while universal conductance fluctuations account for most of the conductance variation observed in these contacts. A larger set of opening and MC traces for Au is shown in the Supplementary Fig. 3. The fact that no clear MC is observed for Au samples proves that the large MC observed in Pt contacts is no artefact of the measurement setup and supports the argument that in Pt mono-atomic chains a local, magnetically ordered state develops.

**Discussion**

Although the experimental observation presented above clearly indicate the formation of magnetic order in atomic Pt contacts and chains, the exact interpretation of the shapes of the MC curves remains a difficult task. We propose a qualitative understanding: following the results of theoretical calculations for atomic contacts of band ferromagnets, we assume that a
parallel alignment of the magnetization gives rise to the highest conductance, whereas a misalignment reduces the conductance\textsuperscript{33}. This approach is similar to Julliere’s model for MTJs\textsuperscript{10}. In contrast to MTJs, where the field scales are set by the coercivity of the electrodes separated by a nonmagnetic barrier, in Pt atomic contacts the magnetism, if present, is created locally on atomic length scales. Different magnetization directions of mono-atomic Pt chains have been predicted with magnetic moments per atom ranging from 0.2 to several 1\mu B (refs 8–10,26,27,34,35). Most calculations agree that the magnetization in the ground state is parallel to the chain axis (insets Fig. 2b), but for certain bond lengths, which can be realized by stretching a chain, the magnetization may become perpendicular to the chain axis\textsuperscript{26,27,35}. Very high values of more than 100 meV were predicted for the magnetic anisotropy energies (MAEs) of the chain and apex atoms, but with varying magnetization directions\textsuperscript{26,27}. Within these calculations, the energetic ground states of the individual atoms for different bonding lengths in the chain were determined separately, before including their interaction. Although the magnetic moment of the central atom is generally pointing in chain direction, those of the apex atoms preferentially lie perpendicular to it. Finally, the coupling among these atoms and the coupling to the electrodes lead to a reduced MAE in the range of several tens of meV for the entire chain, which corresponds to a Zeeman field of \textgtrsim 30 T (refs 26,33). As a result, for small atomic distances, the competing energy scales are of comparable size and might thus lead to a non-collinear magnetization in the ground state (insets Fig. 2c)\textsuperscript{26,36}. Although such a ground state has not been considered explicitly for pure Pt samples\textsuperscript{26}, the mechanism of strong spin-orbit coupling of Pt leading to a spin torque on the magnetic moments of small clusters and chains of 3d band magnets\textsuperscript{36} should be the same for magnetic moments in atomic Pt chains. The structure of magnetic moments in atomic Pt contacts might be even more complex because the calculations did not take into account the influence of the conical leads\textsuperscript{9,27,35}, which have been shown to carry a finite magnetic moment\textsuperscript{37} that is weakly pinned but affects the magnetization of the apex atoms. Furthermore, we assume that the magnetic moments of the chain, apex and central atoms polarize the innermost parts of the electrodes within a range of several atomic layers\textsuperscript{26,38}. Following these considerations, the total MAE is reduced and moderate magnetic fields \textless 10 T should be sufficient to reorient the moments of the electrodes and the apex atoms, while the central chain atoms remain strongly pinned along the chain direction and would not be affected by the magnetic fields accessible experimentally.

Acknowledging the possibility of non-collinear magnetization, we depict a possible scenario which qualitatively describes the MC curve of Fig. 2b: in the ground state, all magnetic moments are aligned along the chain axis. Applying an external field perpendicular to the chain axis results in a relative misalignment between electrodes and chain at field scales corresponding to the anisotropy energy of the electrodes, roughly 400 mT in this case, and thus to negative MC. The conductance change occurs gradually rather than abruptly as in MTJs because several energy scales including the anisotropy energies of the components of the device and the coupling between them are involved. When the magnetic field is increased further to overcome the anisotropy energy of the apex atom (here: 5.5 T), their moments are pulled out of the chain axis and depending on the combined contribution of AMC and conductance change due to misalignment, the conductance may decrease further, remain constant (as, for example, observed in curve V of Fig. 2a) or increase as in Fig. 2b. In the case of a non-collinear ground state, for example, if, at zero field, the chain atoms, one electrode and one apex atom are magnetized in plane, while the other apex atom and the adjacent electrode are magnetized out of plane (inset Fig. 2c), the application of a moderate field initially results in a positive MC, until the electrodes are aligned parallel with respect to each other. At large fields \textgtrsim 6 T, the moment of the in-plane apex atom may reorient out of plane and due to the misalignment with the moments of the central chain atoms, the conductance decreases again. The sweep-direction-dependent conductance values (curves I–III and VI in Fig. 2a) can be explained assuming a non-collinear arrangement of the apex atoms in the ground state. The appearance of additional extremes as exemplified by traces III and IV may be understood by the subtle interplay between the MAEs of apex, chain and electrode moments. Using this simple model, based on the reorientation of magnetic moments in the chain at different field scales, an interpretation of all MC curves in Fig. 2a is presented in Supplementary Fig. 4 and the Supplementary Discussion. In conclusion, our comprehensive set of observations in atomic Pt chains clearly indicates the formation of local magnetic moments in the constriction. The magnetic field influences their coupling and thereby also the conductance characteristics in the magnetotransport that are subject to subtle changes of the atomic configurations. Our findings do not only confirm the longstanding theoretical predictions of local magnetism in atomic chains, but they also reveal why subtle differences in the theoretical model may lead to different predictions for the magnetic order and give valuable input for the still lacking calculations of magnetotransport in these structures. The proposed interpretation model and the occurrence of diverse MC shapes imply that a non-collinear alignment of magnetic moments in atomic Pt contacts is a very common feature. The possibility of creating local magnetic order in ultimate proximity to nonmagnetic electrodes is particularly interesting for the development of atomic-scale magnetoelectronic and spintronic devices, because spin-polarized currents can be created and detected on the atomic scale in ultimate vicinity to nonmagnetic devices.

**Methods**

**MCBJ technique.** The cross-section of the junctions required for the experiments described herein must be continuously tuneable in-situ. The samples need to provide sufficient long-time stability to hold a single-atomic contact over periods up to several days. The first requirement demands for high flexibility, whereas the latter one is mandatory to perform magnetotransport measurements without changing the geometry by mechanical excitation, heating, magnetostriction or other effects. We chose the method of lithographically defined MCBJ because they present a good compromise between tuneability and stability. In principle, also STM-based junctions are suitable candidates with the benefit that information about the atomic arrangements of the surroundings can be determined. However, the low stability of the STM setup, in particular when applying external magnetic fields in various directions, limits this approach to relatively short-term studies of large numbers of contacts. In contrast, the MCBJ technique has the advantage that single-atomic contacts can be stabilized over extended periods of time without large perturbation from external factors, but the high stability also means that statistical measurements become very time consuming.

**Standard lift-off process for Au.** A layer of polyimide (2 \textmu m thick) is spin-coated onto a polished bronze wafer (200 \textmu m in thickness). This layer serves as an electrical insulator and a sacrificial layer in the subsequent etching process. On top of the polyimide, a double layer of electron-beam resists, methyl methacrylate-methacrylic acid (MMA-MAA)/polymethyl methacrylate (PMMA), is deposited by spin coating (maximum 5,000 r.p.m.). Then electron-beam lithography (EBL) is performed on the prepared wafer. After developing the resist, a layer of Au is deposited with a thickness of 80 nm via electron-beam evaporation at a pressure of 10\textsuperscript{-8} mbar. Redundant parts of the metal film are removed by lift-off in acetone at 50°C. To finalize the samples, they are dry etched in oxygen plasma of 50 W in a reactive ion etcher, which results in a free-standing bridge\textsuperscript{19}.

**Subtractive patterning process for Pt.** The fabrication of Pt samples by using the lift-off process is not very reproducible. Because of the high boiling point of Pt, the PMMA is modified during evaporation, causing the lithographically defined constriction to collapse. Cooling the sample with liquid nitrogen during evaporation
process did not result in improved quality of the samples. The main reason is that owing to the different thermal expansions of the polyimide and the metal film, the construction blisters while warming up the sample after evaporation. To overcome this problem, we use subtractive patterning via dry etching. A pristine Pt layer deposited before the EBL process on a polyimide layer via sputter deposition (base pressure ~10^-9 mbar). The double-layer electron-beam resist (MMA-MAA/PMMA) is deposited on the Pt film in order to perform the EBL. After developing the resist, an Al layer with a thickness of 30–35 nm is deposited via thermal evaporation at a pressure of ~10^-6 mbar. After lift-off in acetone at 50°C, the samples are placed in a reactive ion etcher, where a highly anisotropic etching process at 150 W with sulphur hexafluoride (SF6) as a reactive gas is performed. A scheme of the experimental setup and an electron micrograph of a suspended Pt nano-bridge are shown in Fig. 1. The conductance histograms of Pt, embedded in a polyimide host, show a distribution reminiscent of the findings for PtCo (ref. 40). We attribute these effects to structural defects in the constriction of the Pt samples. Without EM, the opening few seconds to remove the residual Al and then dried with N2 gas.

Electromigration treatment. After cool-down, before breaking the first time, we perform a soft electromigration (EM) process to remove impurities and heal out structural defects in the constriction of the Pt samples. Without EM, the opening process did not result in improved quality of the samples. The main reason is that the opening process at 150 W with sulphur hexafluoride (SF6) as a reactive gas is performed. A pristine Pt layer is deposited before the EBL process on a polyimide layer via sputter deposition (base pressure ~10^-9 mbar). The double-layer electron-beam resist (MMA-MAA/PMMA) is deposited on the Pt film in order to perform the EBL. After developing the resist, an Al layer with a thickness of 30–35 nm is deposited via thermal evaporation at a pressure of ~10^-6 mbar. After lift-off in acetone at 50°C, the samples are placed in a reactive ion etcher, where a highly anisotropic etching process at 150 W with sulphur hexafluoride (SF6) as a reactive gas is performed.

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One can identify several regimes in the EM process defined by threshold and power reduction. If the threshold is low or the power is reduced to small values, the constriction simply heats up and cools down at temperature scale too low to trigger structural changes. If the threshold is high or the power is reduced only slightly, the sample heats up very quickly and the process becomes unstable, which in many cases leads to a destroyed sample via melting or evaporation of the constriction. In the intermediate regime, both the desired healing effect and a controlled opening take place. The parameter range varies from sample to sample and may change for a given sample depending on the history of the EM process. We found the healing process to be completed after a resistance change of 5–8% relative to the initial resistance. When continuing to perform the EM, either no further resistance change was observed or an increasing resistance signalling a thinning process without further reduction of impurity concentration.

Magnetotransport measurements. The data set comprises measurements of the MC in two different directions of the applied field with respect to the sample plane and measurements of the AMC in a wide range of conductance values from 50 G0 to 0.001 G0. The measurements are performed at low temperature (~6 K) while continuously changing the contact geometry on the atomic scale. Besides the adjustable field from the external solenoid, the setup features a substrate reconstructions. The optimum constriction is reached again. This procedure is repeated until a total conductance decrease of 6–8 orders of magnitude. A sketch of the MCBJ mechanism is shown in Supplementary Fig. 6. For all measurements, both dc conductance and measurements of the AMC in a wide range of conductance values from 50 G0 to 0.001 G0.

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Author contributions
F.S., C.E. and M.B. performed the experiments. T.P. and E.S. planned the project and advised the students. All authors discussed the results and prepared the manuscript together.

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