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Article
Control of Intramolecular Electron Transport Pathways by Varying Localized Electron Distribution

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Abstract

The control of different electron transport pathways by quantum interference (QI) effects offers a unique opportunity for the modulation of electrical properties in molecular electronic devices and materials. In this work, we propose a chemical way to control the intramolecular electron transport pathways by the localization of the highest occupied molecular orbital (HOMO) distribution. The negative charge injection in para-carbazole by deprotonation exhibited a fourfold suppression of single-molecule conductance, while the conductance is almost the same for meta-carbazole before and after deprotonation. The flicker noise analyses and theoretical simulations revealed the localized distribution of HOMO on the para-carbazole center, leading to the appearance of destructive quantum interference (DQI) effect for the control of electron transport pathway. This strategy of reaction-induced orbital localization offers a new strategy for the control of charge transport through molecular devices and materials.

Introduction

The electron transport through single-molecule junctions represents the overall contributions of all transport pathways.1-4 In different electron transport pathways, the phase of transmission amplitudes is different, resulting in different quantum interference (QI) patterns to form various conductance states.5-8 The QI effects can be regulated through structural modification, such as connection site variation9-11 and side group substitution,12-15 which offers the opportunity for the control of specific pathways. However, because of the strong coupling between the molecular energy level and the
Fermi energy level of electrodes, the frontier orbital levels of the molecule are typically overextended, thus inhibiting the formation of destructive quantum interference (DQI).\textsuperscript{16, 17} To demonstrate an anti-resonance feature from the DQI for the suppression of charge transport,\textsuperscript{18} the localization of electron density distribution of frontier orbitals will be a potential option to avoid this strong coupling.

To reduce the electrode-molecule coupling, the localized distribution can be achieved by chemical design \textit{via} creating the absence of electron density cloud on the connection sites of frontier molecular orbitals.\textsuperscript{19} Recent works suggest that the electrode can be decoupled by localizing the lowest unoccupied molecular orbital (LUMO) distribution to form effective DQI to create highly nonlinear I-V features,\textsuperscript{19} and also to form the electron-hopping pathway responsible for voltage-induced switching.\textsuperscript{20} However, it remains challenging to directly switch between localization and delocalization in single-molecule junctions, the chemical reactions to modify the molecular structure \textit{in-situ} hold a promising way. The chemical reactions, such as isomerization,\textsuperscript{21} acidification,\textsuperscript{22} photothermal reaction,\textsuperscript{23} and redox reaction,\textsuperscript{24} provide potentials to change the structure of the molecule and the distribution of electron density cloud. Therefore, regulating the distribution of molecular orbitals through \textit{in-situ} chemical reactions provides a new strategy for controlling the electron transport pathways.

Herein, we synthesized two series of molecules based on carbazole as the conductive core (Figure \textbf{1a}), and investigated their single-molecule conductance by the scanning tunneling microscope break junction (STM-BJ) technique.\textsuperscript{25, 26} We found that the negative charge injection in \textit{para}-carbazole by deprotonation exhibited a fourfold suppression of single-molecule conductance, while the conductance is almost the same for \textit{meta}-carbazole before and after deprotonation. Further flicker noise measurements and the theoretical simulations, including density functional theory (DFT) combined with non-equilibrium Green’s function (NEGF) and electron transport pathway calculations, demonstrated that the localized distribution of HOMO on the \textit{para}-carbazole center leads to the appearance of DQI effect and the control of intramolecular electron transport pathways.
Results

![Molecular structure and $^1H$ NMR of para-carbazole and meta-carbazole.](image)

**a**, Molecular structure and reaction process of carbazole derivatives. **b**, Schematic of the single-molecule junction, the hafnium dioxide dielectric layer was coated on the gold tip, and the inset indicates two electrical transport pathways in the carbazole ring, C-C and C-N-C. **c**, Monitoring the complete transformation from Para-N-H to Para-N-Me through $^1H$ NMR. **d**, Monitoring the complete transformation from Meta-N-H to Meta-N-Me through $^1H$ NMR.

The carbazole compounds are a class of diphenylamine molecules with an isoelectronic structure, which possesses strong electron-donating and hole transport abilities. The proton on the nitrogen atom can be removed through alkalization to create a localized electron density distribution on the carbazole center, and then recovers by acidification, which acts as an ideal platform for regulating electron transport pathways through an *in-situ* chemical reaction. In addition, to quench the negative charge, methyl trifluoromethanesulfonate (MeOTf) was also used to methylate the carbazole anion derivatives to confirm the formation of the localization further.

Compound Para-N-H and Meta-N-H were prepared through the reported procedures (more details in supporting information (SI) section 1). To accurately achieve the
deprotonation of the carbazole structure, potassium tert-butoxide (t-BuOK) was used as the base to remove the active hydrogen on the nitrogen atom of carbazole, forming a nitrogen anion, which is an extremely hydrogen affinity state. The NMR spectra show that the active hydrogen signal at 11.3 ppm disappears with the addition of base (indicated by the black frame in Figure 1c and 1d), while the chemical shifts of other hydrogens move to high-field, which proves the accomplishment of deprotonation. Afterwards, the negative charge can be quenched by methylation with chemical shifts of other hydrogen returning to their original positions and the signal vanishing at 11.3 ppm meanwhile, suggesting the completion of quenching. Single-molecule conductance measurements were conducted for the para-carbazole and meta-carbazole by using the STM-BJ technique in the solution of N,N-dimethylformamide (DMF) containing 0.1 mM target molecules at the ambient temperatures.

In the investigation of these two states using single-molecule conductance measurement, once high polar DMF as the solvent and a strong base of t-BuOK were used, rendering a strong leakage current in this solution environment between the bare Au tip and the Au substrate. It results in a high electrical background and makes conductance testing impossible (more details in Figure S15a and S15b from SI). Therefore, to realize conductance measurement in such an ionic solution system with high polarity and large tunneling background, the tip coated with hafnium dioxide (HfO$_2$) as the insulating protective layer by Atomic Layer Deposition (ALD) was adopted in the STM-BJ technique. By depositing HfO$_2$ with a high dielectric constant on the Au tips, we achieved intensive tip coating to significantly reduce the leakage current and the electrical background, which resulted in a larger conductance testing window (more details in SI section 4.2).
Figure 2 | Conductance measurement results of carbazole derivatives. One-dimensional (1D) conductance histograms of (a) para-carbazole and (e) meta-carbazole. Two-dimensional (2D) conductance histograms versus relative displacement and corresponding distributions (inset) of (b) Para-N-H, (c) Para-N-e, (d) Para-N-Me, (f) Meta-N-H, (g) Meta-N-e and (h) Meta-N-Me.

Figure 2a and 2e demonstrate the one-dimensional (1D) conductance histogram comparisons of the para-carbazole and meta-carbazole, respectively, where the peak at $G_0$ (quantum conductance, $G_0 = 2e^2/h = 77.5 \ \mu$S) illustrates the formation of Au-Au atomic point contact. From the 1D conductance histogram, the single-molecule conductance is determined to be $10^{-4.36} G_0$ (3.39 nS) for Para-N-H and $10^{-4.96} G_0$ (0.85 nS) for Meta-N-H. After the in-situ deprotonation by injecting 1 eq. of 1 mM t-BuOK, the conductance of deprotonated Meta-N-e becomes $10^{-4.85} G_0$ (1.10 nS), which is slightly higher than that of Meta-N-H. However, the conductance of the deprotonated Para-N-e decreases to $10^{-4.96} G_0$ (0.85 nS), which is only a quarter of that of Para-N-H. To verify that the significant conductance difference is related to the negative charge on nitrogen atoms, the alkylation reagent of 1.5 mM MeOTf was added to the solution. After alkylation, Meta-N-Me and Meta-N-e exhibit almost the same conductance, which is $10^{-4.84} G_0$ (1.12 nS). However, the conductance of Para-N-Me is significantly higher than that of Para-N-e, which is $10^{-4.49} G_0$ (2.51 nS) and almost close to that of Para-N-H. Therefore, this conductance change must be directly related to the negative charge on nitrogen site. Previous researches have shown that the QI effects can be affected by regulating the charge density of the nitrogen atom on the carbazole ring to change molecular conductance. The charge density of nitrogen atoms can be
characterized by the proton binding ability of nitrogen atoms ($p$Ka). Under acidic and neutral conditions, the molecular conductance increases with the increase of $p$Ka. However, in this experiment, after deprotonation, **Para-N-e** and **Meta-N-e** actually have a strong proton binding ability with large $p$Ka. We found that under alkaline conditions, the conductance of *meta*-carbazole after deprotonation moderately changed, and the regulation effect of $p$Ka reached to saturation, while the conductance of *para*-carbazole after deprotonation decreased, showing an opposite trend compared to previous results.

To seek the origins of the conductance variation, the two-dimensional (2D) conductance and displacement histograms of target molecules are analyzed. The displacement distribution histograms show that the junction lengths of molecules **Para-N-H** (Figure 2b), **Para-N-e** (Figure 2c), **Para-N-Me** (Figure 2d), **Meta-N-H** (Figure 2f), **Meta-N-e** (Figure 2g) and **Meta-N-Me** (Figure 2h) are 1.37, 1.36, 1.37, 0.92, 0.94 and 0.91 nm, respectively. With the Au-Au snap-back correction of 0.5 nm, the junction lengths of molecules are 1.87, 1.86, 1.87, 1.42, 1.44 and 1.41 nm, respectively, which sufficiently match the S-S distances in the molecules by theoretical calculations (more details in Table S1 from SI), as shown in Table 1. The junction length correction indicates that the molecular backbone length remains unchanged after deprotonation and alkylation. In addition, we directly synthesized the molecule of **Para-N-Me** and **Meta-N-Me**, by which the test results showed that both the conductance and junction lengths of molecules were consistent with those of the **Para-N-Me** and **Meta-N-Me** formed through *in-situ* alkylation (more details in Figure S16 from SI). Furthermore, to prove the reversibility of this conductance switch, we added an equivalent amount of 1 mM trifluoroacetic acid (TFA) to the solution to neutralize **Para-N-e** into **Para-N-H**, and found that the conductance was reversed. The cycles of protonation and deprotonation can be repeatedly achieved, showing that it is feasible to control the negative charge injection by deprotonation to switch the conductance, as shown in Figure 3a (more details in Figure S17 from SI). Remarkably, the successive addition of t-BuOK and TFA leads to the formation of a large number of ions in the solution, and
experiments show that the HfO$_2$ coated tip is still suitable for measuring molecular conductance signals in this type of solution with highly concentrated salts.

**Figure 3** | The conductance cycles measurement and the flicker noise analysis of *para*-carbazole. **a**, Reversible on-off conductance switching cycles of *Para-N-H* and *Para-N-e* by deprotonation. Two-dimensional histogram of normalized conductance changes versus normalized noise power for **b** *Para-N-H*, **c** *Para-N-e*, **d** *Para-N-Me*.

To further investigate the electrical transport properties before and after deprotonation, the flicker noise measurement was performed. According to the conductance results above, the molecular junction elongation was hovered for 200 ms to extract the conductance signals within the period for flicker noise analysis.$^{42,43}$ We find that the scale of the noise power of *Para-N-H* and *Para-N-Me* is $G^{1.5}$ as shown in Figure 3b and 3d, indicating that the charge transport through *Para-N-H* and *Para-N-Me* is synchronously dominated by through-bond and through-space transport. In contrast, for *Para-N-e*, the noise power of $G^{2.0}$ indicates that the through-bond is dominant as shown in Figure 3c.

The *Para-N-e* and *Meta-N-e* solutions show obvious color changes and different fluorescences before and after deprotonation (more details in Figure S8 from SI). To understand the impact of the injection of negative charge caused by deprotonation on the energy level, we tested the UV-Vis absorption spectra and the fluorescence emission spectra. For *Para-N-e* and *Meta-N-e*, compared with *Para-N-H* and *Meta-N-H*, a new
absorption peak appear at 370 nm and 385 nm (Figure 4a-4b), respectively, and the emission spectra show obvious redshift (Figure 4d-4e), indicating the shrinkage of energy gaps.\cite{28,44} After methylation, the UV absorption peaks and the fluorescence emission peaks of Para-N-Me and Meta-N-Me are basically the same as those of Para-N-e and Meta-N-e.\cite{45} To further reveal the changes in molecular energy levels before and after deprotonation, density functional theory (DFT) calculations were employed to perform theoretical studies.\cite{46} Frontier molecular orbitals, which are directly correlated to the charge transport, were calculated for all the investigated molecules. Overall, the energy of HOMOs and LUMOs increase with the injection of the negative charge as shown in Figure 4c. However, the increase of LUMO energy is less than the increase of HOMO energy, which results in narrower HOMO-LUMO gaps. The calculated energy level changes are consistent with the results of macroscopic spectroscopic measurements.

**Table 1.** Conductance, length and the scale of the noise power from experimental measurements.

| Molecule   | Measured conductance | Length of molecular junctions / nm | The scale of the noise power |
|------------|-----------------------|-----------------------------------|-----------------------------|
| Para-N-H   | $10^{-4.36} G_0$ (3.39 nS) | 1.87                              | 1.5                         |
| Para-N-e   | $10^{-4.96} G_0$ (0.85 nS) | 1.86                              | 2.0                         |
| Para-N-Me  | $10^{-4.49} G_0$ (2.51 nS) | 1.87                              | 1.5                         |
| Meta-N-H   | $10^{-4.96} G_0$ (0.85 nS) | 1.42                              | 1.1                         |
| Meta-N-e   | $10^{-4.85} G_0$ (1.10 nS) | 1.44                              | 1.1                         |
| Meta-N-Me  | $10^{-4.84} G_0$ (1.12 nS) | 1.41                              | 1.1                         |
Figure 4 | The spectroscopic characterization and theoretically calculated frontier energy of para-carbazole and meta-carbazole. UV-Vis absorption spectra of (a) para-carbazole and (b) meta-carbazole. (c) Theoretically calculated frontier energy and energy gaps between the LUMO and HOMO of para-carbazole and meta-carbazole. The fluorescence emission spectra of (d) para-carbazole and (e) meta-carbazole.

To understand how deprotonation affecting the electron transport in the molecule, the theoretical calculation is employed by the combination of density functional theory (DFT) and non-equilibrium Green’s function (NEGF)47-49 (section 5 in SI). As shown in Figure 5a, the transmission of Para-N-e (green line) is lower than that of Para-N-H (purple line) across a wide energy range, which is consistent with the experiments. Meanwhile, the sharp formant between -1 eV and 0 eV, shown in the green curve in Figure 5a, is the HOMO formant. An obvious dip between the HOMO formant and the LUMO formant indicates that there is a distinct DQI effect between HOMO and LUMO.

To further reveal this phenomenon from the perspective of molecular orbital, the spatial distribution of orbital levels of Para-N-H and Para-N-e are analyzed as shown in Figure 5c. For Para-N-H, HOMO and LUMO orbitals are delocalized. In contrast, for Para-N-e formed by deprotonation, the introduction of negative charge causes the localized HOMO and the delocalized LUMO. The localized distribution of electron cloud on HOMO is only on the carbazole fragment, which leads to the weak coupling between HOMO and the electrode Fermi level to some extent. From the Molecular projection self-consistent Hamiltonian (MPSH) states of HOMO for Para-N-e, the fact that HOMO is still predominantly distributed on carbazole fragment again indicates a weak coupling between HOMO and the electrode Fermi levels (more details in Figure
Furthermore, the weak coupling leads to a decrease in the broadening of HOMO formant and a decrease in the resonance energy, resulting in a low and sharp HOMO formant as shown in Figure 5a. In addition, the localized HOMO and the delocalized LUMO even fulfill the theory of Fano resonance that a localized state interacts with a continuum. In contrast, as shown in Figure 5b, the transmission of Meta-N-e (green line) is slightly higher than that of Meta-N-H (purple line) across a wide energy range, which is consistent with the experiments. For Meta-N-e formed by deprotonation, the molecular orbitals are closer to the Fermi level than Meta-N-H, resulting in a slight increase in conductance. In addition, for Meta-N-H and Meta-N-e, the electron cloud on the HOMO orbitals is delocalized and not affected by deprotonation, as shown in Figure 5d. Although the LUMO orbital of Meta-N-H seems to be localized, the transmission spectrum of Meta-N-H shows that the energy of the LUMO formant is high, which means that the LUMO orbital of the molecule is delocalized, thus without the creation of the DQI effect.

Figure 5 | DFT calculated electrical properties of para-carbazole and meta-carbazole. Transmission curves of (a) Para-N-H (purple line) and Para-N-e (green line) and (b) Meta-N-H (purple line) and Meta-N-e (green line). Spatial distribution of orbital levels of (c) Para-N-H (LUMO: -1.14 eV, HOMO: -5.14 eV) and Para-N-e (LUMO: 1.53 eV, HOMO: -1.21 eV) and (d) Meta-N-H (LUMO: -0.80 eV, HOMO: -5.07 eV) and Meta-N-e (LUMO: 1.67 eV, HOMO: -1.31 eV).

To investigate the effect of the introduction of negative charge on the intramolecular electron transport pathway, we calculated the electron transport pathway of Para-N-H
and Para-N-e as shown in Figure 6a-6b and Table S2 in SI.\textsuperscript{50, 51} For Para-N-H, electrons are mainly transmitted through the C-C pathway (more details in Table S2, at the energy position of -1.4 eV, the ratio of current in the C-C pathway reached 71.36%), such a para-connection pathway does not have DQI effect,\textsuperscript{6} thus a higher conductance is revealed, which is also verified on the transmission curve (Figure 5a, purple line). However, for Para-N-e, deprotonation leads to the localization of HOMO orbitals, which further causes the ring current in carbazole center (Figure 6b). The current direction in carbazole center at different energy positions (-0.5 eV and -0.3 eV) is reversed, which is also a sign of DQI,\textsuperscript{52} and the reversed ring current blocks the electron transport pathways of C-C and C-N-C in the molecule, resulting in a significant change of electrical transport properties (more details in Table S2, at the energy of -0.5 eV and -0.3 eV, the current of C-C is capable of that of C-N-C while the direction is opposite). The previous flicker noise measurements also confirmed such changes in intramolecular electron transport pathways. For Para-N-H, which mainly transmits electrons through the C-C pathway, its electron transport mode is coupled by both through-bond and through-space, while for Para-N-e, due to the DQI effect, the through-bond tunneling is suppressed, thus the through-space tunneling becomes dominant, therefore the noise power changed from 1.5 in Para-N-H to 2.0 in Para-N-e.\textsuperscript{53} On the other hand, for meta-carbazole, the electron transport pathways do not significantly change before and after deprotonation, so the electron transport properties remain unchanged. The calculation of transmission pathways shows that electrons in both Meta-N-H and Meta-N-e are transmitted through the C-N-C pathway as shown in Figure 6c-6d (more details in Table S2, at the energy position of -1 eV, the ratio of current in the C-N-C pathway for Meta-N-H and Meta-N-e is 97.77% and 81.90%, respectively). The flicker noise analysis also shows that the noise power of Meta-N-H and Meta-N-e is $G^1$ before and after deprotonation, which means that their electrical transport mode remains through-bond tunneling (more details in Figure S19 from SI). The flicker noise remains unchanged before and after deprotonation for meta-carbazole, which to some extent proved that the change of flicker noise before and after
deprotonation for para-carbazole is due to the change of intramolecular electron transport pathway.

**Figure 6 | Transmission pathways of para-carbazole and meta-carbazole.** Transmission pathways of (a) Para-N-H at the energy of -1.4 eV (upper panel) and 0.5 eV (lower panel), (b) Para-N-e at the energy of -0.5 eV (upper panel) and -0.3 eV (lower panel), (c) Meta-N-H at the energy of -1 eV (upper panel) and 1 eV (lower panel) and (d) Meta-N-e at the energy of -1 eV (upper panel) and 0.4 eV (lower panel).

**Conclusion**

In conclusion, the charge transport through single-molecule junctions of para-carbazole and meta-carbazole before and after deprotonation was investigated using the STM-BJ technique. The results demonstrate that the injection of negative charge via the deprotonation of para-carbazole leads to a reversible fourfold suppression of conductance. The combined theoretical calculations and the flicker noise analysis revealed that the injection of negative charge results in the localization of HOMO orbitals in para-carbazole, leading to the reversed ring current and eventually blocking the intramolecular electrical transport pathways because of DQI. In comparison, the conductance and main electron transport pathway of meta-carbazole remain unchanged before and after deprotonation. Our findings provide an in-situ strategy for designing
reversible molecular switches to manipulate charge transport through the single-molecule junction, which is achieved via blocking the intramolecular transport pathways of electrons by localized electron distribution.

**Methods**

**Materials**

The molecules were synthesized according to the previous report (For more details, see supporting information (SI) section 1). For more details, see supporting information (SI) section 1. The gold substrates were prepared by coating 200 nm Au film on silicon wafers. Gold wire (99.99%, 0.25 mm diameter) was purchased from Beijing Jiaming Platinum Nonferrous Metal Co, Ltd. for the fabrication of the gold tip. The hafnium dioxide coated gold tip was prepared by Atomic Layer Deposition (ALD) (For more details, see SI section 4.2).

**The STM-BJ measurement**

The conductance of molecular junctions was measured using the lab-built scanning tunneling microscope break junction (STM-BJ) technique under ambient conditions (Figure S11). In the STM-BJ measurement, the distance between the gold tip and the substrate is controlled by a stepper motor and a piezo stack. The bias voltage is applied between the tip and substrate, and the current is used as feedback to control the movement of the gold tip. During the repeating opening (tip retracting) and closing (tip approaching) cycles, the conductance versus displacement traces are collected, and the traces of the opening cycles are used for further analysis.

**The flicker noise measurement.**

The flicker noise measurement carried out according to the previous studies. We suspended the tip during the retracting process for 200 ms after the formation of molecular junctions. The noise spectrums were obtained by the fast Fourier transform. We calculated the noise power according to the noise spectrum integration from 100 Hz to 1000 Hz and plotted the noise power against the average conductance.
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J. L., W. H. and Z. N. C. co-supervised the project. J. L., D. Z., P. D., Y. Z. and Q. C. Z wrote the manuscript with output from all authors. P. D., C. L. and K. Q. synthesized and provided the structural characterization of the molecules in Z. N. C.’s lab. D. Z., Y. W. and C. T. carried out the break junction experiments and analyzed the data, J. S. and W. H. built the electrical measurement instrument and wrote the software to control the break junction setup, Y. Z., P. D., L. C. and J. Y. W. performed the theoretical modeling in W. H.’s Lab.

**Competing interests**

The authors declare no competing interests.

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