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Nonequilibrium Bond Forces in Single-Molecule Junctions

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Abstract

Passing a current across two touching C_{60} molecules imposes a nonequilibrium population of bonding and antibonding molecular orbitals, which changes the equilibrium bond character and strength. A current-induced bond force therefore contributes to the total force at chemical-bond distances. The combination of first-principles calculations with scanning probe experiments exploring currents and forces in a wide C_{60} - C_{60} distance range consistently evidences the presence of current-induced attraction that occurs when the two molecules are on the verge of forming a chemical bond. The unique opportunity to arrange matter at the atomic scale with the atomic force and scanning tunneling microscope tip has enabled closely matching molecular junctions in theory and experiment. The findings consequently represent the first report of current-induced bond forces at the single-molecule level and further elucidate the intimate relation between charge transport and force. The results are relevant to molecular electronics and chemical reactions in the presence of a current.

Keywords: density functional theory, nonequilibrium Green's function, atomic force microscopy, scanning tunneling microscopy, single-molecule junction, C_{60}

The measurement of forces at the atomic scale with piconewton resolution using an atomic force microscope (AFM) has tremendously advanced physics research in the last decade. Seminal works discovered the spin texture of magnetic surfaces,^{1,2} explored forces to move single atoms,³ and unraveled the atomic skeleton of molecules.^{4,5} Conventionally, these forces are described as a superposition of long-range van der Waals and electrostatic attraction together with short-range Pauli repulsion, which have to be supplemented in cases of magnetic materials,¹ ionic crystals or polar molecules,^{6,7} and surfaces with low conductivity.⁸

Forces induced by current flow were previously discussed in the context of electromigration,⁹⁻¹² *i. e.*, the transport of matter through conductors in the presence of a dc current. More recently, the impact of a current across atomic wires¹³⁻¹⁵ and single molecules¹⁶⁻²⁴ was

scrutinized, with emphasis on theoretical modeling.^{13–17,19,21–24} The experimental works report local-heating effects of the current without imaging the individual molecule attached to the electrodes.¹⁸ While the rotational state of single-molecule electric motors was controlled by local injection of an electric current and imaged with a scanning tunneling microscope (STM), force measurements were not involved.²⁰

Experiments exploring the interplay between forces and currents at the single-atom or single-molecule level with precise characterization of the junction are very scarce. For atomic contacts it was shown by using an AFM and an STM that short-range forces and junction conductance exhibit an exponential behavior with characteristic decay lengths in the tunneling range²⁵ and are thus correlated. In another experimental work the influence of the tunneling current on the force was unraveled for samples with low conductivity.⁸ Both reports^{8,25} did not consider bond forces.

Here, we go beyond previous reports by combining state-of-the-art density functional theory (DFT) and nonequilibrium Green’s function (NEGF) calculations^{26–31} with AFM-STM experiments on artificially fabricated molecular junctions that closely match the model setup. The calculations unveil a current-induced attraction between the C₆₀ molecules that reflects the nonequilibrium population of bonding and antibonding molecular orbitals. The resulting nonequilibrium bond force is dominant for elevated junction conductance in the Pauli repulsion distance range and exhibits a nearly parabolic evolution with the bias voltage. Force spectroscopy in the noncontact frequency modulation mode of AFM with a C₆₀-terminated tip and C₆₀ molecules adsorbed on Cu(111) ranging from far tunneling to chemical-bond distances is in accordance with the theoretical predictions. After removing van der Waals, electrostatic and Pauli force contributions to the measured total force the residual force component exhibits all characteristics of the calculated current-induced force.

Figure 1a shows the setup used in the calculations. Top (*t*) and bottom (*b*) electrodes are represented by Cu(111) slabs each. Three Cu atoms arranged in a triangle at Cu(111) nearest-neighbor sites are attached to the top electrode. C₆₀ molecules are adsorbed to *t*

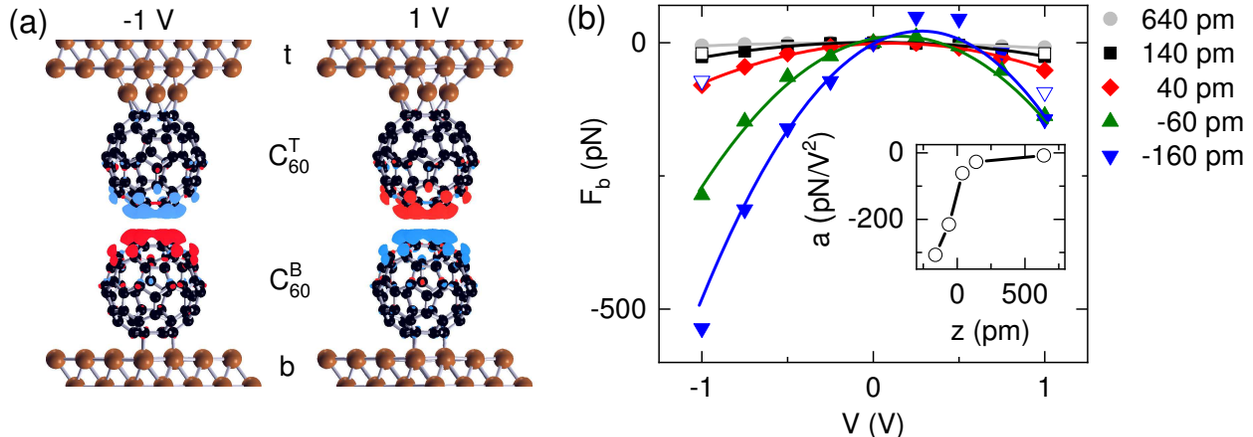


Figure 1: Junction geometry and calculated current-induced bond forces. (a) Cu(111) electrodes (top: t ; bottom: b) are modeled by a (4×4) unit cell with periodic boundaries in each electrode plane. Top C_{60} (C_{60}^T) is attached *via* a C hexagon to 3 Cu atoms arranged in a triangle at nearest-neighbor sites of Cu(111), while bottom C_{60} (C_{60}^B) resides with a C hexagon on Cu(111). The calculated accumulated charge density at bias voltage -1 V (left), 1 V (right) is indicated at each C_{60} . (b) Calculated bond force F_b as a function of the bias voltage V for the indicated unrelaxed C_{60} - C_{60} distances z . Open squares (triangles) depict the electrostatic attraction due to C_{60} charges (a) at a mutual separation of 140 pm (-160 pm). Inset: Curvature of parabolic fit to F_b as a function of z .

and b electrodes with C hexagons facing each other and are referred to as C_{60}^T and C_{60}^B , respectively (the Supporting Information, Figure S1, show results obtained for C_{60}^T exposing a C pentagon to the C hexagon of C_{60}^B). The chemical potentials μ_t and μ_b of, respectively, t and b electrodes (Figure 1a) differ in order to enable an electric current across the molecular junction. Due to different μ_t and μ_b the C_{60}^T and C_{60}^B molecules exhibit different charges (shaded areas in Figure 1a) that are accumulated at the C hexagons oriented towards the vacuum. These charges will be used below to estimate the electrostatic attraction between the molecules.

Using the combined DFT-NEGF formalism^{27,30} (Supporting Information) interatomic forces in the C_{60} - C_{60} junction were determined from the device Hamiltonian \mathbf{H} and the nonequilibrium electron density defined by the density matrix $\mathbf{D} = \int \mathbf{A}^t \cdot n_F(E - \mu_t) - \mathbf{A}^b \cdot n_F(E - \mu_b) dE$ ($\mathbf{A}^{t,b}$: spectral functions of t and b electrodes, n_F : Fermi-Dirac function)

according to²⁷

$$\mathbf{F}_n = -\text{Tr} \left(\frac{\partial \mathbf{H}}{\partial \mathbf{r}_n} \cdot \mathbf{D} \right) \quad (1)$$

with n indicating the atom at position \mathbf{r}_n . Bond forces are defined as the difference of the total forces on C_{60}^T and C_{60}^B projected onto the bond unit vector \mathbf{e} connecting the molecular centers of mass, *i. e.*,

$$F_b = \left(\sum_{n \in T} \mathbf{F}_n - \sum_{n \in B} \mathbf{F}_n \right) \cdot \mathbf{e}. \quad (2)$$

Figure 1b shows the bias voltage dependence of F_b for different C_{60} - C_{60} distances. Calculated data appear as symbols while solid lines represent fits of a second-order polynomial, $aV^2 + bV + c$, to the data and serve as a guide to the eye. Distance $z = 0$ pm is defined at the point of maximum attractive total force in both calculations and experiments (*vide infra*). This point signals the C_{60} - C_{60} chemical bond formation and is referred to as z_c in the following. Distances $z > z_c$ ($z < z_c$) correspond to tunneling (contact) ranges.

According to Figure 1b, F_b is attractive for essentially all investigated C_{60} - C_{60} distances and covered bias voltages. It exhibits a parabolalike evolution with the bias voltage. Figure 1b further reveals that a decreasing mutual molecule distance leads to various effects. First, in the tunneling distance range, $z > 140$ pm, F_b is comparable to the electrostatic force, F_{el} (open symbols in Figure 1b), which results from the accumulated charge at C_{60}^T and C_{60}^B (shaded areas in Figure 1a). The upper estimate of F_{el} shown in Figure 1b was obtained using a plate capacitor model in which the calculated charge densities were attached to discs defined by the planes of the outermost C hexagons of the two C_{60} . Importantly, entering further into the range of chemical-bond distances, $z < 140$ pm, F_b outweighs F_{el} appreciably for finite V . Second, with decreasing C_{60} - C_{60} separation the parabola apex shifts toward positive V giving rise to an asymmetric behavior of F_b . This asymmetry is due to the asymmetric junction setup, *i. e.*, to C_{60}^T adsorbed on a triangular Cu cluster and C_{60}^B adsorbed on planar Cu(111). Third, the parabolic curvature, a , decreases with decreasing C_{60} - C_{60} distance (inset to Figure 1b). A particularly rapid decrease is observed upon entering into distance

ranges in the vicinity of chemical-bond formation, $z < 140$ pm.

After describing the properties of the calculated bond force F_b , its nature will now be elucidated in terms of electron transport across a molecular junction *via* antibonding and bonding molecular orbitals.^{32,33} The electric current induces a nonequilibrium population of these orbitals and, thus, alters the bond character and strength. The antibonding or bonding character of the bond-forming states is identified by the crystal orbital overlap population (COOP).³⁴ It is calculated as the energy-resolved charge density in the intermolecular bond,

$$\text{COOP}(E) = 2 \sum_{i \in T, j \in B} S_{ij} A_{ij}(E) \quad (3)$$

(S : orbital overlap, A : spectral function in atomic orbital basis), where i and j indicate all C_{60}^T and C_{60}^B orbitals, respectively. The spectral functions may furthermore be partitioned to consider contributions of t and b electrodes ($\alpha = t, b$) to the origin of scattering states, resulting in decomposed $\text{COOP}^\alpha(E) = 2 \sum_{i \in T, j \in B} S_{ij} A_{ij}^\alpha(E)$. The COOP sign determines whether the states at energy E contributing to the bond have bonding (positive) or antibonding (negative) character. The integration of the COOP^α weighted by n_F ,

$$O_{ij}^\alpha = S_{ij} \int A_{ij}^\alpha(E) n_F(E - \mu_\alpha) dE, \quad (4)$$

yields the overlap population or bond charge, $\text{OP} = \sum_{\alpha=t,b} \sum_{i \in T, j \in B} O_{ij}^\alpha$, which provides a measure of the C_{60} - C_{60} bond charge and strength.³⁵ The V -dependent change of OP with respect to $V = 0$ V reads

$$\Delta \text{OP} = \text{OP}(V) - \text{OP}(0). \quad (5)$$

Figures 2a,b illustrate the concept of COOP for a C_{60} - C_{60} junction at $(\mu_t - \mu_b)/e = -1$ V (e : elementary charge). For positive bias voltage t and b electrodes interchange roles. Figure 2a shows the relevant C_{60} orbitals involved in the overlap, while Figure 2b discloses their bonding or antibonding character, indicated by the shaded areas for, respectively, $E > 0$ eV

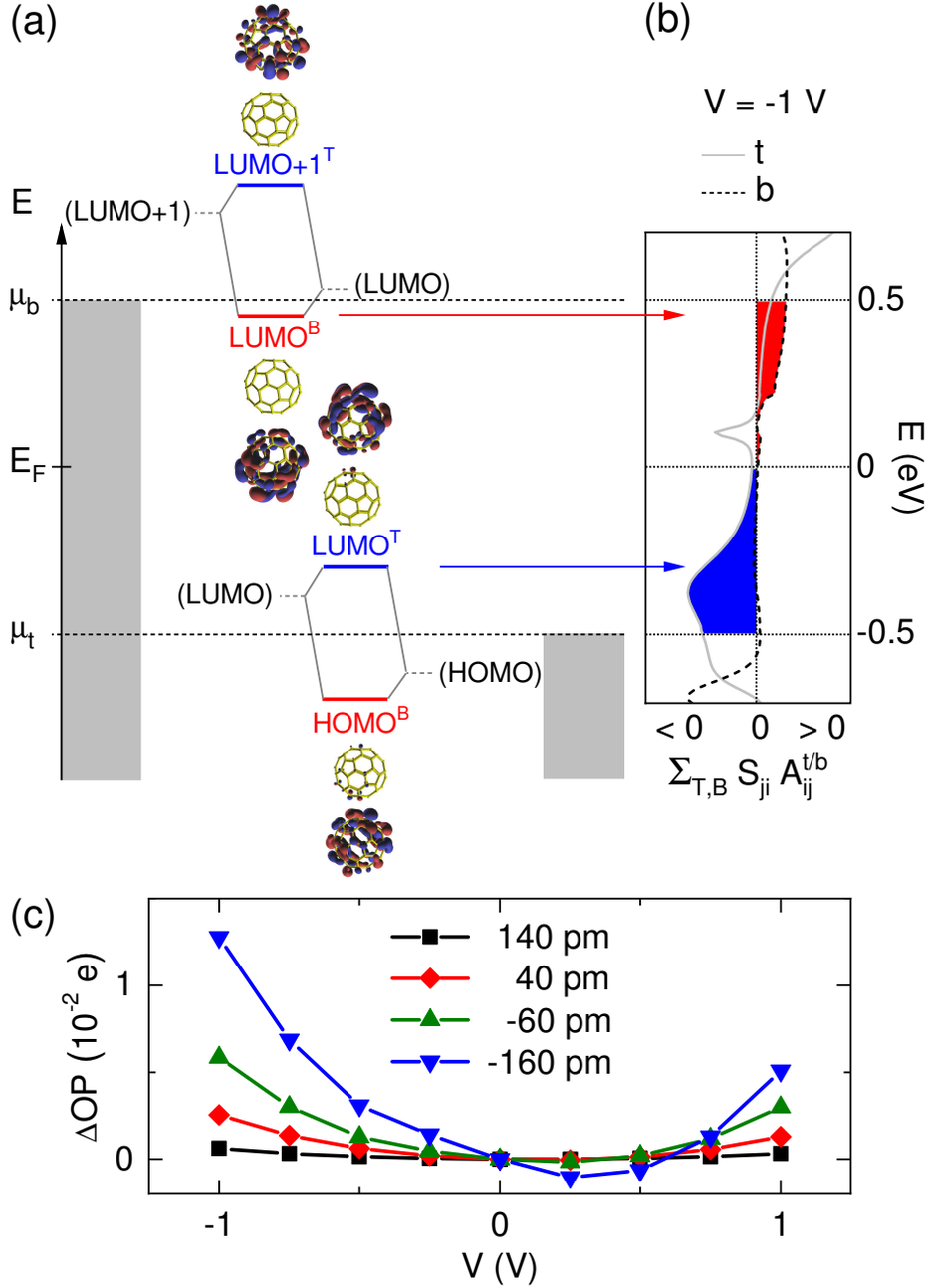


Figure 2: Concept of the crystal orbital overlap population (COOP). (a) Energy levels of the coupled C_{60} - C_{60} system ($HOMO^B$, $LUMO^T$, $LUMO^B$, $LUMO+1^T$) and the schematic relation to the uncoupled counterparts ($HOMO$, $LUMO$, $LUMO+1$ in parentheses) in a range spanned by the chemical potentials μ_t and μ_b of t and b electrodes for $V = (\mu_t - \mu_b)/e = -1$ V. E_F is the equilibrium Fermi energy. Antibonding orbital of C_{60}^T , $LUMO^T$, and bonding orbital of C_{60}^B , $LUMO^B$, fall into the voltage window. (b) Calculated electrode-decomposed COOP (eq 3) of $LUMO^T$ and $LUMO^B$ at $z = -60$ pm revealing the bonding (antibonding) character indicated by red (blue) color. $E = 0$ eV corresponds to E_F . The lowering of μ_t depletes the antibonding $LUMO^T$ (blue area), while the increase of μ_b populates the bonding $LUMO^B$ (red area), both resulting in an increase of ΔOP . (c) ΔOP (eq 5) as a function of V for indicated z .

and $E < 0$ eV. Molecular orbitals projected onto the coupled C_{60} molecules³⁶ with energies in the interval $\mu_t \leq E_F \leq \mu_b$ (E_F : equilibrium Fermi energy) are labeled $LUMO^T$ and $LUMO^B$ because of their resemblance with the lowest unoccupied molecular orbital (LUMO) of isolated C_{60} . The resulting electrode-resolved $COOP^\alpha(E)$ is plotted for t and b electrodes in Figure 2b as solid and dashed lines, respectively. The highest occupied molecular orbital (HOMO) as well as the LUMO+1 do not participate in the COOP for the considered energies. The scenario depicted in Figures 2a,b is conceptionally close to the simplified model presented previously.³²

Figure 2c shows ΔOP (eq 5) as a function of V . For small C_{60} - C_{60} distances, ΔOP mirrors the nearly quadratic behavior and characteristic asymmetry in the bias voltage polarity as observed from F_b (Figure 1b). At large C_{60} - C_{60} distances, however, ΔOP is no longer related to F_b reflecting the dominant influence of F_{el} in that distance range.

Given the akin V dependence of ΔOP (Figure 2c) and F_b (Figure 1b) at the different C_{60} - C_{60} distances, a close correlation between ΔOP and F_b is established on the basis of which we attribute the origin of F_b to the bond charge. The V -dependent changes in F_b are due to the difference in the chemical potentials, *i. e.*, to the nonequilibrium electron distribution and, thus, to the current. For, *e. g.*, $V = -1$ V (Figure 2b), $\mu_b > \mu_t$ leads to an increased population of the bonding $LUMO^B$ (red area) and a concomitantly decreased population of the antibonding $LUMO^T$ (blue area) with respect to the equilibrium situation at 0 V, which both increase ΔOP and give rise to a current-induced C_{60} - C_{60} attraction. Moreover, the junction electric field affects the level alignment^{37,38} and effectively increases the electronic coupling and, thus, enhances the discussed effect with increasing V .

Combined AFM and STM experiments (Supporting Information) were carried out to test the theoretical results. Using a surface science approach, junctions were assembled *in situ* to mimic the setup in the model calculations as closely as possible. To this end, C_{60} was adsorbed on Cu(111) (Figure 3a). Ordered molecular islands with monolayer (ML) height were obtained after annealing the C_{60} -covered sample. Separate C_{60} molecules were detached

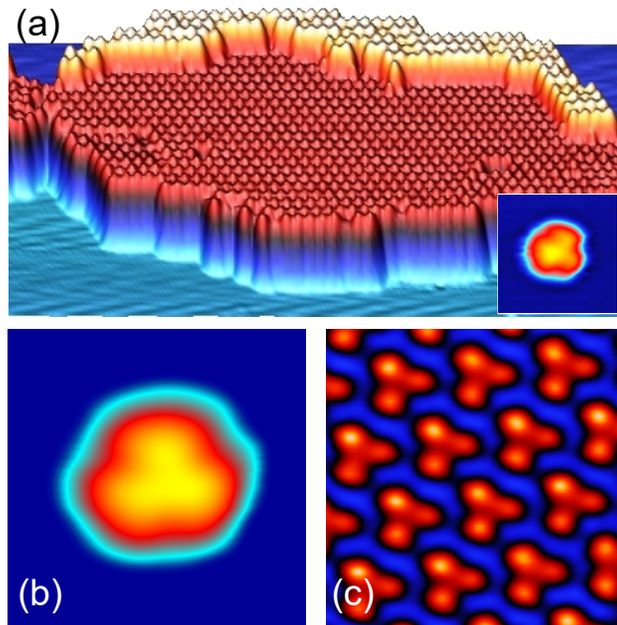


Figure 3: STM topographic data of C_{60} -covered Cu(111) and of a C_{60} -terminated tip. (a) Constant-current STM image of a C_{60} island on Cu(111) (0.7 V, 100 pA, $50 \times 25 \text{ nm}^2$). The color scale ranges from 0 pm (blue) to 830 pm (white). Inset: STM image of a C_{60} -terminated tip using an atomic protrusion on Cu(111) (-2.1 V , 50 pA, $2.2 \times 2.2 \text{ nm}^2$). The color scale ranges from 0 pm (dark blue) to 100 pm (bright yellow). (b) Close-up view of a separated C_{60} (1.7 V, 50 pA, $4 \times 4 \text{ nm}^2$). (c) Close-up view of the inner part of a C_{60} island (2 V, 50 pA, $3 \times 3 \text{ nm}^2$). For (b), (c) the same color scale as in (a) is used. STM data were processed using WSXM.³⁹

from the step edges of ML molecular islands and dragged across the substrate terrace with the tip of the microscope. Close-up views of separate C_{60} molecules (Figure 3b) and ML islands (Figure 3c) reveal a trefoil-like structural motif of adsorbed C_{60} , which indicates a C hexagon exposed to vacuum and to the surface, in agreement with a previous report.⁴⁰ The STM data of Figure 3 were acquired with a PtIr tip whose apex had presumably been coated with Cu substrate material due to the applied tip preparation protocol (Supporting Information). For the force measurements to be discussed next, the tip was terminated with a single C_{60} molecule using a standard procedure.^{41,42} The successful transfer to the tip was verified by imaging the modified tip apex with an adsorbed Cu atom that had been transferred from the Cu-coated tip to Cu(111)⁴³⁻⁴⁶ prior to C_{60} termination of the tip apex. The inset to Figure 3a shows an STM image revealing that C_{60} at the tip is likewise oriented with a C hexagon to the vacuum.

Figure 4a presents the changes in the tuning fork resonance frequency Δf during the approach of a C_{60} -terminated tip to a single separated C_{60} , *i. e.*, for decreasing z . The same plot contains the vertical force, F_z , extracted from Δf according to previously reported algorithms,^{47,48} which led to virtually identical results. The evolution of the simultaneously recorded current, I , is shown in Figure 4b.

$F_z(z)$ (Figure 4a) adopts the expected Lennard-Jones-type^{49,50} behavior. At large z , $F_z(z)$ is dominated by the long-range and slowly varying van der Waals force and electrostatic attraction between the electrodes. The minimum of F_z at $z_c^{(F)}$ signals the point of maximum attraction and defines the distance range for chemical-bond formation. For $z < z_c^{(F)}$ the short-range Pauli repulsion progressively strengthens and, thereby, weakens the C_{60} - C_{60} attraction. Figure 4b shows the current evolution with z around $z = 0$ pm. $I(z)$ increases exponentially with decreasing C_{60} - C_{60} separation, more strongly for $z > z_c^{(I)}$ than for $z < z_c^{(I)}$. The intersection of the extrapolated exponential fits to the data (dashed lines in Figure 4b) approximates the contact point, $z_c^{(I)}$.^{43,45,51,52} Both data sets, $F_z(z)$ and $I(z)$, provide a consistent definition of the contact point, *i. e.*, $z_c \equiv z_c^{(F)} \approx z_c^{(I)}$. Similar behavior was

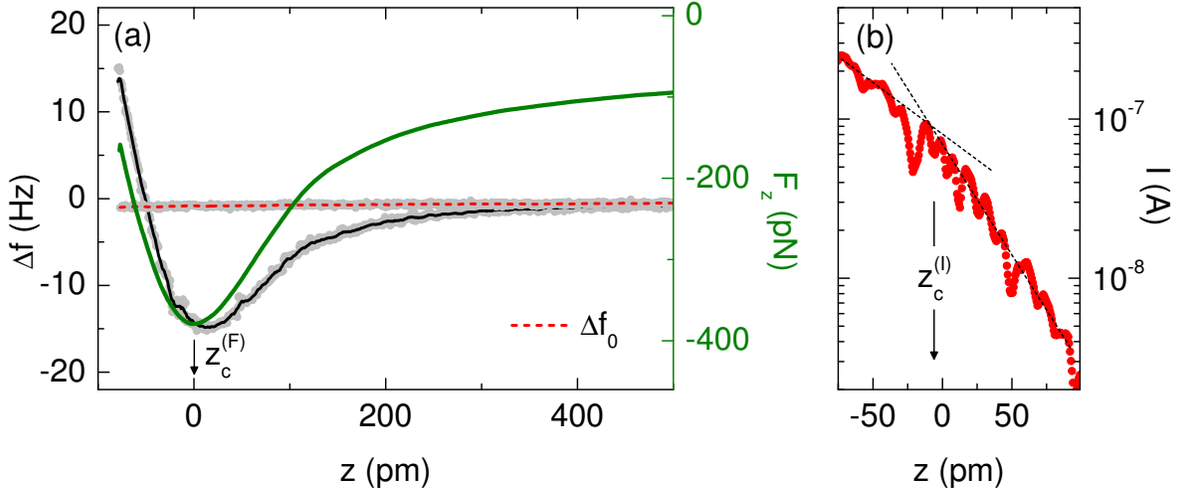


Figure 4: Force and current data obtained from C_{60} - C_{60} junctions. (a) Resonance frequency change Δf (gray dots and superimposed smoothed data as a black line) and vertical force F_z (green line) as a function of z measured at 0.6 V for a C_{60} -terminated tip and a single C_{60} on Cu(111). The minimum of F_z at $z_c^{(F)}$ signals C_{60} - C_{60} contact formation. Δf_0 data (gray dots and superimposed smoothed data as a dashed line) were obtained with the same C_{60} -terminated tip on clean Cu(111). An additional z offset was applied to guarantee the same absolute tip-surface distance as for $\Delta f(z)$ measurements atop adsorbed C_{60} . (b) Current I (dots) across the C_{60} - C_{60} junction of (a) acquired simultaneously with Δf . Dashed lines are exponential fits to $I(z)$ data for $z < z_c^{(I)}$ and $z > z_c^{(I)}$ whose intersection indicate the point of contact at $z_c^{(I)}$. In (a), (b) the feedback loop had been disabled at 1 V, 50 pA and the bias voltage was subsequently ramped to 0.6 V.

observed from C_{60} molecules embedded in ML molecular islands (not shown).

In order to access the predicted bond forces F_b (Figure 1b) long-range, slowly varying attractive forces between the full tip structure and surface must be removed from the experimentally obtained total force F_z (Figure 4a). The following approximation has to be made to this end. The measured total force acting on the tip, F_z , can be decomposed into a short-ranged force between the two C_{60} molecules, F_z^* , and a residual long-ranged and slowly varying force, F_0 , acting between the tip including the attached C_{60} and the metal substrate. One may expect F_z^* and its bias voltage dependence to be negligible at large distance, while at chemical-bond distances only F_z^* will reflect the charge redistributions occurring at the C_{60} molecules in the course of the current-induced population of bonding and antibonding orbitals. In contrast, F_0 takes into account charges well separated from the C_{60} - C_{60} junction and is independent of the presence of C_{60} adsorbed on Cu(111).

To obtain F_z^* and test this approximation, we follow a previously suggested and successfully applied procedure^{25,53,54} where the resonance frequency changes were acquired with a C_{60} -terminated tip on an extended pristine Cu(111) surface region ($\approx 200 \times 200 \text{ nm}^2$) at the same absolute tip-surface distance as used for the $\Delta f(z)$ measurements atop adsorbed C_{60} on Cu(111). These frequency changes are referred to as Δf_0 (dashed line in Figure 4a) and reflect the long-ranged slowly varying force F_0 (*vide supra*). The subtraction of the long-range and slowly varying F_0 from the total force F_z is performed by transforming $\Delta f - \Delta f_0$ into a force,^{47,48} F_z^* . This approach was used for correcting Δf and F_z obtained for separated C_{60} as well as for molecules embedded in an ML island. In Figure 5 F_z^* is plotted as a function of V and different C_{60} - C_{60} distances for the single (Figure 5a) and ML (Figure 5b) C_{60} . The vertical offset is due to the distance-dependent short-range force.

In all cases F_z^* is essentially independent of V for large mutual C_{60} distances (top), which reflects the appropriate subtraction of long-range and slowly varying forces, as expected. With decreasing C_{60} - C_{60} distance F_z^* exhibits a progressively pronounced parabolalike behavior (middle and bottom). In order to facilitate the comparison with the calculated results

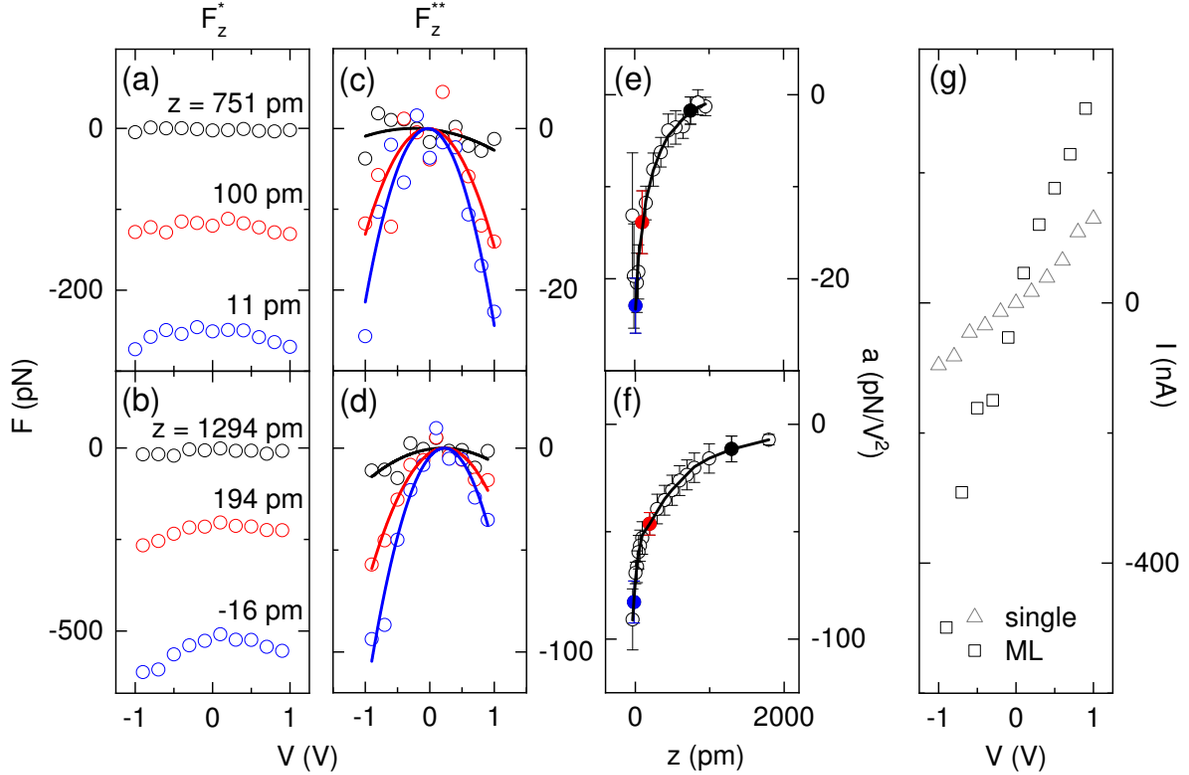


Figure 5: Experimental evidence for current-induced forces. (a), (b) Residual force F_z^* for (a) single and (b) ML C_{60} obtained for the indicated distances z after subtracting long-range forces from F_z (see text). (c), (d) Plots of $F_z^{**}(V) = F_z^*(V) - F_z^*(0)$ for (c) single and (d) ML C_{60} . Solid lines depict parabolic fits to F_z^{**} . (e), (f) Parabola curvatures of (c), (d) for different z . Dots correspond to the data presented in (a)–(d). (g) I - V characteristics for single (triangles) and ML (squares) C_{60} acquired at $z = 0$ pm.

(Figure 1b) F_z^* is referred to its value at the parabola maximum close to 0 V, resulting in $F_z^{**}(V) = F_z^*(V) - F_z^*(V \approx 0 \text{ V})$ plotted for single and ML C₆₀ in Figure 5c and Figure 5d, respectively. The solid lines represent parabolic fits to the data with the curvature plotted as a function of z in Figure 5e,f.

How do the experimental data compare with the theoretical findings? The experimentally extracted F_z^{**} is in qualitative agreement with the calculated F_b . Both F_z^{**} and F_b are attractive and exhibit a nearly parabolic evolution with V (Figure 1b, Figure 5c,d). Their curvature is negative, weak in the far tunneling range and increasingly pronounced for smaller C₆₀-C₆₀ distances. Moreover, the curvature rapidly decreases upon entering into the range of chemical-bond distances in both cases (inset to Figure 1b, Figure 5e,f). Quantitatively, however, there are some deviations between experiment and theory. The calculated properties generally tend to overestimate the experimental results since only the C₆₀-C₆₀ bond force is accounted for. The additional compliance due to relaxations of tip and surface structures is not considered. In addition, periodic boundary conditions are used in the simulations that inevitably lead to multiple tips in the model setup, which contribute to the overestimation of currents and forces (Supporting Information, Figure S2). The junction conductance in the calculations is approximately two times larger than the experimentally observed conductance, which entails a correspondingly larger bond force in the model than in the experiments. For instance, in the vicinity of chemical-bond distances, $z \approx z_c$, F_z^{**} varies between 0 pN and ≈ -25 pN for single C₆₀ (Figure 5c) and between 0 pN and ≈ -100 pN for ML C₆₀ in the voltage interval $-1 \text{ V} \leq V \leq 1 \text{ V}$. The corresponding variation of F_b covers the range from 0 pN to ≈ -100 pN and thus is a factor ≈ 4 larger than observed for single C₆₀, while it matches the range of F_z^{**} for ML C₆₀ very well. In addition, the evolution of the $F_b(V)$ curvature is more pronounced in the calculations than for the measured $F_z^{**}(V)$. The abrupt decrease of the calculated curvature upon entering chemical-bond distances is smoothed in the experiments due to, supposedly, a less stiff junction in which relaxations of tip-C₆₀ and C₆₀-surface geometries are involved. Because of the overall qualitative and

in part quantitative accordance between F_z^{**} and F_b ample evidence is provided by the experiment for the presence of the calculated current-induced forces.

The importance of the current is likewise evidenced by the comparison of I - V characteristics (Figure 5g) with F_z^{**} . In the relevant range of bias voltages, $-1 \text{ V} \leq V \leq 1 \text{ V}$, the current across single C_{60} (triangles) is a factor ≈ 4 lower than across C_{60} embedded in ML islands (squares), which matches well the ratio of F_z^{**} data obtained for these junctions (Figure 5c,d). The difference in the I - V characteristics is most likely due to the stronger C_{60} - $\text{Cu}(111)$ hybridization in the molecular island where the substrate surface exhibits a C_{60} -induced reconstruction.⁵⁵ As a consequence, the C_{60} LUMO is located at the Fermi energy⁵⁵ (Supporting Information, Figure S3), which entails a higher junction conductance than observed for single C_{60} . Furthermore, I - V data for ML C_{60} exhibit a more pronounced asymmetry with the polarity of the bias voltage. This behavior is likewise present for F_z^{**} . In the calculations the F_b asymmetry with the bias voltage polarity was assigned to the asymmetric coupling of C_{60} to the tip and to the substrate. However, the degree of asymmetry in the molecule-electrode coupling may considerably be reduced in the experiments. The metal tip may be less sharp than assumed in the calculations. In this case it would therefore represent a rather flat adsorption region for C_{60} , which is similar to the adsorption situation for C_{60} on the counter electrode, $\text{Cu}(111)$.

Before concluding it is noteworthy that deviations from C_{60} hexagon orientations at the tip apex did not change the characteristics of the bond force significantly. In the Supporting Information (Figure S4) F_z^* and F_z^{**} are presented for a tip terminated by a C_{60} molecule exposing a C pentagon to the surface. The weak dependence of F_z^* , F_z^{**} on the C_{60} orientation is in agreement with simulations (Supporting Information, Figure S1). The akin behavior may tentatively be attributed to the spatial distribution of the C_{60} LUMO, which is most important for contact formation (Figure 2a). It exhibits a toruslike shape centered at the C pentagon.^{45,51} The C hexagon orientation of C_{60} still shows good parts of the LUMO of the adjacent three C pentagons (Figure 3b,c). Therefore, the LUMO overlap may be

comparable in case of the different C_{60} orientations considered here and result in the similar behavior of the bond force. Another rationale may be a favored contact geometry; that is, in the Pauli repulsion distance range the reaction partners adopt a preferred orientation for bond formation that is independent of the initial orientation prior to contact. Forthcoming experiments will involve C_{60} on Cu(100) where several orientations of adsorbed C_{60} are available.⁵¹

In conclusion, theory and experiment unambiguously unveil current-induced bond forces in a molecular contact and contribute to the understanding of the relation between force and current in nanometer-scaled junctions. Owing to the assembly of the junction with precise control of the electrode composition and structure, model calculations and force spectroscopy experiments are based on closely matching contact geometries and, thus, enable a meaningful comparison of calculated and measured data. The current-induced bond forces in a biased conductive single-molecule junction result from the nonequilibrium population of bonding and antibonding molecular orbitals. These forces exhibit a nearly parabolic bias voltage behavior and are dominant for molecule–molecule separations in the chemical-bond distance range. Their magnitude depends on the orbital overlap in the energy range defined by the electrode chemical potentials. The reported theoretical and experimental findings are relevant to miniaturized circuits in molecular electronics and offer a fresh view on the formation of chemical bonds at the single-molecule level in the presence of a current.

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Supporting Information Available

The Supporting Information is available free of charge on the ACS Publications website at DOI: [hyperlink DOI]

Details on DFT modeling, NEGF calculation, data for different C₆₀ orientations, sample and tip preparation, spectroscopy of d/dV

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Graphical TOC Entry

