Evidence for temporary and local transition of sp² graphite-type to sp³ diamond-type bonding induced by the tip of an atomic force microscope

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Evidence for temporary and local transition of sp\(^2\) graphite-type to sp\(^3\) diamond-type bonding induced by the tip of an atomic force microscope

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Abstract
Artificial diamond is created by exposing graphite to pressures on the order of 10 GPa and temperatures of about 2000 K. Here, we provide evidence that the pressure exerted by the tip of an atomic force microscope onto graphene over the carbon buffer layer of silicon carbide can lead to a temporary transition of graphite to diamond on the atomic scale. We perform atomic force microscopy with CO terminated tips and copper oxide (CuOx) tips to image graphene and to induce the structural transition. A local transition induced by the force of the tip is accompanied by local rehybridization from an sp\(^2\)-bonded to an sp\(^3\)-bonded local structure. Density functional theory predicts that a repulsive threshold of \(\approx 13\) nN, followed by a force reduction by \(\approx 4\) nN is overcome when inducing the graphite-diamond transition. The experimental observation of the third harmonic with a magnitude of about 200 fm fits well to overcoming a force barrier of \(F_{\text{barrier}} \approx 5\) nN, followed by a force reduction by \(-F_{\text{barrier}}\) and an upswing by \(F_{\text{barrier}}\) for decreasing distances. Experimental evidence for this transition is provided by the emergence of third harmonics in the cantilever oscillation when the laterally flexible CO terminated tip exerts a large repulsive force. Probing the sample with rigid CuOx tips in the strong repulsive regime shows a strong difference in the yielding of the A versus B sites to the pressure of the tip. The large repulsive overall force of \(\approx 10\) nN is only compatible with the experimental data if one assumes that the repulsive force acting on the tip when inducing the transition is compensated by a heavily increased van-der-Waals attraction of the tip due to form fitting of tip and sample by local indentation. The experiment also shows that atomic force microscopy allows to perform high pressure physics on the atomic scale.

Contents
1. Diamond versus graphite and transitions ............................................................ 2
2. Density functional theory calculations on the effect of a high load of an atomic contact to bilayer graphene on SiC ......................................................... 3
3. Experimental setup ......................................................................................... 6
   3.1. Scanning probe microscopy ..................................................................... 6
   3.2. Experimental observables ....................................................................... 7
   3.3. Sample preparation ............................................................................... 8
4. Experimental data .......................................................................................... 10
   4.1. AFM with inert trimer tips and CO terminated tips ..................................... 11
1. Diamond versus graphite and transitions

'Deep within the earth diamonds grow. Diamonds the size of footballs, diamonds the size of watermelons—countless billions of tons of diamonds wait for eternity a 100 miles beyond our reach.'—this is the opening sentence in chapter I of Robert M Hazen’s stunning account on the history of making artificial diamond [1] as reported in [2, 3]. The pressure in the mantle of the earth at a depth of 100 miles is about 10 GPa, and fractions of these football-sized diamonds can survive the trip to the earth’s surface only when transported by volcanic eruptions of great speed [1]. Most of these ‘watermelon-sized diamonds’ emerge on the surface of the earth as graphite, the stable phase of carbon for pressures below about 2 GPa at low temperatures. A schematic phase diagram that shows the large coexistence area shaded in gray—both graphite and diamond are stable in ambient conditions, is presented in figure 1 after data from [4].

Graphite and diamond coexist over a large range of pressures and temperatures, indicated by the gray zone in the phase diagram of figure 1 created with data from [1, 4]. Therefore, transforming graphite to synthetic diamond requires to push the material through this wide gray area utilizing massive and ingenious reactor cells that sustain a pressure on the order of 10 GPa at temperatures around 2000 K [1]. At ambient conditions, graphite is slightly more stable than diamond by about 3 kJ mol$^{-1}$ or 30 meV per atom [5, 6], while at 0 K, diamond appears to be more stable than graphite by an even smaller energy [5]. The two phases separated by a barrier of about 0.5 eV per atom (figure 4(a) in [6]), and it was already found in 1962 [7] that diamond starts to transform into graphite both inside a crystal as well as on the surface by heating it in a vacuum to 1700 °C. This temperature corresponds to $k_B \cdot T \approx 172$ meV. While this energy is significantly below the barrier of 0.5 eV, the attempt frequency is expected to be around the frequency of optical phonons in diamond of about 40 THz, yielding a significant transition rate even for thermal energies below the barrier.

Recently, the tip of an atomic force microscope (AFM) [8] was used to induce a transition of bilayer graphene to diamond [9] or 2D boron nitride to a diamond-like structure by AFM [10]. These nanoindentation experiments applied loads of up to 100 nN to a blunt spherical Si probe with, compared to atomic dimensions, a large contact area, which lead to these sp$^2$ type bonding to diamond-like sp$^3$ type bonding conversions [9]. Recently, such transitions have also been observed in boron nitrate on a SiO$_2$ substrate [10].

Because of the increasing relative stability of matter with decreasing size, one might even expect that it is easier to induce this transition with an atomically sharp tip. The strength of materials with a given cross section is typically much smaller than the strength of an individual bond times the number of bonds that bridge the cross section. The reason is the slipping of bonds one by one under the influence of shear forces (see e.g. vol II, chapter 30-7 in [11]).

The tip of an AFM can create enormous pressure when its repulsive force is confined on a single unit cell. A surface unit cell in graphite spans an area of 246 pm $\times$ 246 pm $\times$ sin($\pi/3$) and forms a honeycomb mesh with two basis atoms spaced at 142 pm. In AFM, forces between the front atom of the tip and the sample...
atom next to it can reach forces in the nanonewton regime without creating damage. A small repulsive force of merely 525 pN over one unit cell already invokes a huge pressure of 10 GPa.

A transition of the bonding type between a CO terminated tip and a Fe adatom on Cu(111) has recently been observed [12]. In this experiment, approaching the CO terminated tip to the Fe adatom resulted first in a local van-der-Waals (vdW) minimum of about $-10$ pN at a distance of 400 pm, followed by a repulsive barrier of about $+20$ pN at 300 pm ending in a weak covalent bond with an absolute minimum of about $-400$ pN at a distance of 260 pm. Here, we attempt to modify the bonding character between the graphene surface layer and the buffer layer (BL) underneath on the scale of a few atoms with an atomically sharp tip.

2. Density functional theory calculations on the effect of a high load of an atomic contact to bilayer graphene on SiC

The creation of diamond in pressure cells differs largely from exerting pressure by a single front atom of an AFM tip. Density functional theory (DFT) can simulate the effect of impinging an atomically sharp CO tip on bilayer graphene on SiC. For simplicity, we only consider the CO molecule pointing with the O atom towards the sample as the AFM tip. DFT within the generalized gradient approximation of Perdew et al [13], as implemented in the Fritz-Haber-Institute-aims code package [15] is used here to calculate the effect of a CO terminated tip that is located in the center of a graphene hexagon on SiC(0001). We build on a previous DFT study of this sample system by Nemec et al [14]. The calculations were performed at zero bias voltage and a temperature of 0 K.

Figure 2 shows the results of a DFT calculation where a CO terminated tip is placed above the center of a graphene hexagon and the repulsive force on the CO tip termination is plotted against the interlayer distance between the monolayer graphene and the carbon BL for three different sizes of unit cells.

The black curve corresponds to a $1 \times 1$ structure where a CO terminated tip is placed over each unit cell as indicated in figure 2(b). The force starts at zero at the equilibrium interlayer distance of 335 pm and then rises sharply to a value of about 2 nN at a nearest inter-graphene-layer distance of 235 pm. The next stable data point is at about 160 pm, close to the bond length in diamond, where the force rises sharply again for even smaller distances.

The blue curve refers to a $2 \times 2$ structure, corresponding to one CO tip in four unit cells as depicted in figure 2(c). The force again starts at zero at the equilibrium interlayer distance of 335 pm and then rises even more sharply to a value of about 7 nN where it declines until the nearest inter-graphene-layer distance reaches 160 pm, again increasing profoundly upon further distance reduction.

Finally, the red curve refers to a $2\sqrt{3} \times 2\sqrt{3}$ structure, corresponding to one CO tip in 12 unit cells. Here, the force even rises up to a value of about 13 nN before it declines again until a distance of 150 pm. The calculated curve does not cover the even smaller distances that are expected to show another rising repulsive force.
Figure 2. A CO molecule is located over the hollow cite (center) of a hexagonal unit cell of bilayer graphene on SiC. (a) DFT calculated force spectra that occur during rehybridization for monolayer graphene on a graphene BL on SiC as a function of interlayer distance \( d_{g-g} \) as illustrated in (g). Three different sizes of unit cells are shown: (b) in the black curve, a CO molecule presses on every unit cell (1 × 1 structure), (c) the blue curve refers to one CO molecule for four unit cells (2 × 2 structure), and (d) finally one CO molecule for 12 unit cells (2√3 × 2√3 structure). (e) The slab is shown under high load where the transition has taken place. (f) Same slab with CO 20 pm further away. (g) Close up of the tip region of (e): the CO tip molecule is centered over the carbon hexagon, and one of the three A sites of the graphene surface layer moves significantly towards the BL and forms a covalent bond with a \( A_{BL} \) atom.

It should be noted, however, to exert such a high force (pressure) from the CO tip, in our simulations, the CO molecule has to be fixed vertically in the simulation. Without doing so, the CO molecule will be tilted away from the upright position before reaching such a high force. How the CO tip might be stabilized under the experimental condition will be discussed further below.

While the increase of unit cell sizes has led to increased force thresholds that induce the graphite to diamond transition, we expect that the force threshold saturates for large enough unit cells. Experimental and theoretical evidence shows that when graphene is exposed to forces from the tip of a scanning probe microscope close by, the largest part of the bending occurs within nanometers from the tip [16].

Table 1 summarizes the results of figure 2 and lists the corresponding average pressures. As expected, the average pressure falls for larger unit cells. However, the calculated pressure where a transition occurs for the 1 × 1 structure at \( T = 0 \) K is at 41 GPa surprisingly high, given that the empirically found pressure needed for artificial diamond creation is about 10 GPa.

A possible reason for this discrepancy is provided by Grochala, who notes that thermal activation is important in particular when transforming graphite to diamond, ‘which results in immeasurably sluggish transformation at \( T < 1000 \) K even when the pressures involved are larger than necessary for transformation’ (p 3682 in [5]).
Table 1. Rehybridization thresholds as a function of unit cell size.

| Number of unit cells | Surface vectors | $F_{\text{rehybridization}}$ (eV Å$^{-1}$) | $F_{\text{rehybridization}}$ (nN) | Pressure (GPa) |
|----------------------|----------------|--------------------------------------|-------------------------------|----------------|
| 1                    | $1 \times 1$  | 1.33                                 | 2                             | 41             |
| 4                    | $2 \times 2$  | 4.7                                  | 7                             | 36             |
| 12                   | $2\sqrt{3} \times 2\sqrt{3}$ | 8.0                                  | 13                            | 20             |

Figure 3. (a) Bonding energies of graphene and diamond after figure 4(a) of [6], showing the energetic minima at the bond lengths of graphene at 142 pm and diamond at 155 pm. (b) Energy profile of (a) rescaled to the vertical bond length of bilayer graphene of about 330 pm. (c) Force in vertical direction derived from (b). Note that (b) and (c) are only qualitative derivations from graphene and diamond data.

Although the sample is at a temperature of about 4 K in our experiment, it is conceivable that the oscillating motion of our tip with its amplitude of $A = 50$ pm shakes the lattice in a similar fashion as thermal excitation and thus helps the atoms to find their new equilibrium positions as the thermal motion at the high temperatures required in artificial diamond creation. One may argue that the oscillation of the cantilever is very slow in frequency (40 kHz) with respect to optical phonon frequencies (40 THz) and should therefore considered to be adiabatic. However, as we will see below, dissipation is observed in the experiment, proving that the oscillation of the tip to and from the sample is not a conservative process, but runs through a hysteresis loop as explained by Sasaki and Tsukada (figure 2 in [17]). The jumps at the edges of the hysteresis loop are limited by phonon dynamics and are therefore very similar to direct thermal excitation.

The equipartition theorem predicts that the thermal motion of an atom with mass $m$ held by bonds with an atomic stiffness $k_{\text{bond}}$ has a thermally excited motion $\delta x$ in every spatial degree of freedom of

$$\frac{1}{2} k_{\text{bond}} \langle \delta x^2 \rangle = \frac{1}{2} \frac{k_b T}{m},$$

(1)


where $k_B = 1.38 \times 10^{-23}$ J K$^{-1}$ is Boltzmann’s constant. At a temperature of $T \approx 2000$ K and an estimated value for $k_{\text{bond}} \approx 300$ N m$^{-1}$, a total thermal motion of about 10 pm is estimated. This motion will certainly help to find the new local energy minimum that the diamond structure provides under high pressure.

Figure 3(a) displays the bonding energies of graphene and diamond after figure 4(a) of [6]. Note that the bonding energy of graphite and graphene are very similar, the one of graphite is lower by the exfoliation energy. This energy is required to release the vdW bond between two layers and amounts to approximately 30 meV per atom. Chemical bonds are typically two orders of magnitude larger.

The minimum for graphene is located at a distance of 142 pm, the bonding distance of the sp$^2$ bonds in planar graphene. For diamond, the minimum is at 155 pm, the bonding distance of the sp$^3$ bonds in diamond.

In bilayer graphene, the layers are spaced by 330 pm. Figure 3(a) plots the bonding energy of graphene and diamond as a function of the bonding distance, where the minimum of the graphene curve is at the bonding length of 142 pm and the minimum for diamond at 155 pm. Bilayer graphene has a vertical bonding distance of 330 pm. When rescaling the energy profile of figure 3(a) to the vertical dimension, we again have a local minimum of at 155 pm for diamond and one at 330 pm for bilayer graphene, obtaining figure 3(b). We can now take the derivative with respect to distance and obtain the force profile in figure 3(c). Note that this procedure is not exact, it is only an approximative method to obtain a qualitative idea for a force versus distance curve in the vertical direction.

3. Experimental setup

3.1. Scanning probe microscopy

We use combined scanning tunneling microscopy (STM) and AFM to probe the charge density at the Fermi level by STM and the bonding properties of the sample to the various tips by AFM in a single experiment, utilizing a combined scanning tunneling and AFM operating in ultrahigh vacuum at a temperature of 4.4 K (LT-SPM by Scienta Omicron Nanotechnology, Taunusstein, Germany) employing qPlus force sensors [18, 19].

The data with CO tip termination were recorded with a sensor type S1.0 (see table I in [19] for detailed specifications) with a stiffness of $k = 1800$ N m$^{-1}$, an eigenfrequency of $f_0 = 26 660.3$ Hz, an amplitude of
A = 50 pm and a quality factor of $Q = 42,844$. The bias voltage was 0.1 V unless noted otherwise. The data with the copper oxide (CuOx) tip were recorded with a sensor type M4 (table I in [19]) with $k = 1850$ N m$^{-1}$, $f_0 = 46,597.3$ Hz, $A = 50$ pm and $Q = 482,321$.

The instrument is controlled by a Nanonis BP4 SPM electronics with an OC4 oscillation controller (Specs-Nanonis, Zurich, Switzerland) and the higher harmonics were detected with a HF2 lock-in amplifier (Zurich Instruments, Zurich, Switzerland).

The tip was made of etched tungsten, cleaned by in situ field emission and formed by poking into a Cu(111) surface followed by checking the apex of the tip by carbon monoxide front atom identification (Zurich Instruments, Zurich, Switzerland) and the higher harmonics were detected with a HF2 lock-in amplifier.

An averaged gradient of the tip sample force is measured by frequency modulation (FM)—AFM [29].

The COFI method for tip characterization uses a CO molecule adsorbed on Cu(111) [20, 21] or Cu(111) surface followed by checking the apex of the tip by carbon monoxide front atom identification (Zurich Instruments, Zurich, Switzerland).

Here, we use single atom metal tips, trimer metal tips, single atom tips terminated by a CO molecule [23, 24] and CuOx tips [25–27] that have all been characterized by COFI. Information on the preparation of COFI. The COFI method creates a portrait of the tip that shows if a tip has a single atom, a dimer, trimer [20–22] or even a flake of graphene as in figure 21 at the front.

3.2. Experimental observables

We record a set of seven observables in each image in forward and backward scan direction that are used to create images and to analyze the interaction in detail:

(a) A tunneling current ⟨I⟩, where the averaging occurs over many oscillation cycles of the sensor that covers a vertical distance of twice the oscillation amplitude $A$. The tunneling current has an exponential distance dependence with gap width $z$ given by

$$I(z) = G_0 V_{\text{bias}} \exp(-z \kappa_1),$$

where $G_0$ refers to the conductance quantum $G_0 = 2e^2/h \approx 1/(12,906 \Omega)$, $e = 1.6022 \times 10^{-19}$ C is the elementary charge of the electron and $h = 6.626 \times 10^{-34}$ Js is Planck’s constant. The decay constant $\kappa_1$ is given by

$$\kappa_1 = \frac{\sqrt{2m_e \Phi}}{\hbar},$$

with electron mass $m_e = 9.11 \times 10^{-31}$ kg and work function $\Phi$ as explained in [28]. A typical decay rate of the tunneling current with distance is a reduction to 1/10 for every distance increase of 100 pm or 1 Å [28].

(b) An averaged gradient of the tip sample force is measured by frequency modulation (FM)—AFM [29]. The frequency shift is given by $\Delta f = f - f_0$, where $f$ is the actual oscillation frequency and the frequency shift $\Delta f = f_0 \cdot (k_0z)/(2k)$ [30], caused by the tip sample force $F_{ts}$ or more precisely an average over its gradient $k_{ts} = -\partial F_{ts}/\partial z$ as shown in equation (4).

If the perturbation of the tip-sample force to an oscillating cantilever is weak, the frequency shift is given by

$$\Delta f(z) = \frac{f_0}{k} \int_{-\infty}^{\infty} k_{ts}(z + uA) \sqrt{1 - u^2} \, du$$

that shows that the average tip sample force gradient $\langle k_{ts} \rangle$ is computed by a convolution of $k_{ts}$ over a semicircular weight function with a radius of the oscillation amplitude $A$ [31]. Experimental noise on this frequency shift measurement can be kept low by experimental precautions and choosing a low acquisition bandwidth [19, 32]. Note that all distance data $z$ have an arbitrary offset and $z = 0$ refers to the minimal distance of the experimental data.

The tip sample force $F_{ts}$ can be recovered from the frequency shift spectrum, using e.g. the Sader–Jarvis formula [33] or the matrix approach [31]. Recently, it was discovered that the presence of inflection points in the force curve requires to meet certain conditions regarding the oscillation amplitude $A$ in order to obtain a well-posed force deconvolution [34–36]. The deconvolutions performed here have been verified to be well-posed and thus valid.

(c) Higher harmonic amplitudes $a_n$ of order $n$ from 2 to 5, where the deflection $q$ of the cantilever is expressed as a Fourier series with $q(t) = \sum_{n=0}^{\infty} a_n \cos(n2\pi ft)$ with $a_0 = A = 50$ pm and $f = f_0 + \Delta f$. These higher harmonics arise when the tip-sample force $F_{ts}$ is not strictly linear with distance $z$. Dürrig [37] showed that the $n$th harmonic $a_n$ can be expressed as a convolution of the force curve with a Chebyshev function and proposed that the full force curve, separated into conservative and dissipative components, can be recovered from the amplitudes and phases of the higher harmonics in FM-AFM. Kawai et al have used this method to experimentally verify the force field over a KBr(001) surface [38].
The calculation of the higher harmonics as outlined in section VII A.1 of reference [39], and \( a_n \) can be expressed in terms of the Fourier components of the tip-sample force \( F_{ts} \) for \( n \neq 1 \):

\[
a_n = \frac{1}{1 - n^2} \frac{1}{\pi (1 + \delta_{n0})} \frac{1}{k} \int_0^{2\pi} F_{ts}(z + A \cos(\phi)) \cos(n\phi) d\phi. \tag{5}
\]

The constant cantilever deflection \( a_0 \) thus is given by the average force divided by the spring constant \( a_0 = \langle F_{ts} \rangle / k \), the amplitude at the fundamental oscillation frequency is \( a_1 = A \) and for \( n \geq 2 \), we find

\[
a_n = \frac{1}{1 - n^2} \frac{F_{ts}(n\omega)}{k}, \tag{6}
\]

where \( F_{ts}(n\omega) \) is the Fourier component of the time dependent tip-sample force \( F_{ts}(z + A \cos(\omega t)) \) at angular frequency \( n\omega \). For example, for \( F_{ts}(z_0 + z) = -1.8 \text{ nN} \cdot \cos(2 \arccos(z/A)) \) for \(-A \leq z \leq A\), we find \( a_2 = 333 \text{ fm} \) for a sensor stiffness of \( k = 1800 \text{ N m}^{-1} \), while for \( F_{ts}(z - z_0) = -1.8 \text{ nN} \cdot \cos(3 \arccos(z/A)) \) for \(-A \leq z \leq A\), we find \( a_3 = 125 \text{ fm} \). Figure 4 displays force versus distance laws that are needed to generate a frequency shift of -10 Hz and higher harmonics of order 2, 3 and 4 with a magnitude of 160 fm. It is noted, that measureable frequency shifts require only small force variations, while higher harmonics that rise above the experimental noise level require quite large force variations. Mathematically, the \( n \)th harmonic \( a_n \) can also be evaluated as a convolution of the \( n \)th force gradient with bell-shaped weight functions as presented in [40].

(d) A drive signal \( V_{exc} \), that excites and maintains a constant oscillation amplitude \( A \) of the cantilever. Dissipative tip-sample interactions lead to an increase of the drive signal from \( V_{exc} \) to \( V'_{exc} \). The dissipation per oscillation cycle \( \Delta E_{cs} \) is given by [19]:

\[
\Delta E_{cs} = \frac{\pi k A^2}{Q} \left( \frac{V'_{exc}}{V_{exc}} - 1 \right). \tag{7}
\]

3.3. Sample preparation
To study the graphite to diamond transition induced by an AFM, graphene [41, 42] on a carbon BL as in graphene grown on the Si side of silicon carbide [43] is an ideal sample, because the graphene top layer is stacked in an AB order [44] on top of the rigid SiC substrate. Graphene samples were produced by the group.
Figure 6. Top row: (a) left: calculated STM image of graphene for a metal tip (charge density at the Fermi level, arbitrary units). Right: calculated AFM image for a metal tip (given by total charge density in electrons per nm$^3$, scale inverted). (b) Calculated STM image of graphene for a CO terminated tip (see figure 9 for an explanation, arbitrary units). Right: calculated AFM image of graphene for a CO terminated tip (given by the total charge density expressed in electrons per nm$^3$). (c) Experimental constant height STM image (left), simultaneously recorded with the corresponding AFM image on the right recorded with a metal tip. (d) Experimental constant height STM image (left), simultaneously recorded with the corresponding AFM image on the right recorded with a CO terminated tip. Insets show COFI images of these tips, verifying that a metallic trimer tip was used in (c) and a CO-terminated tip was used in (d). Bottom row: force versus distance curves for the metallic trimer tip (e) and for the CO terminated tip (f). The insets show how the CO tip termination acts as a spacer, leading to only 1/5 of the vdW force in (f) compared to the trimer metal tip in (e).

Figure 7. Proposed mechanism where the CO tip penetrates into the surface, leading to a very large vdW attraction $F_{vdW}$ that is compensated by a large repulsive force $F_{CO}$, leading to a very small tip-sample force $F_{ts}$.

of Thomas Seyller by heating silicon carbide in an argon atmosphere [43]. The samples were inserted into a vacuum chamber, heated in situ for cleaning and transferred into the low temperature scanning probe microscope. This preparation led to a flat sample with large areas of single layer graphene as illustrated in figure 5. Two more variations of graphene have been imaged here: (a) the surface layer of highly oriented pyrolytic graphite (HOPG) that was prepared by cleaving with the scotch tape cleaving technique and (b) monolayer graphene flakes that are known to form from traces of carbon that are naturally dissolved in copper. Heating a Cu (110) crystal to approximately 800 K leads to migration of carbon to the surface and subsequent graphene formation.
4. Experimental data

Figures 5(a)–(c) shows the structure of monolayer graphene on a BL on top of a SiC crystal. The stacking of graphene over the BL leads to two inequivalent lattice sites A and B within the hexagonal unit cell (figure 5(a)), where A sits on top of an ABL site on the BL, while B is over a hollow site of the BL. The BBL site on the BL is underneath a hollow site of the surface graphene layer.

The SiC crystal is atomically flat with terraces with a width of a few μm separated by atomic steps and covered predominantly by monolayer graphene with patches of bi- and tri-layer graphene (figure 5(b)). Topographic STM images in figures 5(d) and (e) show the typical $6 \times 6$ domains and $6\sqrt{3} \times 6\sqrt{3}$ unit cells.
4.1. AFM with inert trimer tips and CO terminated tips

Figure 6 shows simulations and experimental STM and AFM data for a metal tip (left two columns) and for a CO terminated tip (right two columns), respectively. The STM contrast is calculated using the Tersoff–Hamann approximation [45] for $s$-type tips in (a) and considering the $p_x/p_y$ symmetry [46, 47] of the CO terminated tip in (b), see figure 9. Carbon atoms appear bright in the STM channel for metal tips (figures 6(a) and (c)) and dark for CO tips (figures 6(b) and (d)).

The conductance of undoped graphene is very poor as the density of states at the Fermi level is zero because of the Dirac cone electronic structure. However, the Fermi level is pinned at 400 meV for epitaxial graphene on the silicon side of silicon carbide such that even for small tunneling bias, a sizeable tunneling current can flow.

Forces between metal tips and the carbon atoms are attractive [48], therefore the calculated AFM image in figure 6(a) and the experimental AFM image in figure 6(d) show the carbon sites as minima. Figure 6(b) is a simulation of STM and AFM data for a CO terminated tip. The calculated AFM images use the total charge density as a measure of the attractive force for metal atoms [48] and the repulsive force in case of graphene and CO tips [49].

Figure 6(c) shows the experimental STM (left) and AFM (right) images of graphene using a trimer tip with the COFI portrait [20, 21, 50] of the tip in the inset of the AFM data. The data shows that the trimer tip is indeed sufficiently inert to allow for atomic imaging in both STM and AFM modes. Although graphite is a lubricant and two graphene layers even show superlubricity [51], single atom metal tips form strong bonds to graphene [48]. The formation of a strong bond between the metal tip and the graphene leads to rehybridization of the $sp^2$ bonds in graphene to $sp^3$ bonds [52], resulting in unstable imaging and even to ripping the graphene layer off the surface and covering the tip (see experimental data in the appendix A.1). Just as metal trimers on surfaces are much less reactive than monomers [53], trimer tips show much less short range attraction than single atom metal tips and thus enable stable imaging of graphene on SiC. The AFM image appears inverted with respect to the STM image, as the carbon atoms attract the reactive tip and attraction leads to a more negative frequency shift that is displayed dark.

Figure 6(d) displays the experimental STM (left) and AFM (right) images of graphene using a CO terminated tip, again with the COFI portrait of the tip in the inset of the AFM data. The STM data shows...
Figure 10. Effect of CO bending on image contrast and force spectroscopy. (a)–(c) Constant height AFM images of the average tip–sample force gradient \( \langle k_{ts} \rangle = 2k \Delta y / f_0 \) recorded at three distances where the \( z' \) scale corresponds to the \( z \) scale in figure 8.

(d) Contour line at larger distance where little CO bending occurs, yielding regular contrast where atoms appear bright.

(e) Contour line at close distance, where the O termination of the CO tip remains in the center of the graphene hexagon all the time, leading to strong CO bending and contrast inversion where hollow sites appear bright.

(f) Force versus distance spectra in the center of the hexagon or hollow site (open circles) and over an atom position (solid diamonds), showing a kink followed by a smaller slope at a distance of \( \approx 40 \) pm caused by CO bending.

maxima at the centers of the hexagons as discussed below while the AFM data shows an image of graphene that resembles the total charge density of graphene as expected for a tip that interacts by Pauli repulsion.

Overall, the agreement between calculated data in the top row and experimental data in the middle row of figure 6 is very good, deviations can be explained by the unavoidable asymmetries of the trimer tip and the tilt of the CO tip as discussed in figure 20.

Figures 6(e) and (f) display the force versus distance curves for the two different tips. Figure 6(e) shows a strong attraction between a Cu trimer tip and graphene that reaches 2 nN where the largest contribution is due to vdW interaction. The strong attraction between graphene and metal tips is a serious challenge for imaging graphene particularly for single atom metal tips (see figure 20 in the appendix). Even for trimer tips, it is possible that graphene sticks stronger to the tip than to the BL in SiC graphene. To maintain the integrity of the tip, we did not go close enough to probe the force minimum when recording figure 6(e).

Figure 6(f) shows a force versus distance curve that is observed for CO terminated tips. The maximal attraction reaches only 200 pN before repulsion between the CO terminated tip and graphene becomes noticeable. The AFM image in figure 6(d) right displays an experimental image of graphene that looks very similar to the calculated total charge density of graphene as, e.g. shown in figure 3D of [54]. The shallow increase of the magnitude of the attractive force with decreasing distance, indicated by blue lines in figures 6(e) and (f) points to its vdW origin. Highly reactive tips that interact by covalent chemical bonding would show an exponential increase of force with decreasing distance \( F \propto \exp(-z/\lambda) \) with \( \lambda \approx 50 \) pm. Thus, the CO terminated tips that have proven to image organic molecules such as pentacene very well [24] are also suited well for imaging graphene.
Figure 11. Effect of CO bending on image contrast and force spectroscopy. (a)–(c) Constant height AFM images of the average tip-sample force gradient $\langle k_{\text{ts}} \rangle = 2k \Delta f / f_0$ recorded at three distances where the $z$' scale corresponds to the $z$−scale in figure 8. (d) Contour line at larger distance where little CO bending occurs, yielding regular contrast where atoms appear bright. (e) Contour line at close distance, where the O termination of the CO tip remains in the center of the graphene hexagon all the time, leading to strong CO bending and contrast inversion where hollow sites appear bright. (f) Force versus distance spectra in the center of the hexagon or hollow site (open circles) and over an atom position (solid diamonds), showing a kink followed by a smaller slope at a distance of $\approx 40$ pm caused by CO bending.

Both force spectra are dominated by vdW forces. The vdW force between a sphere of radius $R$ and a flat surface at distance $z$ is given by [55]:

$$F_{\text{vdW sphere}}(z) = -\frac{A_H R}{6z^2}, \quad (8)$$

where $A_H$ is the Hamaker constant which has a magnitude of about 1 eV for many combinations of solids [55]. A somewhat weaker distance dependence $\propto 1/z$ results for conical and pyramidal tips, where the vdW force for a conical tip with full angle $\alpha$ is given by [30]:

$$F_{\text{vdW cone}}(z) = -\frac{A_H \tan^2(\alpha/2)}{6z}. \quad (9)$$

For spherical and conical tips, the vdW force diverges for $z \to 0$. However, $z$ cannot reach zero as it refers to the distance between the center of the front atom of the tip and the plane connecting the atomic nuclei of the surface layers, yielding a minimal value for $z$ of about 200 pm. For blunt metal tips, e.g. a spherical tip with a radius of $R = 30$ nm or a conical tip with an opening angle of $170^\circ$, equations (8) and (9) yield a maximal vdW attraction of $F_{\text{vdW}}(z = 200 \text{ pm}) \approx -20 \text{ nN}$ for $A_H = 1 \text{ eV}$.

The force spectra in figures 6(e) and (f) reveal that the difference in short range force is about 200 pN between the hollow and the atomic sites for metallic trimer tips in (e), thus we conclude that the 2 nN attraction is mainly due to vdW forces. For CO terminated tips, the vdW forces are much smaller as the CO molecule acts like a separator that keeps the blunt metal tip further away from the flat sample. For the CO terminated tip, the hollow- and atom sites are barely distinguishable in the force spectra for distances of $z > 75$ pm, merging into a difference of about 20 pN for smaller distances once the Pauli repulsion is in effect.

Experimentally, we can only measure the total tip-sample force $F_{\text{ts}}$ by AFM, with

$$F_{\text{ts}} = F_{\text{front atom}} + F_{\text{background}}, \quad (10)$$
where the background force usually has a long range and is mainly due to vdW forces $F_{\text{vdW}}$, unless a large bias voltage is applied that would add electrostatic long range forces. Hence, it is not possible to separate the long-range vdW part from the short range interactions that originate from the front atom of the tip. When studying individual adatoms, the on–off technique as introduced in [56] can be used. In this technique, the short-range force is given by the difference between the tip-sample force spectrum over the adatom and the force spectrum on a flat section of the sample nearby [56]. However, this technique is not applicable for flat surfaces, where the vdW contribution has to be guessed from the force characteristics before making contact (blue lines in figures 6(e) and (f)).

Because of the divergence of the vdW force for small distances, the estimated vdW contribution can be grossly in error for small distances as will be discussed further below. Figure 7 illustrates where a relatively small total tip-sample force $F_{ts}$ is composed of a strong attractive background force $F_{\text{background}} = F_{\text{vdW}}$ and a repulsive force of the front atom $F_{\text{front atom}} = F_{\text{CO}}$.

Resolving graphene atomically by STM is straightforward [41] although the large forces between metallic STM tips and graphene can pull the graphene significantly out of equilibrium, as demonstrated in graphene drumheads [16]. Atomically resolved imaging by AFM with metallic tips is more challenging because the atomic spacing of 142 pm in graphene is only about half of the diameter of a metal tip atom. Simultaneously recorded STM and AFM images of graphite with tungsten tips have shown large lateral offsets between the current and force channels [54]. This lateral shift has been attributed to the large diameter of metal front atoms and their varying contributions to current and force of different atomic orbitals. For CO terminated tips, where the diameter of the O front atom is only 40% of the diameter of a metal atom, the registry between STM and AFM images is close to perfect as evident from figures 6(d) and 8.

Carbon monoxide terminated tips mounted on qPlus force sensors [19] are now a standard in AFM [57] after they have successfully been employed to image a pentacene molecule [24], other organic molecules

Figure 12. (a) Third harmonic amplitude $a_3$ for six different heights spaced by 6 pm. A noticeable amplitude $a_3$ first occurs in the center of the graphene hexagons in (b), turning to rings for closer distances in (d) with increasing diameters $d_{\text{HH}}$ (e) and (f). Note that the scan frame is not perfectly parallel to the sample surface and the $z$-distances at the lower edge are approximately 4 pm smaller than at the top edge. (g)--(l) Schematic view showing a tilted CO tip termination. (m) and (n) The ring diameter $d_{\text{HH}}$ follows a simple Pythagorean relation $\left(\frac{d_{\text{HH}}}{2}\right)^2 = r_{\text{Cu+CO}}^2 - z'^2$. 

14
Figure 13. (a) High resolution image of $a_3$ close to the distance where two adjacent higher-harmonic rings merge. Parameters: $V_{tip} = 0$ V, image size $0.5 \times 0.5$ nm$^2$ at $64 \times 64$ pxl, scanning speed $125$ pm s$^{-1}$. (b) Distance spectrum of the third harmonic amplitude $a_3$ recorded over the hollow site. (c) Geometry of the CO terminated tip trapped with its O termination in the hollow site (see text).

Figure 14. Geometry of the SiC substrate, covered by a BL of graphene and an additional graphene layer. The CO terminated tip oscillates at a reduced amplitude $A'$ once the repulsive forces become significant (see text).

[58, 59], graphene [49] and even molecules that extend into the third dimension normal to the surface [60]. Here, we find that while CO terminated probes truthfully image graphene, CO bending strongly affects the imaging process on the picometer scale.

4.2. Constant height data sets recorded with a CO terminated tip

Figure 8 shows the detailed contrast development in a set of six constant height images of tunneling current, frequency shift $\Delta f$ and third harmonic amplitude $a_3$. Distances get consecutively closer by decrements of $12$ pm over a range of $60$ pm, starting at (a) the largest distance of $z = 60$ pm, at (b) $z = 48$ pm etc. The left column shows constant height images of the tunneling current, the center column is the frequency shift and the right column shows the third harmonic amplitude. The contrast in the tunneling current (left column) does not change significantly over the small distance variation of $60$ pm from (a) to (f). The current maximum is located at the hollow site for all distances as predicted by the calculation presented in figures 6(b) left and 9. The frequency shift produces an image of graphene similar to those obtained by transmission electron microscopy in figure 1 of [62] or as expected from the total charge density calculation in figure 1(b) at $z = 60$ pm. For smaller distances, the bonds appear to become sharper from figure 8(a) center down to figure 8(c) center at $z = 36$ pm. This sharpening effect has been observed in imaging organic molecules and explained already in 2012 [59], it is caused by CO bending.

At $z = 24$ pm, the contrast starts to invert and the third harmonic $a_3$ rises over its noise level at the hollow sites. Interestingly, of the six monitored higher harmonic amplitudes $a_2 - a_7$ [40], only $a_3$ rises above the noise level of about $100$ fm (see appendix with figure 23). At $z = 12$ pm, the contrast inversion is fully developed and the higher harmonics show a ring-like structure. The contrast inversion in the frequency shift as well as the rise of the higher harmonics starting at the hollow sites is explained in the following. When the tip is above a hollow site, the oxygen end of the CO molecule becomes trapped in the hollow site and cannot bend laterally.

Our hypothesis, supported by additional considerations further below, is as follows. When the local pressure reaches the value that is required to induce a carbon A atom from the graphene surface layer and a carbon ABL atom from the underlying BL to rehybridize from three sp$^2$ and one p$_z$ orbital to four sp$^3$ orbitals, a strong anharmonic force component arises that leads to the rise of $a_3$. Because the magnitude of
Figure 15. (a)–(d) Third harmonic amplitude $a_3$ (left) and energy dissipation of the sensor per oscillation cycle (right) at distance decrements of merely 5 pm per frame, i.e. the whole $z$–range covered here is only 20 pm. The onset of a few meV of dissipation for very small distances indicates a hysteresis in the force versus distance curve. Parameters: $V_{\text{tip}} = 0$ V, image size $500 \times 500$ pm$^2$ at $64 \times 64$ pxl, scanning speed $125$ pm s$^{-1}$.

$a_3$ is only on the order of 100 fm (only 2/1000 of the oscillation amplitude $A = a_1$), the lock-in amplifier that measures $a_3$ has to be set to a fairly low bandwidth (here, 5 Hz) to obtain a good signal-to-noise ratio [61].

The frequency shift is more positive over the carbon atoms from frame (a) to (c), pointing to repulsive interactions while a contrast inversion starts to develop at (d) and is fully developed at (e) and (f). The contrast inversion is due to a lateral bending of the CO molecule. The third column, the third harmonic amplitude, becomes noticeable at the height where contrast inversion starts to emerge, explained in section 2. The apparent bond sharpening that becomes noticeable from (c) in the frequency shift channel is caused by CO bending as discussed in figure 22 and pertaining text.

4.3. Interpretation of the constant height data

4.3.1. Calculation of STM data for graphene with CO tips

The calculation of the STM current for CO terminated tips with its, in comparison to metal tips, inverted contrast as well as the influence of a tilt of the CO tip out of an exact vertical orientation is addressed in figure 9. In graphene, electron energies rise linearly with momentum at the Fermi energy rather than
Figure 16. Distance dependence of (a) frequency shift $\Delta f(z)$, (b) tip-sample force $F_{ts}(z)$, (c) logarithm of tunneling current in $\text{pA}$ and (d) third harmonic amplitude $a_3(z)$ as a function of vertical distance $z$. The data was recorded in the center of a hexagon (hollow site), see figure 5(a).

4.3.2. Simulating frequency shift data with the probe particle model

The interpretation of AFM data on organic materials recorded with CO terminated tips is often performed using the probe-particle model after Hapala et al [65]. This model assumes a Lennard–Jones interaction between the O front atom of tip and allows for a lateral bending of the CO termination. Standard parameters after [65] were used to plot $\Delta f$ images in steps of 12 pm. For the simulations (as in the experiment), $k = 1800 \text{ N m}^{-1}$ and $f_0 = 30\,300 \text{ Hz}$. The graphene is a flat sheet of carbon that was not allowed to relax. No electrostatics were included in the calculation.

A comparison of the experimental frequency shift data in figure 8 with the probe-particle model simulations in figure 10 shows a convincing agreement, starting with an image that resembles the total charge density of graphene in (a), and followed by apparent bond sharpening from (b)–(d). This apparent
bond sharpening with decreasing distances was already observed by Gross et al in 2012 in figure 2 of [59]. Further distance reduction leads to incipient and complete contrast reversal from figures 10(d)–(f). CO bending also is responsible for a crossover in local force versus distance spectra.

4.3.3. CO bending leads to crossover in force versus distance spectra

When the CO terminated tip is directly above an atom and interacts via Pauli repulsion, it is in an unstable equilibrium. As the lateral stiffness of CO terminated tips is small, only about 0.25 N m\(^{-1}\) [66], small lateral forces will bend the CO tip termination to the side.

Figure 11 shows two consequences of CO bending. First, we again observe apparent bond sharpening in figure 11(a), transiting into incipient (b) and full (c) contrast reversal. The mechanism for this is illustrated in figures 11(d) and (e). For intermediate distances, the CO tip termination stays almost perfectly upright and the contour of the frequency shift at constant height would show a positive value right above the carbon atoms as sketched in figure 11(d). When moving very close, the oxygen termination of the CO tip is caught in the hollow site of the graphene hexagon and the CO termination pivots around the hollow site of graphene, causing a contrast inversion in the constant height frequency shift images in figures 11(a)–(c).

The second consequence of CO bending is apparent in the force versus distance spectra shown in figure 11(f). The force versus distance spectrum over the center of the hexagon results in a smooth force distance curve because the oxygen end of the CO terminated tip remains stable in the center of the hexagon while local repulsive forces start to act. Note that although the net force on the tip is still attractive at a distance of 100 pm, the force on the CO tip is already repulsive, i.e. positive. The strong vdW attraction of the tip that is larger in magnitude than the repulsion to the CO tip leads to a net negative force on the tip. When performing the force spectrum over the atom position in figure 11(f), the CO molecule at the tip starts to bend at a distance of around 40 pm, leading to a crossover of the two force curves.

An inverted crossover, caused by a completely different reason, has been observed by Boneschanscher et al in figure 1(c) of [49] for graphene on iridium using a metallic tip. The crossover region in figure 1(c) of [49] is smooth and was explained by turning an initially attractive short range interaction between the tip and the carbon atoms in graphene to repulsion for smaller distances. COFI images of the tip are not presented in [49], and as these authors did not report instability issues as those observed here in figure 9, we suspect that either graphene adheres much stronger to Ir(111) than to the BL on SiC, or their tip was also a fairly inert metallic trimer tip similar to the one used in figure 6 rather than a highly reactive single atom metal tip as presented in figure 9.

4.3.4. The emergence of higher harmonics for close distances

The right column of figure 8 shows that the third harmonic \(a_3\) starts to become noticeable at a distance (row figure 8(d)) where the contrast inversion starts to emerge and the repulsive force between the CO tip termination becomes large enough to cause considerable tip bending.

The emergence of a third harmonic requires the tip to go through a force profile as plotted in figure 4(c), i.e. overcoming a force barrier, followed by a reduction in force and again followed by an increasing force. The qualitative figure 3(c) would provide a similar force profile, suggesting that the emergence of \(a_3\) is related to overcoming a strong force barrier such as occurs in the graphite to diamond transition.

While \(a_3\) starts to emerge at the hollow sites, further approach leads to a ring-like appearance. We now look closer at this emergence of rings in \(a_3\) in figure 12. The weak lateral stiffness of the CO terminated tip allows it to pivot around the center of the metallic front atom of the tip with an arm of \(d = r_{\text{Cu}} + r_{\text{CO}}\) as displayed in figure 12.

Figure 12(a) shows the evolution of third harmonics \(a_3\) with decreasing distance at distance decrements of merely 6 pm. Note that there is a slight tilt of about 0.5\(^\circ\) in the 500 pm \(\times\) 500 pm scan frames—the top line is approximately 4 pm closer than the bottom line, therefore \(a_3\) becomes noticeable first at the bottom of figure 12(b). The third harmonics start to emerge at the hollow sites and evolve into ring-like structures for closer distances, in accordance with the model of their origin presented in figure 12(m) that shows that the diameters of the ring-like structures follow a simple geometric relation \((d_{\text{HH}}/2)^2 = r_{\text{Cu+CO}}^2 - x^2\) plotted in figure 12(n).

Figure 13(a) shows a constant height image of the third harmonic \(a_3\) at a close distance where the rings of adjacent unit cells merge, leaving a triangular local minimum at one of the two lattice sites A and B. The small width \(\Delta x \approx 20\) pm of the roughly circular profile of \(a_3\) shows that the distance dependence of \(a_3\) at the hollow site plotted in figure 13(b) even overestimates the \(z\) width of \(a_3\). With an estimate for the arm length \(d\), we find a maximal deflection \(\phi \approx 18^\circ\), and with the geometry considerations in figure 13(c) we find \(\Delta z \approx \Delta x \times \tan\phi \approx 7\) pm.

Thus, the third harmonic occurs over a very thin slice in \(z\) direction of only about 7 pm. Given that the weight function to derive \(a_3\) (see equation (5)) has a width of about the actual oscillation amplitude \(A'\) at
which the front atom of the tip and its next sample atom oscillate, it shows that this actual oscillation amplitude $A'$ between the tip’s front atom and graphene in the squeezed material must be much smaller than the oscillation amplitude of the sensor itself with $A = 50$ pm. This reduced amplitude can be explained by the repulsive load of the tip over the relatively soft bilayer graphene. Figure 14 is a schematic view of the sample and tip arrangement. Although the tip oscillates at amplitude $A$, the distance between the graphene surface layer and the tip will oscillate at a much smaller amplitude $A'$ once the repulsive load becomes large enough to compress the soft bonds between the BL and the graphene.
Figure 18. Force versus distance spectra for a CuOx tip over a hollow site (a) and a carbon atom (b). (a) The sharp fall of the spectrum taken at the hollow site at $z=30$ pm points to a possible onset of a graphite to diamond transition. (b) When taking the spectrum over a carbon atom, we expect that both sample- and tip deformations lead to a lateral displacement, finally ending with the oxygen front atom in the hollow site, causing a delayed force induced transition.

Figure 15 shows another high resolution image of $a_3$ supplemented by dissipation data on the right column. The presence of dissipation after equation (7) shows that the tip oscillates non-adiabatically, exciting bursts of phonons of a larger amplitude than those present at the sample temperature of about 4 K for each oscillation cycle.

In a single graphene layer, A and B sites are equal. As mentioned above, for graphene on the BL of SiC, A and B sites are different, because A sites have $A_{BL}$ atoms in the BL underneath while B sites are above hollow sites. If the hypothesis about the graphite to diamond bonding transition is correct, an asymmetry between A and B sites is expected. We expect the A sites to be reduced in height with respect to the B sites. Figure 15 shows high resolution data of $a_3$ and the energy dissipation per oscillation cycle for four different heights spaced by merely 5 pm. The third harmonic $a_3$ shows clear differences between A and B sites in figures 15(c) and (d), and the dissipation data shows these differences clearly in (c) and in particular in (d), supporting the bonding transition hypothesis.

4.4. Distance spectra

The physical nature of the tip-sample interaction is revealed by the distance dependence of the pertinent physical observables. Figure 16(a) shows the distance dependence of the frequency shift recorded with a CO terminated tip that was placed over a hollow site, i.e. the center, of a graphene unit cell. The deconvoluted force in figure 16(b) has only a very small magnitude, reaching merely $-300$ pN. However, this inconspicuous force curve is incompatible with the distance dependence of two more observables, the averaged tunneling current and the third harmonic $a_3$.

The averaged tunneling current $\langle I \rangle$ is displayed in a logarithmic scale in figure 16(c), it is at or below the noise level of about 1 pA for $z > 200$ pm, rising in the expected exponential fashion after equation (2) until $z$ reaches about 70 pm, where the slope becomes noticeably smaller. This reduction in slope points to the emergence of repulsive forces, as previously observed in local tunneling versus distance spectroscopy by Schull et al [67] on buckyballs.

The emergence of a strong third harmonic in figure 16(d) is interesting and possibly caused by the graphite to diamond transition discussed earlier. The force versus distance relation that is needed to generate a third harmonic of a magnitude of 0.2 pm, again derived from equation (5), is a force that swings up and down by almost 3 nN as displayed in figure 4.

5. Images of graphene using CuOx tips

The introduction of CuOx tips [25] provided tips that are as inert as CO terminated tips but more stiff.

The termination of AFM tips with a CO molecule has three profound benefits: first, CO terminated tips are chemically highly inert, interacting mainly by Pauli repulsion [68], where so far, our group only found Cu and Fe as exceptions [12]. Second, the atomic radius of O is small. Therefore, they allow high spatial resolution. The lateral softness can lead to imaging artifacts, although sometimes the flexibility of the tips provides greater contrast. Third, the long CO molecule acts like a distance holder that keeps the blunt metal tip further from the flat surface, reducing vdW attraction.
Figure 19. Calculation of total charge density for graphene (a) and high resolution constant height images of graphene using a CO terminated tip (b) and a CuOx tip (c). Image size 0.6 nm by 0.6 nm. Parameters: (a) the total charge density is calculated at a height of 200 pm above the plane that connects the nuclei of the carbon atoms. The charge density ranges from 2 (dark) to 4 (bright) electrons per nm$^3$, the contour lines are spaced by 0.2 electrons per nm$^3$. (b) Sensor type S1.0 (see [19] for specifications) with $k = 1800$ N m$^{-1}$, $f_0 = 26660.3$ Hz, $A = 50$ pm and $Q = 42844$. The contour lines are spaced by 0.1 Hz, the frequency shift ranges from $-8.2$ Hz (dark) to $-4.9$ Hz (bright). (c) Sensor type M4 (see [19] for specifications) with $k = 1850$ N m$^{-1}$, $f_0 = 46597.3$ Hz, $A = 50$ pm and $Q = 482321$. The contour lines are spaced by 0.1 Hz, the frequency shift ranges from $-33.5$ Hz (dark) to $-30.2$ Hz (bright).

In 2016, Mönig et al [25] introduced CuOx tips, tips that share the first and second benefit of CO tips yet providing greater lateral stability. We have used those tips to image graphene on SiC as well.

Figure 17 shows again a constant height set where the tunneling current and frequency shift data look similar to the data set taken with a CO terminated tip of figure 8 for relatively weak repulsive forces up to (b). However, they do not show significant third harmonics, possibly because the bluntness of CuOx tips does not allow them to push a single unit cell of graphene significantly closer to the BL as other parts of the blunt tip are expected to touch neighboring unit cells. However, one of the two lattice sites A, B is significantly harder than the other, as evident from the emergence of a strong A–B site difference for heights below 20 pm from figure 17(d).

Also, the force versus distance curve over the hollow site displayed in figure 18(a) shows a pronounced step at a $z$ value of 30 pm that points to a collapse of the lattice and a possible start of rehybridization. When performing the spectrum over an atom site in figure 18(b), the transition is softer and at closer $z$ values as expected.

6. Images of graphene under weak repulsive forces for CO-terminated and CuOx tips

In scanning transmission electron microscopy, progress in collimating the electron beam to a diameter close to 130 pm enabled the resolution of the atomic lattice and defects in graphene [43]. Graphene on SiC has been studied before by FM-AFM in vacuum at low temperatures [49], room temperature [69] and in ambient conditions [70]. The data here observed in a slight repulsive regime reveals a very subtle detail. Figure 19(a) shows a charge density calculation of graphene at a height of 200 pm over the planes connecting the C nuclei. Slater-type orbitals were used for the calculation as in [54], showing weak but noticeable local maxima over the atom sites. When operating AFM with CO or CuOx tips at small repulsive forces, the experimental images clearly show the local maximum of charge density over the carbon atoms as displayed in figure 19(b) for CO terminated tips and in figure 19(c) for CuOx tips. The overall contrast is similar for CO and CuOx tips, but the absolute value of the frequency shift is more negative for CuOx tips due to the larger vdW attraction.

7. Summary

To summarize the unification of the experimental observation of higher harmonics and the DFT data, we can explain the higher harmonics as a consequence of local and temporary transition of graphene layers into a diamond structure. However, we can only explain the fivefold increase of vdW attraction to values up to 10 nN by a runaway increase of vdW attraction due to indented graphene snuggling up closely against a relatively blunt metal tip. We also note that while the data was perfectly reproducible with that one tip, other tips that were possibly sharper did not produce the higher harmonics. Early data from 2004 [40], a time where tip preparation of AFMs was much less sophisticated and tips were known to be very blunt indicated by large negative frequency shifts showed even much higher magnitudes of higher harmonics, pointing to
the importance of large runaway vdW attraction for the induction of a graphite to diamond transition and the subsequent generation of higher harmonics.

We note that as we do not yet have a recipe on how to prepare tips that reproduce the observed data, we cannot consider our data to be a proof for our suggested interpretation. In principle, some very odd tip artifact might also cause strong $a_3$'s. However, we routinely record higher harmonics on our low temperature microscopes and we have not ever observed such clear and strong higher harmonic signals with other CO terminated tips. One of the reviewers suggested that a tip that slides out of the local energy minimum in the center of the graphene hexagon might result in a force curve similar to the one plotted in figure 4(c). Also, one could speculate that a breaking of the bond between the CO molecule and the metallic front atom, followed by rebonding to a metal atom in the second atomic layer of the tip might result in a similar force curve. However, we doubt that the current and frequency shift data would continue in such a smooth way over distance reduction in figures 8, 12 and 15.

In summary, we have found strong additional evidence to references [9, 10] that the tip of an AFM can cause a pressure induced transition from an $sp^2$ type bonding symmetry to a diamond-like $sp^3$ type bonding. The picometer scale study allows us to describe this process on the atomic scale, opening up high-pressure material science to the atomic dimension. In the future, it would be desirable to supplement these studies with high speed optics to learn about chemical bond evolution in the time domain.

8. Remarks about the time sequence of these data

Indications for the proposed transition from $sp^2$ graphene to $sp^3$ diamond bonding induced by the pressure of an AFM tip may have been unknowingly observed decades ago. In 1987, Binnig observed the flipping rate of graphene in a low temperature STM operated in liquid helium [72], showing strong A-B asymmetry of the flipping rate in figure 3(b) of [72]. This could have been caused by a similar outward-motion of C atoms induced by an $sp^2$ to $sp^3$ transition as found here in figures (e)–(g). In 2004, Hembacher et al [40] used graphene to image a tungsten tip, where the front atom of a tungsten tip was imaged by either the A or B atoms of the graphite sample using strong creation of higher harmonics in the sensor oscillation. A local transition to diamond would explain the experimental observation of only one front atom image per unit cell of graphite, i.e. a dangling $sp^3$ bond that reaches from rehybridized B atoms into the vacuum.

The experimental observation of a strong third harmonic that marks overcoming a force threshold with subsequent lowering and increase of the force has only been observed with one CO terminated tip so far. Although this tip lasted for about 34 h, generated about 900 images (64 constant height datasets with 7 data channels and two scan directions) and experienced $\approx 9 \times 10^9$ approach-retract cycles over a $2A = 100$ pm distance range (oscillation at 26.6 kHz for 34 h), we did not succeed so far to prepare a CO terminated tip that showed the same resilience.

As outlined further below, a CO terminated tip needs to fulfill two criteria to be able to induce a graphite-to-diamond transition and to generate a large third harmonic in the cantilever oscillation: (1) the CO terminated tip needs to sustain a repulsive force on the order of 10 nN; (2) it needs to have a fairly flat metal base to generate a strong vdW attraction. A recent study has shown that CO terminated tips that are all formed by attaching CO to a single metallic front atom of the tip still show a significant variation in there force versus distance curves over metal clusters (figure 1 in [53]).

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Data availability statement

All data that support the findings of this study are included within the article (and any supplementary files).
Figure 20. STM and AFM data with a reactive single metal tip. (a) Atomically resolved STM data, while the AFM data in (b) does not provide atomic resolution. (c) Constant height STM data where the graphene flake jumps towards the tip by about 150 pm, enabling atomic contrast in the AFM channel in (d) as well. (e) STM and (f) AFM data with atomic resolution with an unstable tip junction. (g) Constant height STM data and trace (h) showing a jump in current of more than 10, indicating a distance reduction of about 150 pm. (i) Scan area of (a)–(g).

Appendix

A.1. Combined STM and AFM with reactive single atom metal tips

The tip is crucial for image contrast in STM and AFM. It appears to be sensible to use a sharp single atom metal tip. However, Ondrácek et al [48] calculated that single metal tips are very reactive even with respect to graphene and a tungsten tip can form a chemical bond with graphene where the attractive force reaches 2 nN [48]. This may come as a surprise, as graphite is a widely used lubricant. The graphene layers in graphite slide with very low friction on each other, even leading to superlubricity when those graphene layers are rotated as reported by Dienwiebel et al [51].

Careful experiments have shown that the highly reactive single atom metal tips allow to obtain atomically resolved images as depicted in figure 20. At large distance, the attractive forces to the tip are small enough such that atomically resolved STM data is obtainable in figure 20(a), while the AFM data in figure 20(b) does not show atomic contrast yet. Figures 20(c), (d), (g) and (h) record an event, where the graphene surface layer apparently moves about 150 pm closer to the tip as indicated by the 30-fold jump in current in figure 20(h). At these close distances, atomic resolution is observed in both the STM channel in figure 20(e) and the AFM channel in figure 20(f).

Recently, it was verified experimentally that single metal adatoms on a surface are more than twice as reactive as trimer clusters [53]. Specifically, a single Fe atom on Cu(111) can exert up to 450 pN of attraction to a CO terminated tip, while a Fe trimer only exerts a maximal force of 200 pN [53]. We expect a related effect for single atom versus trimer tips and the reduced reactivity of trimers versus monomers on surfaces is consistent with metallic trimer tips allowing atomic imaging of graphene, while single atom tips pose a danger to pull the graphene layers towards the tip.

Figure 21(a) shows a topographic STM image where a monolayer graphene on silicon carbide sample was scanned and the attractive forces between tip and sample were so large that the tip picked up a part of the graphene surface layer causing strong noise in the upper half of figure 21(a). This pickup of graphene...
flakes by a reactive metal tip, illustrated in figure 21(b), occurred quite frequently and leads to a metal tip that has a graphene cover wrapped around it as shown in figure 21(c).

Inspection of these compromised tips by COFI [20] is used to study and image the tip in figure 21(d) after the event that occurred in the midst of recording figure 21(a). Similar events occurred many times when starting with a monoatomic metal tip and the tips were always covered by graphene after these instabilities as shown in figures 21(e) and (f). While it was still possible to image graphene atomically with these graphene covered tips, the force versus distance spectra showed pronounced shoulders as evident in figure 21(g). The appearance of the shoulder in figure 21(g) is caused by a change in the dominating vertex of the graphene sheet that covers the tip induced by the reduction of distance, also accompanied by a large lateral shift of constant height data for small height variations (not shown here, see figure A.2 in [71] for details). Graphene terminated tips provided noisy images in the STM channel (figure 21(h)) and also in the AFM channel (figure 21(i)), and cleaner data can be obtained with metallic trimer tips as shown in the following.

A.2. Correction of CO bending
A further demonstration of the effects of CO bending is provided by figure 22, where we first use the normal force data from figures 22(a)–(c) to compute the interaction energy between tip and sample and then take the lateral derivative to calculate the CO deflection and to correct the image [63]. By utilizing the energy profile, calculating lateral forces from it and determining the lateral CO deflection [63, 66], the CO deflection can be determined and the images can be corrected as shown in figure 22. The correction...
algorithms computes the lateral forces acting on the CO terminated probe tip, calculates its lateral bending and corrects the ‘true’ lateral position of the image data, correcting the apparent bond sharpening and leading to warped edges of the corrected images. The correction algorithm shows that the apparent sharpening of the images for closer distances is an artifact caused by the lateral bending of the CO terminated tip.

A.3. Higher harmonics of a different order and illustration of higher harmonics generation

One striking character of the higher harmonics is their very strong distance dependence. Another interesting property is that among the higher harmonics from $a_2$, $a_3$, $a_4$ and $a_5$, only $a_3$ rises significantly above the noise, as displayed in figure 23. Although there is a weak signal in $a_2$ in figures 23(a)–(c), only $a_3$ is large enough to allow for a detailed analysis.

This particular representation of the higher harmonics allows for a descriptive graphic representation in figure 24—$a_m$ is the $m$th Fourier coefficient of the force when expressed a function of $z + A \cos \phi$.

In particular, for $m = 3$ we find:

$$a_3 = -\frac{1}{8k \cdot \pi} \int_0^{2\pi} F_s(z + A \cos \phi) \cos(3\phi) d\phi.$$  \hspace{1cm} (11)

The magnitude of the higher harmonics is very small: for a force sensor with $k = 1800$ N m$^{-1}$, a third order amplitude $a_3 = 200$ fm corresponds to a Fourier component of 2.88 nN. This also implies that in practice, higher harmonics can only be detected at a very low detection bandwidth $B$. The noise in the higher harmonics is given by $\delta a_m = n_q(f_0) \cdot \sqrt{B}$, where $n_q(f)$ is the frequency dependent deflection noise density that varies little with frequency (see section IV B in [19]) and is approximately 100 fm (\sqrt{Hz})$^{-1}$.

Figure 22. Raw data (a), (c), and (e) and corrected data (b), (d), and (f) for short range potential energy plots deconvoluted from constant height frequency shift images. (g) The CO terminated tip is deflected laterally by an angle $\alpha$ that ranges from 0 to 3.7$^\circ$ as shown in (h).
Figure 23. Constant-height images of tunneling current, frequency shift and higher harmonics $a_2$, $a_3$, $a_4$, and $a_5$ for a CO terminated tip over a distance range of 48 pm from (a) to (d) with decrements of 12 pm. Among the higher harmonics, $a_2$ may show a weak signal over the noise in (a)–(c), but $a_3$ rises very significantly above the experimental noise [61].

Figure 24. Generation of higher harmonics $a_n$ of order $n$. The simple relation $a_n = F_{nf}/(k(1 - n^2))$ where $F_{nf}$ is the Fourier component of the tip-sample force at frequency $n \cdot f$ yields a straightforward interpretation. Pseudo-3D representations of force fields $F(z)$ with $z = A \cos \phi$, $\phi = 2\pi ft$ with oscillation frequency $f$ and time $t$ and $\zeta = A \sin \phi$ where the force is plotted in the vertical axis. (a) Force $F \propto z$ that generates a frequency shift, but no higher harmonics. (b) Force $F \propto \cos(2 \arccos(z/A))$ that generates a nonzero $a_2$, but neither a frequency shift nor higher harmonics of an order other than 2. (c) Force $F \propto \cos(3 \arccos(z/A))$ that generates a nonzero $a_3$, but neither a frequency shift nor higher harmonics of an order other than 3. (d) Force $F \propto \cos(4 \arccos(z/A))$ that generates a nonzero $a_4$, but neither a frequency shift nor higher harmonics of an order other than 4. (e)–(h) 3D representations of the corresponding force fields $F(z)$. 
Figure 25. (a)–(d) Tunneling current, frequency shift and third harmonic amplitude $a_3$ for HOPG for decreasing distances. (e)–(h) Tunneling current, frequency shift and third harmonic amplitude $a_3$ for a small flake of graphene on Cu(110) for decreasing distances. The contrast of tunneling current (left column) and frequency shift (center column) shows a similar distance dependence as for graphene on SiC shown in figure 8. However, the higher harmonics $a_3$ remain below the noise level of about 100 fm.

Figure 26. Constant-height image of small graphene flakes that formed by thermal annealing of Cu(110), accompanied by two adsorbed CO molecules. The square frame shows the scan area imaged in figures 25(e)–(h).

A.4. Examples of imaging graphene with CO tips on other substrates

In figures 25(a)–(d), we look at HOPG with a CO tip and observe the same evolution in contrast for the tunneling current and frequency shift as observed for graphene on SiC above, but no higher harmonics. The same happens for graphene on Cu(110) as shown in figures 25(e)–(h).

Figure 26 shows images of the graphene flakes with two random CO molecules that formed on Cu(110), where their small size explains why the STM data in figures 25(e)–(h) is highly irregular, related to the irregularities that are observed in STM on multilayer graphene [73].

Figure 27 is an extended constant height set of figure 17 showing current, frequency shift and third harmonic over graphene with CuOx tips where the $z$ distance samples are spaced by merely 5 pm.
Figure 27. Constant-height images of tunneling current, frequency shift and third harmonic $a_3$ for a CuO$_x$ tip over a distance range of 50 pm from (a) to (f) with decrements of 5 pm. At $z = 50$ pm, the frequency shift image is close to the total charge density of graphene, i.e. a natural image of the surface. For closer distances, distortions start to appear before a pronounced asymmetry between A and B sites emerges at distances at and below $z = 10$ pm.

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References

[1] Hazen R M 1999 The Diamond Makers (Cambridge: Cambridge University Press)
[2] Bundy F P, Hall H T, Strong H M and Wentorf R H 1955 Man-made diamonds Nature 176 51
[3] Bovenkerk H P, Bundy F P, Hall H T, Strong H M and Wentorf R H 1959 Preparation of diamond Nature 184 1094
[4] Bundy F P 1989 Pressure-temperature phase diagram of elemental carbon Physica A 156 169
[5] Grochala W 2014 Diamond: electronic ground state of carbon at temperatures approaching 0 K Angew. Chem., Int. Ed. 53 3680
[6] Popov I V, Görnè A L, Thoureen C L and Dronskowski R 2019 Relative stability of diamond and graphite as seen through bonds and hybridizations Phys. Chem. Chem. Phys. 21 10961
[7] Howes V R 1962 The graphitization of diamond Proc. Phys. Soc. 80 648
[8] Binnig G, Quate C F and Gerber C 1986 Atomic force microscope Phys. Rev. Lett. 56 930
