Elastic coupling between layers in two-dimensional materials

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large scientific and technological effort is underway to understand and control the properties of 2D materials because of their potential technological applications¹⁻⁷. The most studied 2D material is graphene, existing as a single laver of graphite⁸ or a few-layer-thick epitaxial graphene (EG) film⁹. Graphene possesses a large in-plane Young's modulus¹⁰ (~1 TPa) as well as high intrinsic carrier mobility^{11,12} and high in-plane thermal conductivity¹³. Besides graphene, also 2D films of graphene oxide¹⁴⁻¹⁷ (GO), hexagonal boron nitride¹⁸ and transition metal dichalcogenides¹⁹⁻²¹ such as MoS₂ exhibit unique and excellent properties and hold great promise for nanotechnology applications. One of the main characteristics of 2D materials is the high anisotropy between the in-plane and perpendicular-to-the-plane properties. For example, in graphite, owing to the strong covalent bonds between atoms in the plane, and the weak Van der Waals interlayer interaction, the in-plane Young's modulus is $E_{\parallel} = 1$ TPa (ref. 22), whereas the interlayer perpendicular-to-the-plane Young's modulus is only $E_{\perp} = 36$ GPa (ref. 22). Recent studies have suggested that the mechanical properties of 2D materials are strongly correlated to their structure and properties^{1,10,23-25}. The in-plane Young's modulus^{1,10,25-27} of exfoliated graphene and MoS₂ has been widely studied in bending experiments where a film is suspended on trenches or holes, and an atomic force microscopy (AFM) tip is used to bend the suspended film with deformations of tens and hundreds of nanometres. On the other hand, very little is known about the elasticity perpendicular to the planes, hereafter called perpendicular or interlayer elasticity, of 2D materials composed of very few atomic layers. Recent calculations have investigated the out-of-plane shear and Young's modulus of carbon nanotubes and graphene²⁸. Experimentally, resonance ultrasound spectroscopy was used to study the elastic constants and the anisotropy between the in-plane and perpendicular-to-the-plane directions of thin films²⁹. Investigations of the perpendicular-to-the-plane elasticity of fewlayer-thick 2D films have not been reported to the best of our knowledge, and remain an experimental challenge because they require indentations on supported—as opposed to suspended—2D films, where the indentation should remain smaller than the films' interlayer distance, that is, less than a few ångströms. Nevertheless, the interlayer elastic coupling is particularly important because it is related to the thermal³⁰, electronic³¹, tribological^{24,32} and optical³³ properties of 2D films. The perpendicular elasticity is expected to be affected by the structure and chemistry of the layers, the presence of stacking and intrinsic defects, and intercalation, which is a critical process for doping and tuning mechanical and electronic properties in 2D films. Mapping the interlayer elastic coupling in 2D films is therefore an important technological and scientific advancement.

Modulated nanoindentation experiments

Here, an unconventional AFM-based method allowing for subångström-resolution indentations coupled with a semi-analytical method (SAM) is used to measure the elasticity of 2D films in the direction perpendicular to the layers, for a number of layers as small as two. This AFM-based method is capable of maintaining an indentation depth smaller than the interlayer distance, with a resolution of 0.1 Å. We report on AFM–SAM investigations of the perpendicular elastic modulus of EG, epitaxial graphene oxide (EGO), and conventional GO films at varying ambient humidity. These studies, together with density functional theory (DFT) calculations, highlight how the interlayer elastic coupling in graphene and GO films is affected by the films' chemistry, structure, water intercalation, and number of layers. Interestingly, the perpendicular Young's modulus of GO increases by increasing

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Figure 1 | Modulated nanoindentation experiments. **a**, Schematic diagram of the experimental set-up, where a spherical AFM tip vibrates while indenting a few-layer-thick film of graphene or GO. **b**, Experimentally measured indentation curves for single-crystal SiC, 10-layer-thick EG, and 10-layer-thick EGO. All three curves were obtained with the same AFM tip, R=114 nm.

the amount of water trapped in between the layers until a full monolayer is produced. Then, when the second water layer is forming, the modulus decreases. Finally, our studies show that when using sub-interlayer-distance indentations in 2D films thicker than a few layers, the AFM indentation curves are very sensitive to the elastic modulus perpendicular to the layers, E_{\perp} , and almost independent of the value of the in-plane Young's modulus, E_{\parallel} (see Fig. 1a). As a consequence, the indentation force curves can be fitted with a simple modified Hertz model as if the film is an isotropic material, where Young's modulus of the film is indeed the perpendicular modulus E_{\perp} . This work is therefore offering a new experimental and theoretical framework to investigate the interlayer elastic coupling in 2D films.

The experimental measurement of the perpendicular-to-theplane elasticity of 2D films is here achieved by using an unconventional AFM-based method that we call modulated nanoindentation^{14,34,35} (MoNI). In particular, we have investigated highly oriented pyrolytic graphite (HOPG), 4H-SiC, EG films grown on a SiC substrate^{2,9}, EGO films grown on a SiC substrate³⁶, and regular GO films deposited with the conventional Hummers' method37 on a Si substrate. More details about indentation experiments, sample preparation and properties of the films are provided in Methods and Supplementary Information (see also Supplementary Figs 1-5). During a typical MoNI experiment, an AFM tip, which is vertically oscillated at a fixed frequency with a ~0.1 Å amplitude, applies an increasing pressure to a 2D film surface, in the z-direction perpendicular to the film surface (see Fig. 1a). The oscillations are applied and controlled by a lockin amplifier, while a constant normal force F_z between the tip and the 2D film is maintained constant by the feedback loop of the AFM. By working with a constant force, any thermal drift is avoided. Furthermore, using a lock-in detection system together with a differential measurement allows us to measure very shallow indentations, usually smaller than 1-3 Å, with a resolution of 0.1 Å. Indeed, instead of measuring directly the normal force as a function of the indentation, z_{indent} , we measure the slope of the force versus indentation curve at each constant normal force, namely $k_{\text{cont}}(F_z)$. Force versus indentation curves are then obtained by integrating the equation $dF_z = k_{cont}(F_z) \cdot dz_{indent}$ as follows:

$$z_{\text{indent}}(F_z) = \int_0^{F_z} \frac{\mathrm{d}F_z}{k_{\text{cont}}(F_z)} \tag{1}$$

This experimental set-up is therefore uniquely suited to investigate the perpendicular elastic modulus of 2D films, which are only a few layers thick. We note that the lower limit of the integral in equation (1) is zero only when there are no adhesion forces. In the case of an adhesive contact, the lower limit is the pull-off force, F_{po} , defined as the negative load at which the AFM tip loses the contact with the sample and can be easily determined experimentally (Supplementary Figs 2 and 3 for more details)³⁸⁻⁴⁰. Furthermore, for SiC and the 2D materials investigated here, when the load is equal to F_{po} the contact area is indeed zero and z_{indent} (F_{po}) = 0 (see Supplementary Information). As shown in Supplementary Fig. 3, $k_{cont}(F_z)$ —which is proportional to the contact radius and the squared root of the indentation depth-drops to zero when the load reaches F_{po} . We underline that the F_z versus z_{indent} curves could in principle be shifted along the z_{indent} axis in very soft and adhesive materials depending on the value of $z_{indent}(F_{po})$; however, the shape of these curves does not depend on the type of adhesive contact. We also note that in Supplementary Figs 2 and 3 the pulloff force can change significantly from curve to curve; however, ΔF_{z} remains consistent. This shift in the absolute value of the normal force is due to humidity and the drifting of the laser position on the photo-detector during the different individual experiments (we take hundreds of curves for each sample and condition). More detailed discussions can be found in the Supplementary Information and Supplementary Figs 4 and 5. However, the clear determination of the pull-off force during each indentation experiment allows one to always determine the corrected 'absolute' normal load as $F_z = (F_z - F_{po})$, and to compare different curves independently of the laser position shifting and presence of adhesion forces. In the Supplementary Information we discuss in detail why for the materials investigated here we can account for the adhesion force by simply adding to the load the pull-off force. This adhesive contact mechanics model is called the Derjaguin-Muller-Toporov (DMT) model^{40,41} and it describes well stiff materials and small tip radii. Figure 1b shows the measured (and offset-corrected) F_z versus z_{indent} at ambient humidity for different 2D films, namely EG and EGO. We remark that both EG and EGO in Fig. 1b are 10 layers thick. Figure 1b also shows the force versus indentation curve for a single crystal of silicon carbide, the substrate over which EG and EGO are grown. The indentation curves, all performed with the same AFM tip, clearly indicate a larger stiffness for 10-layer EG than 10layer EGO. As expected from previous studies, SiC shows a very large stiffness. 4H-SiC has a hexagonal crystal structure; however, its elastic behaviour is very isotropic, and therefore a simple Hertzian offset-corrected contact model can be used to extract from the force versus indentation curves Young's modulus of the material. For the configuration of a sphere (the AFM tip) pressing on a flat surface of an isotropic half-space with a normal force F_z the Hertz model gives:

$$F_z = \frac{4}{3} E^*(R)^{1/2} z_{\text{indent}}^{3/2}$$
(2)

where $E^* = ((1 - (v_{\text{Hertz}}^{\text{sample}})^2 / E_{\text{Hertz}}^{\text{sample}}) + (1 - (v_{\text{Hertz}}^{\text{tip}})^2 / E_{\text{Hertz}}^{\text{tip}}))$, with $v_{\text{Hertz}}^{\text{sample,tip}}$ and $E_{\text{Hertz}}^{\text{sample,tip}}$ being Poisson's ratio and Young's moduli of, respectively, the investigated sample and the AFM tip, with R the AFM-tip radius⁴². For the silicon tip used in the MoNI measurements, $E_{\text{Hertz}}^{\text{tip}} = 169 \text{ GPa}$ and $v_{\text{Hertz}}^{\text{tip}} = 0.27$. The elastic modulus can therefore be obtained by using the relationship (2) to fit the experimental measurements of F_z versus z_{indent} , as shown in Fig. 1b for a quasi-isotropic sample such as SiC. Indeed, the fitting procedure gives $E_{\text{Hertz}}^{\text{SiC}} = 400 \text{ GPa}$, for a tip radius R = 114 nm(the details of tip radius measurement can be found in the Supplementary Information and Supplementary Fig. 6), which is in excellent agreement with the literature. Similar experiments have also been performed on other standard samples such as ZnO single crystals to ensure the ability of MoNI to obtain reliable results (see details in the Supplementary Information and Supplementary Fig. 2). However, the Hertz relationship in (2) is in principle not well suited to model the contact mechanics between an AFM spherical tip and an anisotropic material such as graphite and other 2D materials because the Hertz model was originally valid only for isotropic half-spaces^{42,43}.



Figure 2 | **Experimental, SAM-simulated and Hertz indentation curves. a**, Experimentally measured indentation curves in HOPG (filled circles), semi-analytical model simulations of indentation in graphite (open circles), and Hertzian fitting (continuum line) of the indentation curves on HOPG. The indenting tip radius was 100 nm. b, Contact-pressure distribution profiles for Hertz contacts and SAM simulations of indentation in graphite. Note that for bulk graphite and for a graphite film 50 nm thick, the SAM simulations and the contact distribution profiles almost overlap. c, Experimental indentation curves on 10-layer-thick EG, 1-layer-thick EG, buffer-layer EG, and SiC. **d**, Statistical analysis of exponent number *b* in the fitting function $F_z = C^* z_{indent}^b$. For EGO and GO, the RH is indicated. $b_{average}$ is 1.40.

Semi-analytical methods

A better insight into the contact-pressure distribution as a function of the material elastic constants can be obtained using simulations with SAMs, which have proved their efficiency in describing the contact mechanics of anisotropic materials⁴⁴⁻⁴⁶. Here, we use SAM to simulate the force versus indentation curves in graphite. We use graphite elastic constants found in the literature²² (see Methods), and we model the indentation of an AFM silicon tip (R = 100 nm)in a graphite sample deposited on SiC. We use this configuration because the 2D materials studied here have been deposited either on SiC (EG and EGO) or Si (conventional GO). Figure 2a shows the results from the SAM simulations on graphite along with the experimental curves obtained by MoNI on a bulk sample of HOPG. In Fig. 2a, the SAM-simulated curve agrees extremely well with the experiments on HOPG; it is important to note that in the SAM simulations for graphite we use, according to the literature, as in-plane Young's modulus $E_{\parallel} = 1.046$ TPa, and as *z*-axis (perpendicular to the planes) Young's modulus $E_{\perp} = (36.4 \pm 1)$ GPa (more details are given in the Methods). Interestingly, when the Hertz model in (2) is applied to fit the experimental indentation curves measured on graphite, as if graphite was an isotropic material, the result of the fitting procedure gives as the single isotropic modulus $E_{\text{Hertz}}^{\text{HOPG}} = (33 \pm 3)$ GPa, for R = 100 nm like the AFM tip radius. The Hertz model fitting curve is also reported in Fig. 2a to show the perfect agreement with experiments and SAM simulations. The Hertz model is therefore able with a simple fitting procedure to obtain a value of Young's modulus that is equal, within an error of 10%, to the most accepted value for the perpendicular-to-the-plane Young's modulus of graphite, that is, $E_{\perp} = 36$ GPa.

The excellent consistency between experiment, simulation and Hertz model when studying the indentation of 2D films with extremely small indentation depths is a direct consequence of the following SAM observations. If we consider sub-interlayer distance indentations in a transversally isotropic (orthotropic) material having $E_{\perp} = E_{\text{Hertz}}$ and E_{\parallel} varying up to one order of magnitude compared with E_{Hertz} , we find that the contact pressure and contact area for a given pressure remain almost the same as in an isotropic material having $E = E_{\text{Hertz}}$. On the other hand, the contact pressure changes markedly compared with the isotropic case when varying E_{\perp} in the same range and maintaining $E_{\parallel} = E_{\text{Hertz}}$ (ref. 44). Overall these results indicate that for sub-nanometre AFM indentations, much smaller than the film's thickness, the force versus indentation curves are very sensitive to E_{\perp} , and almost independent of the value of E_{\parallel} . This result can be understood with a back-of-theenvelope calculation of the in-plane and perpendicular-to-theplane stress distribution in a layered material when indenting the material perpendicular to the planes (for details see Supplementary Information and Supplementary Fig. 8). From these calculations it seems clear that the key parameter controlling sub-nanometre indentations is the ratio $a^2/l_0 z_{\text{indent}}$, where a is the contact radius, and l_0 is the length of an sp^2 bond. For example, when indenting graphite with an AFM tip with $z_{\text{indent}} < 0.3 \text{ nm}$, this simple model shows that we are sensing only E_{\perp} . A more precise analysis can be developed by using SAM calculations. In particular, the contact pressure distribution profiles in the case of sub-nanometre indentations for the Hertz model in the case of an isotropic material with $E_{\text{Hertz}} = 33 \text{ GPa}$ (the value of E_{\perp} in graphite), and in the case of $E_{\text{Hertz}} = 1.046$ TPa (the value of E_{\parallel} in graphite), are plotted in Fig. 2b along with the SAM simulations for bulk graphite and

Table 1 | Summary of the experimental results of E_{\perp} (GPa) at different RHs.

Relative humidity	10 ± 2%	15 ± 3%	25±3%	35±3%	50 ± 3%
10-layer EGO	22 ± 3	-	23 ± 4	19 ± 3	22 ± 3
Conventional GO	21 ± 6	26 ± 6	35 ± 10	-	23 ± 7
10-layer EG	-	-	36 ± 3	-	-
HOPG	-	-	33 ± 3	-	-

50-nm-thick graphite. It can be easily concluded that the pressure distributions for an isotropic material having E = 33 GPa, bulk graphite, and 50-nm-thick graphite are almost all the same. On the other hand, the pressure distribution changes markedly when considering an isotropic material with E = 1.046 TPa. We conclude that when studying 2D materials with sub-nanometre indentations, the Hertz model is an extremely simple and accurate model to fit the experimental AFM indentation curves and obtain the perpendicular Young's modulus E_{\perp} of 2D films having a thickness larger than the indentation depth.

Perpendicular-to-the-plane elasticity

After having tested and gathered a better understanding of subnanometre indentations in orthotropic layered materials such as bulk graphite, we have then used MoNI to investigate the interlayer elasticity of very thin films of layered materials. Initially, MoNI is performed on a film of 10-layer-thick EG grown on SiC, as shown in Fig. 2c. We remark that 0.1 nm indentations in supported graphene films with a thickness of ~ 10 layers are not very sensitive to the presence of the SiC substrate, as also shown by SAM calculations. Similarly to the fitting procedure used in Fig. 2a, we use the Hertz model to fit the indentations curves on 10-layer EG. We have repeated several measurements, and the Hertz fit provides for 10-layer EG a modulus perpendicular to the planes equal to $E_{\perp} = (36 \pm 3)$ GPa, the same value as ideal graphite, and larger than the value measured on HOPG ($E_{\perp} = 33 \pm 3$ GPa). It is worth noting that in Fig. 2a for the SAM simulations and the MoNI experiments on HOPG we use a tip radius of $R = (100 \pm 10)$ nm, whereas the MoNI experiment on 10-layer EG in Fig. 2c was performed with an AFM tip of $R = (114 \pm 10)$ nm; therefore, to compare these curves we need to rescale them by a factor of $R^{1/2}$ (see equation (2)).

To investigate the role of the substrate and number of layers, we have then focused our experiments on the first layers of EG grown on the Si face of SiC (0001). More details on these experiments are reported in the Methods, in the Supplementary Information and in Supplementary Fig. 7. Much attention in the scientific community is at present directed towards single-layer EG grown in the Si face of SiC (refs 31,47). This system is very interesting because between the SiC substrate and the first 'well-decoupled' 1-layer graphene, there is a honeycomb carbon layer called the buffer layer³¹ that has different properties from graphene, and it has a strong interaction with the SiC substrate. This substrate interaction strongly modifies the electronic structure of the carbon buffer layer, which is in fact not conducting. Extensive research is underway about the properties of the decoupled first 1-layer graphene and the buffer layer on SiC. Here, we have performed MoNI experiments on the decoupled first 1-layer graphene and on the buffer layer grown on the Si face of SiC, as shown in Fig. 2c and Supplementary Fig. 7. We find that the buffer layer produces indentation curves very similar to the ones on bare SiC (also reported in Fig. 2c). This result is in agreement with the notion that C atoms in the buffer layer have a strong interaction with the SiC substrate; indeed this interaction is often described as covalent bonding³¹. On the other hand, we obtain stiffer indentation curves on the decoupled 1-layer graphene on SiC. These curves are

stiffer than those ones measured on the buffer layer, and on 10layer EG, as clear in Fig. 2c and Supplementary Fig. 7. This result can be understood by the following considerations. For a film of 10layer graphene, we are probing only E_{\perp} , which is ~36 GPa, because the 0.1 nm indentation is easily distributed among the 0.01 nm displacements per layer, supposing an infinitely rigid substrate. In contrast, when the AFM tip is indenting $(z_{indent} \sim 0.1 \text{ nm})$ 1-layer EG on an infinitely rigid substrate, the whole indentation is carried on by that single layer sitting at an interatomic distance of 0.3 nm from the substrate, and strongly resisting such a huge deformation. Therefore, for 1-layer graphene on SiC we are now probing a complex convolution of effects and properties, which include E_{\parallel} (~1 TPa), E_{\perp} , and the elasticity of the SiC substrate ($E \sim 400$ GPa). In conclusion, these experiments show that MoNI measurements can be sensitive to the substrate interaction and to the number of layers, although more calculations will be required to fully explain the results for 1-layer graphene.

The ability of the Hertz relationship in equation (2) to model sub-nanometre indentations in transversally isotropic 2D materials is further confirmed by Fig. 2d. We have fitted all (except for 1-layer EG) the measured $F_z(z_{indent})$ curves with Hertz equation (2) leaving as a free fitting parameter the exponent, b, of z_{indent} , that is, by using the function $F_z = C \cdot z_{indent}^b$, where C is a constant. The results are reported in Fig. 2d, and they show that b is on average equal to (1.4 ± 0.1) , a value in very close agreement with the exponent expected for a perfect Hertzian contact where b = 1.5 (see equation (2)). By fitting with the Hertz equation (2) all of the experimental indentation curves for the different 2D materials films in ambient atmosphere (see Table 1), we find that as mentioned above the perpendicular elastic modulus is larger in 10-layer EG films than in HOPG; this difference is probably related to the absence of defects in 10-layer EG compared with HOPG. We also find that 10-layer EGO is much softer than 10-layer EG; this result can be explained by the increase of the interlayer distance d, in EGO compared with EG, precisely from d = 3.4 Å to d = 9.3 Å (ref. 48) as also observed in X-ray diffraction spectroscopy measurements⁴⁹. The more surprising result is the difference in the perpendicular elasticity between EGO films and GO films produced with the conventional exfoliation/filtration/deposition method (see Methods). In ambient humidity we precisely obtain: $E_{\perp} = (35 \pm 10)$ GPa (for conventional GO), $E_{\perp} = (36 \pm 3)$ GPa (for 10-layer EG), $E_{\perp} = (33 \pm 3)$ GPa (for HOPG), and $E_{\perp} = (23 \pm 4)$ GPa (for EGO). To understand the origin of the large interlayer perpendicular elasticity in conventional GO compared with EGO, we have performed DFT calculations on the elasticity of GO with a different amount of intercalated water. We have then compared these calculations with MoNI experiments on GO and EGO performed at different ambient humidities.

Density functional theory study of perpendicular elasticity

DFT calculations have been performed on model structures of GO consisting of periodic stacks of graphene layers fully oxidized by either hydroxyl or epoxide groups, including increasing concentrations of water molecules, and presenting AA stacking (see Supplementary Information and Table 1). For each model, we used a DFT-D2 (refs 49,50) scheme (technical details and the full list of calculations are discussed in Supplementary Information, and Supplementary Figs 9-12) to perform a full structural optimization and determine the zero-temperature interlayer spacing. Subsequently, we applied a pressure and used DFT-D2 to estimate the z-axis Young's modulus from the energy versus displacement curves. A summary of the DFT results reporting E_{\perp} and interlayer distance as a function of intercalated water percentage for GO structures fully oxidized with hydroxyls is reported in Table 2. Figure 3a shows the calculated F_z versus displacement curves at varying intercalated water content for the case of graphene fully oxidized with hydroxyl groups, because



Figure 3 | **DFT** and **experimental results for conventional GO films. a**, The DFT-calculated F_z versus displacement curves for different water contents in graphene fully oxidized with hydroxyl groups. **b**, Experimental F_z versus indentation depth curves at different RHs in conventional GO. All of the curves were obtained with the same AFM tip. **c**,**d**, Experimental and DFT results of E_{\perp} of GO as a function of water content and RH, respectively. The insets are schematic diagrams of the corresponding atomistic structures showing how water molecules fill the interlayer spacing. Each experimental point of E_{\perp} is an average value of more than 30 different measurements.

previous experiments have shown that conventional GO films are mainly composed of hydroxyls groups³⁶. In comparison, Fig. 3b presents the experimental MoNI results on conventional GO at different relative humidities (RHs). Figure 3c,d shows the resulting E_{\perp} as a function of intercalated water percentage compared with carbon (for DFT calculations) and RH (for MoNI experiments), respectively. A summary of the experimental results is also reported in Table 1. The agreement between experiments and DFT calculations is striking. DFT calculations and experiments show that E_{\perp} increases with the amount of intercalated H₂O molecules (and RH), reaching a maximum value of about 31 GPa at 25% H₂O, and 35 GPa at RH = 25%, for the DFT calculations and indentation experiments, respectively. The perpendicular elastic modulus then decreases down to 20 GPa at 50% water, and 23 GPa at 50% RH, for the DFT calculations and indentation experiments, respectively. We remark that the presence of a maximum in the perpendicular modulus found in both experiments and calculations is a key result; however, the excellent quantitative agreement between water content (from DFT) and RH (from experiments) might be only a coincidence. The DFT calculations clearly give an insight into the atomistic origin of the behaviour of E_{\perp} as a function of intercalated water. DFT calculations show that the interlayer distance and perpendicular Young's modulus in GO change abruptly between the case of dry layers and a multilayer film including a small amount of H_2O (<6%), whereas more gradual variations of these physical properties are obtained for increasing the water content between 6.25% and 25% (Fig. 3c, and Table 2). In particular, E_{\perp} drops from about 35 GPa (close to EG) in dry films to about 11-14 GPa when the layered structure includes 6.25% H₂O. This behaviour can be understood by considering that when the amount of water is only a few percentage of the carbon amount, H₂O molecules swell the graphene structure, increasing the interlayer distance from 3.4 Å to about 6.2 Å, but leaving the interlayer space mainly empty and therefore producing a soft structure with a low perpendicular elastic modulus. This interlayer modulus increases with increasing amount of water, which fills the interlayer space without changing too much the interlayer distance (Table 2). However, at 25% water, H₂O molecules have completely filled a water layer in between the layers, and this situation corresponds with the maximum in perpendicular elastic modulus. Above 25% water, the perpendicular elastic modulus decreases because a second water layer starts to form in between the layers, further swelling and softening the GO structure. Owing to this explanation, it is now possible to understand the different values of E_{\perp} in EGO compared with conventional GO. EGO is not a porous structure, and water intercalation is minimal⁴⁹ and independent of humidity. For this reason we find that E_{\perp} in EGO remains constant and \sim 22 GPa for all RHs. On the other hand, conventional GO is a porous structure³ where the amount of intercalated water can change depending on the humidity; therefore, in agreement with the DFT calculations we observe a maximum in conventional GO when varying the RH.

Outlook

In conclusion, we have presented a new methodology, which combines sub-ångström-resolution indentation measurements and SAMs to study the elasticity perpendicular to the plane of few-layerthick 2D materials. The comparison between indentation experiments and SAMs has also demonstrated that a simpler approach to interpret experimental sub-nanometre indentation curves in fewlayer-thick 2D films is to use the Hertz model. We showed that the fitting of the experimental indentation curves with the Hertz model provides, to a good approximation, the value of Young's modulus **Table 2** | Summary of the DFT results reporting interlayer distance and E_{\perp} as a function of different fractions of intercalated water for GO structures fully oxidized with hydroxyls.

Water fraction (%)	Interlayer distance (Å)	E⊥ (GPa)
0	6.2	34.9
6.25	8.6	13.4
12.5	8.7	18.2
25	8.9	31.3
50	11.3	20.7

perpendicular to the plane of 2D films. The experimental study of EG and different types of GO film, combined with DFT calculations, has demonstrated that the interlayer elasticity is extremely sensitive to the presence of intercalated molecules in between the planes. In particular, these studies show that intercalated water in GO can markedly change the interlayer elastic modulus, which at first decreases when a small amount of water is intercalated in between the layers and the structure is swelled, and then increases with increasing amount of water until H₂O molecules have completely filled a water layer in between the layers. Above this point, the perpendicular elastic modulus decreases because a second water layer starts to form in between the layers, further swelling and softening the GO structure. This understanding can also explain the different behaviour of E_{\perp} in conventional GO and EGO. Finally, the here proposed sub-ångström-resolution indentation measurements applied to EG are shown to be sensitive to the substrate interaction and to the number of layers.

The results reported here provide a new path to study the interlayer elastic coupling and the Van der Waals forces in few-layerthick 2D materials, and shed new light on the use of the Hertz model in investigating the perpendicular-to-the-plane Young's modulus of 2D films. This study will impact a variety of fields, from electronics to phononics, allowing new investigations and understanding of the relationship between molecular structure, thermal conductivity, electronic properties, and phonon propagation in layered materials. For example, through local measurements of the elastic modulus, MoNI could probe interlayer and substrate interaction, as well as the presence of dopants/intercalates, which are extremely important for modulating the electronic properties of 2D materials. Furthermore, the interlayer elasticity is strictly connected with the out-ofplane thermal properties of layered materials. The here discussed interlayer elasticity measurements could help in the understanding of the origin of the extremely low out-of-plane thermal conductivity found in thin films of transition metal dichalcogenides³⁰.

Methods

Methods and any associated references are available in the online version of the paper.

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Author contributions

Y.G., S.K. and H-C.C. performed nanomechanics experiments and data analysis. S.Z. carried out DFT calculations. D.N. performed the SAM calculations. C.B., and W.d.H. synthesized the EG and EGO samples. L.P. and R.S. synthesized the GO samples. A.B. conceived and designed the theory and analysed the data. E.R. conceived and designed the experiments and analysed the data. All authors contributed to write the article.

Additional information

Supplementary information is available in the online version of the paper. Correspondence and requests for materials should be addressed to A.B. or E.R.

Competing financial interests

The authors declare no competing financial interests.

Methods

Modulated nanoindentation measurements. The oscillations are applied at 1 kHz to the atomic force microscopy (AFM) tip by a piezoelectric stage rigidly attached to the AFM cantilever-tip system, and controlled by a lock-in amplifier (Stanford Research Systems, SR830), while a constant normal force F_z between the tip and the 2D films is maintained constant by the feedback loop of the AFM (see Supplementary Information for more details). To maintain the linear elastic regime, the piezo-stage oscillations are chosen to be only ~ 0.1 Å. During the indentation, the driving fixed piezo-stage oscillation amplitude Δz_{piezo} is equal to the sum of the cantilever bending and tip-2D film normal deformation. Under such circumstances, the AFM cantilever and the tip-film contact can be considered as two springs connected in series: the cantilever with stiffness k_{lev} and the tip-film contact with stiffness k_{cont} . The force required to stretch these two springs in series with a total displacement Δz_{piezo} is equal to the normal force variation ΔF_z . This experimental configuration allows us to measure the total stiffness k_{tot} at each normal load F_z , fixed by the feedback loop of the AFM, from the following relation: $(\Delta F_z / \Delta z_{\text{piezo}}) = k_{\text{tot}}(F_z)((1/k_{\text{lev}}) + (1/k_{\text{cont}}))^{-1}$ where ΔF_z is the variation of the normal force caused by the piezo-stage oscillation and is much smaller than F_z , k_{lev} is the spring constant of the cantilever, and k_{cont} is the tip-film contact stiffness. As $k_{\rm lev}$ can be measured independently, the measurement of $\Delta F_z / \Delta z_{\rm piezo}$ at different normal loads F_z allows us to acquire the tip-film contact stiffness k_{cont} as a function of F_z . Force versus indentation curves are obtained by integrating $dF_z = k_{cont}(F_z) \cdot dz_{indent}$. The data reported in Table 1 are averaged values obtained from 10, 210, 90 and 12 measurements on highly oriented pyrolytic graphite, graphene oxide (GO), epitaxial graphene oxide (EGO) and epitaxial graphene (EG) respectively. More details on the MoNI experiments are reported in the Supplementary Information.

2D materials. We have studied 10-layer graphene grown epitaxially on the C face of a 4H-SiC substrate². We have also investigated the buffer layer and the first decoupled 1-layer graphene, both grown epitaxially on the Si face of 4H-SiC (0001) (refs 31,47; see Supplementary Fig. 7).

To prepare EGO samples, the 10-layer EG samples on SiC chips are then directly oxidized shortly after preparation by using a milder Hummers' method⁴⁹, which avoids graphene exfoliation and dispersion in solution. Once the reactions are terminated, EGO films on the SiC chips are picked up from the solution and rinsed with deionized water for 1 min. The EGO films, 10 layers thick, are finally blow-dried by nitrogen gas.

GO films are prepared by drop-casting colloidal GO dispersion on a Si chip and leaving it to dry at 80 °C. Stable colloidal GO dispersions are produced by a modified Hummers' method³⁷. Once the reaction is terminated, the oxidized portion is separated and cleaned from unoxidized graphite and other residual species. The cleaning is performed by centrifugation of the obtained suspension at 7,197g for 30 min to remove both the acidic content and ions. The solid content is collected and redispersed with deionized H₂O. This operation is repeated in sequence until the pH of the supernatant is close to neutrality. At that point, the exfoliation of graphite oxide is performed by prolonged and vigorous shaking, forming a brownish colloidal suspension of GO flakes. The subsequent collection of the purified supernatant results in stable aqueous GO suspensions, which are drop-casted on Si to form a film with a thickness of 40–50 nm (ref. 51).

Semi-analytical method. The theoretical model used here to predict the indentation curves as well as the distribution of contact pressure and the contact area is based on what is known as semi-analytical methods. SAM is an alternative to the finite element method, quite close to the boundary element method, suitable for continuum mechanics problems. The main advantage of the boundary element method over SAM is that the first one is more versatile because it uses surface or volume integrals whatever the shape of the volume of interest. Conversely, SAM uses analytical solutions of these integrals for simple geometries (such as sphere or cuboid), which reduces the computation to a simple summation of elementary solutions. SAM consists of the numerical summation of elementary analytical solutions, such as the effect of a point load on a layered half-space, as far as they are known. The difficulty for anisotropic elastic coating and/or substrate is that the set of equations to be solved requires manipulating complex numbers with conjugate pairs, leading to double roots when the material tends to behave as elastically

isotropic. One of the advantages of SAMs when applied to the contact problem compared with more widely used numerical techniques is the computing time, which is at least one or two orders of magnitude shorter. For more details on SAM for the numerical procedure used to solve the contact problem for an elastic substrate coated with an anisotropic layer, see refs 44,45.

For transversely isotropic material, there are 5 independent elastic constants in the elasticity stiffness tensor: C_{11} , C_{12} , C_{13} , C_{33} and C_{44} . These 5 elastic constants can be transformed to the well-known engineering notation in the following way.

$$E_{\parallel} = E_x = E_y = (C_{11} - C_{12})(C_{11}C_{33} + C_{12}C_{33} - 2C_{13}C_{13})/$$

$$(C_{11}C_{33} - C_{13}C_{13})$$

$$E_{\perp} = E_z = C_{33} - 2C_{13}C_{13}/(C_{11} + C_{12})$$

$$v_{zx} = C_{13}/(C_{11} + C_{12})$$

$$G_{xz} = C_{44}$$

$$G_{xy} = (C_{11} - C_{12})/2 \quad v_{yy} = v_{yy} = (E_x/G_{yy}) - 1$$

Note that G_{xy} and v_{xy} are correlated, which means that only one of them is independent. Graphite's elastic constants used for SAM have been previously reported in the literature²² and they are $C_{11} = (1,060 \pm 20)$ GPa, $C_{12} = (180 \pm 20)$ GPa, $C_{13} = (15 \pm 5)$ GPa, $C_{33} = (36.5 \pm 1)$ GPa and $C_{44} = (4.5 \pm 0.5)$ GPa. For graphite, this gives $E_{\parallel} = (1.06 \pm 0.02)$ TPa, and $E_{\perp} = (36.4 \pm 1)$ GPa.

The SiC substrate was approximated as cubic with $E\,{=}\,450$ GPa, $\nu\,{=}\,0.17$ and $G\,{=}\,E/2(1+\nu)\,{=}\,192.31$ GPa.

DFT calculation. Density functional theory (DFT) calculations are performed by using the PWscf code of the QUANTUM Espresso toolkit⁵⁰. We use a plane-wave basis set with an energy cutoff of 120 Ry to represent the Kohn-Sham wavefunction, norm-conserving pseudo-potentials for all atomic species⁵², and the exchange-correlation functional proposed by Perdew, Burke and Ernzerhof⁵³. To model hydrated GO, we consider two ordered structures-graphene sheets fully covered by either hydroxyl or epoxide groups, with various amounts of water molecules intercalated in between the oxide layers^{36,48,49}. The GO sheets are arranged in either 'AA' or 'AB' stacking geometry. We use the primitive unit cell of each model as the supercell. We use $3 \times 3 \times 3$ and $3 \times 3 \times 1$ gamma-centred Monkhorst-Pack meshes to sample the Brillouin zone of the supercells including one and two oxide layers, respectively. To account for the London dispersion forces in multilayer GO, we adopt the semi-empirical DFT-D2 approach proposed previously⁵⁴. The GO models are fully optimized for both the electronic and cell degrees of freedom using the DFT-D2 scheme. To calculate the perpendicular-to-the-plane Young's modulus, we started from the equilibrium structure of each model, and we vary the lattice parameter along the z direction with a strain step of \sim 0.2% up to a 2% strain. The ionic positions of GO models are then gently relaxed to obtain the total energy and stress tensor under each strain. The E_{\perp} value of each model is calculated by interpolating the pressure along the z direction with respect to the volume using the equation: $E_{\perp} = -V dP/dV$, where P is the pressure along the z direction, and V is the volume of GO models. More details are reported in the Supplementary Information.

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