Second layer of H$_2$ and D$_2$ adsorbed on graphene

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We report diffusion Monte Carlo calculations on the phase diagrams of para-H$_2$ and ortho-D$_2$ adsorbed on top of a first layer of the same substances on graphene. We found that the ground state of the second layer is a triangular incommensurate solid for both isotopes. The densities for promotion to a second layer and for the onset of a two-dimensional solid on that second layer compare favorably to available experimental data in both cases.

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I. INTRODUCTION

Graphite and graphene are closely related forms of carbon in which the atoms are located in the nodes of a honeycomb lattice. The main difference is that graphite is formed by a whole stack of the two-dimensional carbon sheets that constitute a single graphene layer. Adsorption of gases on top of graphite and graphene are expected to show similar trends. This is in fact what has been shown by computer simulations of the phase diagrams of adsorbed quantum gases on graphite, both in the first and second layers. In general, those calculations render phase diagrams which are very close to the experimental ones on graphite. The only appreciable difference is the binding energy of the adsorbate species on top of the carbon structure, bigger in the graphite case. Unfortunately, to our knowledge, there are no experimental data yet on adsorption of quantum gases on graphene to be compared to.

Most of that work, both from the experimental and theoretical sides, has been devoted to the determination of the phase diagram of the first layer of quantum gases and solids adsorbed on top of graphite. However, there has been some measurements of the properties of a second He sheet on top of an incommensurate helium layer, directly in contact with a graphite surface, work that can be compared to the simulations on the same subject. On the other hand, the second layer of molecular hydrogen on top of graphite has been less explored. Experimentally, the second layer of molecular hydrogen and deuterium adsorbed both on graphite and MgO has been studied using calorimetric measurements. In particular, Ref. studies the promotion to a second layer and the phase diagram of pure para-H$_2$ (p-H$_2$) and ortho-D$_2$ (o-D$_2$) second layers on graphite. One of the main conclusions of that work is that the ground state of both isotopes in the second layer is a quasi two-dimensional solid. Those calorimetric measurements suggest triple points at T $\sim$ 6 K for H$_2$ and T $\sim$ 11 K for D$_2$. For lower temperatures and densities those solids seem to coexist with infinitely diluted gases. The main goal of our present work is to perform diffusion Monte Carlo calculations to determine the phase diagram of the second layer of H$_2$ adsorbed on a first layer of H$_2$ on graphene, and of the second layer of D$_2$ adsorbed on the first layer of D$_2$, also on graphene. The results so obtained will be compared to the experimental ones for the same systems on graphite the only ones available, in order to assess the possible differences. In particular, we try to ascertain if the oblique structure suggested for D$_2$ in Refs. from neutron diffraction experiments, and in Ref. from low-energy electron-diffraction (LEED) measurements, is more stable than an arrangement consisting of two incommensurate triangular solids of different densities.

In the next Section, we describe the theoretical method used in the microscopic study of the adsorbed phases. The results obtained for both p-H$_2$ and o-D$_2$ on graphene are shown in Sec. III. Finally, the main conclusions are discussed in Sec. IV.

II. METHOD

The diffusion Monte Carlo (DMC) method is a stochastic technique that allows us to obtain the ground state of a zero-temperature system of bosons. Since the species adsorbed (p-H$_2$ and o-D$_2$) on graphene are both bosons, we can obtain through DMC the arrangement of molecules with the lowest energy for each surface density. In order to reduce the statistical variance of the many-body problem, the algorithm uses a guiding wave function $\Phi$ which enhances the occupation probability in places where the hydrogen density is expected to be large. In general, $\Phi$ depends on the coordinates of all atoms or molecules in the simulation cell. However, in this work, we will consider that the carbon atoms of the graphene layer are kept in fixed positions, what means that their overall effect on the hydrogen molecules can be described as an external potential.
proximation, the guiding wave function depends only on the positions of the \( N \) hydrogen molecules \((r_1, \ldots, r_N)\). We chose,

\[
\Phi(r_1, \ldots, r_N) = \Phi_J(r_1, \ldots, r_N)\Phi_1(r_1, \ldots, r_{N_1}) \\
\quad \times \Phi_2(r_{N_1+1}, \ldots, r_N),
\]

where \( N_1 (N_2 = N - N_1) \) is the number of molecules adsorbed in the first (second) layer, and \( \Phi_J \) is a Jastrow wave function used to take into account the \( H_2-H_2 \) and \( D_2-D_2 \) correlations. In particular,

\[
\Phi_J(r_1, \ldots, r_N) = \prod_{i=1}^{N} \exp \left[ -\frac{1}{2} \left( \frac{b}{r_{ij}} \right)^5 \right],
\]

\( r_{ij} \) being the distance between the center of mass of hydrogen molecules (considered as spheres and interacting through the isotropic Silvera and Goldman potential). This potential was developed to describe the hydrogen-hydrogen interaction in bulk and it was successfully used to reproduce the phase diagram of the first layer of both \( H_2 \) and \( D_2 \) on graphene. The variational parameter \( b \) in Eq. (2) was taken to be 3.195 Å for both \( H_2 \) and \( D_2 \), in agreement with variational optimizations done in previous simulations of similar systems.

The purpose of the remaining terms in Eq. (1), \( \Phi_1 \) and \( \Phi_2 \), is to describe the localization of hydrogen molecules in the first and second layers on top of graphene, respectively, and also the correlations due to the carbon-hydrogen interactions. Concretely,

\[
\Phi_n(r_1, r_2, \ldots, r_{N_n}) = \prod_{i}^{N_n} \prod_{J}^{N_C} \exp \left[ -\frac{1}{2} \left( \frac{bc}{r_{ij}} \right)^5 \right] \\
\quad \times \prod_{i}^{N_n} \exp(-a_n(z_i - z_n)^2)
\]

with \( n = 1, 2 \) for the first and second hydrogen layer, respectively, and \( N_n \) the number of molecules in each layer. \( r_{ij} \) represents the distance between the center or mass of each molecule, \( i \), and each of the \( N_C \) carbon atoms, \( J \), in the graphene layer. Each hydrogen molecule interacts with each of those carbon atoms by a potential of Lennard-Jones type, whose parameters are taken from Ref. [2]. This part of the guiding wave function is again similar to the one used in previous works to describe the first layer of \( H_2 \) (Ref. [3]) and \( D_2 \) (Ref. [7]) on graphene, to the point that we took the same \( a_1 \) and \( z_1 \) parameters as in those works. Thus, \( a_1 = 3.06 \, \text{Å}^{-2} \) for \( H_2 \) and \( a_1 = 5.2 \, \text{Å}^{-2} \) for \( D_2 \). \( bC = 2.3 \) Åand \( z_1 = 2.9 \) Åfor both hydrogen isotopes. For hydrogen molecules on the second layer, \( bC \) was kept constant, and \( a_2 \) and \( z_2 \) were variationally optimized. The optimal values were \( a_2 = 1.53 \, \text{Å}^{-2} \) for both \( H_2 \) and \( D_2 \), and \( z_2 = 6 \) Å(\( H_2 \)) and 5.8 Å(\( D_2 \)).

When the phase to be described is a quasi two-dimensional solid, \( \Phi_n \) is multiplied by a Nosanow term,

\[
\prod \exp\{-c_n[(x_i - x_{site})^2 + (y_i - y_{site})^2]\},
\]

where \((x_{site}, y_{site})\) are the crystallographic positions of the solid lattice. The \( c_n \) parameters were taken to be the same for \( n = 1, 2 \), i.e., a linear interpolation between the values corresponding to densities in the range 0.08 \( \text{Å}^{-2} \) (\( c_n = 0.61 \, \text{Å}^{-2} \)) and 0.10 \( \text{Å}^{-2} \) (\( c_n = 1.38 \, \text{Å}^{-2} \)) for \( H_2 \), and between 0.08 \( \text{Å}^{-2} \) (\( c_n = 1.11 \, \text{Å}^{-2} \)) and 0.11 \( \text{Å}^{-2} \) (\( c_n = 2.93 \, \text{Å}^{-2} \)) in the case of \( D_2 \). If the hydrogen density within the considered layer was not in those ranges, we used the linear extrapolated \( c_n \) value.

To model the second layer of molecular hydrogen on top of a first layer of the same substance, we followed closely the prescription of Ref. [4] in which a second layer of \( ^4\text{He} \) on graphene was simulated. Basically, for a fixed total hydrogen density, we considered only the arrangement for which the total energy per molecule was lower. In practice, this means that we have to take a fixed solid density for the first layer, and to change the number of molecules (if we have a liquid) or the lattice constant (if we have a solid) in the second layer. The structure of the second layer solid was assumed to be incommensurate with respect to the one on the first layer, i.e., the phase diagram was assumed to be of the same type as that of helium on graphene. However, to verify that this was so, we considered also the oblique commensurate structure proposed for \( D_2 \) from neutron diffraction and LEED. We checked the stability of that structure for both isotopes. To be able to treat incommensurate second layers, we used different periodic boundary conditions for the first and second layers. No exchange of molecules between the first and second layer was allowed. Importantly, we did not fix the positions of the molecules closest to the graphene surface, i.e., we took into account the zero point motion of all the hydrogen molecules. For both isotopes, we considered hydrogen densities up to those of promotion to a third layer, obtained experimentally for the same systems on graphite.

### III. RESULTS

#### A. \( H_2 \)

The phase diagram of the second layer of \( H_2 \) on graphene can be extracted from the data displayed in Fig. [1]. The full squares correspond to the energy per molecule of a triangular incommensurate solid adsorbed on a single layer on top of graphene, and were taken from Ref. [4]. When we considered only a hydrogen layer, incommensurate means that there is no registry of the hydrogen molecules with respect to the carbon sheet. The third-order polynomial fit displayed on top of them is a guide-to-the eye. To study second layer structures, we put on top of one of these incommensurate structures a set of hydrogen molecules described by a guiding wave function with \( c_2 = 0 \) (4). The dimensions of the simulation cells were determined by the density of incommensurate solid in the first layer since, in all cases, the simulation cells comprised \( 120 \, H_2 \) molecules close to the
graphene surface. Then, we put enough hydrogen on top of them to produce surface areas (the inverse of density) in the range displayed in Fig. 1. To be sure that the energy per molecule considering both hydrogen sheets was the minimum one, we performed several sets of simulations with different incommensurate solid densities (in the first layer). In particular, we considered 0.090, 0.095, 0.10 and 0.105 Å⁻². Our results indicate that on the second layer, the arrangement with lowest energy per H₂ molecule is a liquid on top of a solid whose density is 0.095 Å⁻². Those results are displayed in Fig. 1 as full circles. The lowest limit of the surface area corresponds approximately to the experimental value for H₂ promotion to a third layer (5.80 Å²).²³

We modeled the second layer incommensurate triangular solid on the same principles, i.e., we considered the same densities as for the liquid case in the first hydrogen sheet, and distributed the atoms on the second layer as to produce the total densities displayed as surface areas in Fig. 1. Here, incommensurate means that the second layer is registered neither with respect to the first layer nor with the underlying graphene. The minimum energies per H₂ molecule corresponded to an arrangement in which the lower layer density was 0.10 Å⁻²; those data are displayed in Fig. 1 as open squares. The line on top of them represents also a third-order polynomial fit. We considered also two second layer commensurate (with respect to the first layer) solids: a 4/7 lattice and an oblique bilayer structure.²⁸ both proposed originally to describe second layers of ⁴He on graphite. The latter was suggested to be stable for the second layer of H₂ and D₂ on graphite. The energy per H₂ molecule of those registered phases is represented by a full and open triangle, respectively. We can see that both structures are metastable with respect to a set of two incommensurate layers, since their energies per molecule are larger. Their corresponding energies per H₂ molecule are -274.9 ± 0.1 K (oblique structure), and -304.6 ± 0.1 K (4/7), versus -293.0 ± 0.1 K and -309.38 ± 0.08 K of the corresponding incommensurate structures of the same densities (0.173 and 0.157 Å⁻², respectively). A 7/12 commensurate solid, also proposed to be stable for helium,³⁰ and not displayed for simplicity, was also considered and found to be unstable with respect to a second layer incommensurate solid.

With all of that in mind, we can draw a double-tangent Maxwell construction to obtain the phase diagram of the second layer of H₂ on graphene. The slope of that line is minus the internal pressure at which the transition takes place, and has to be positive for stable arrangements. In addition, if several transitions are possible, one has to consider only the corresponding to the lowest pressure. The straight line displayed in Fig. 1 that joints a single layer incommensurate solid and its second layer counterpart fulfills all the necessary requirements. We can see that the energy per H₂ molecule of a second layer liquid (full circles) is always above the double-tangent line. This means that a second layer liquid is unstable with respect to a mixture of a single layer solid and another with two incommensurate sheets. The surface areas at which the slopes of both three-order polynomial fits are the same define the stability limits for the phases involved in the transition. In our case, those correspond to an upper density limit for a single layer solid (equivalent to a second layer promotion) of 0.094 ± 0.002 Å⁻², and a lowest density value of 0.173 Å⁻². Both results are in excellent agreement with the calorimetric values of H₂ on graphite:²³ 0.094 and 0.165 Å⁻². The chemical potential of H₂ at the monolayer-bilayer transition derived from our simulations was -161 ± 2 K. This means that a second layer of H₂ is still stable with respect to the formation of a bulk H₂ crystal, whose ground state chemical potential is -92.3 K.³⁰ This also means that the second layer solidifies at H₂ densities as low as 0.060 Å⁻². As in ⁴He,¹¹,¹² the first layer suffers a compression that produces a ~ 6% increase of its density upon the adsorption of the second hydrogen sheet.

We also compared our simulation results for H₂ on a second layer to similar data obtained for a pure two-dimensional (2D) system. In Fig. 2 we show the energy per H₂ molecule as a function of density, but only for the molecules on that second sheet. This energies are taken from the same set of simulation results as the ones dis-

![FIG. 1: (Color online) Maxwell construction (straight line) to determine the limits of the phase diagram of a second layer of H₂ on top of the same substrate on graphene. Full squares, energy per H₂ molecule on a first layer triangular incommensurate solid; full circles, same data for a second layer liquid on a triangular solid; open squares, a triangular incommensurate solid on top of first layer solid. The open triangle represents the energy per H₂ molecule of the oblique structure proposed in Ref. ²³, while the full triangle is the energy of a second layer 4/7 structure. The curves are three-order polynomial fits to the simulation results.](image-url)
played in Fig. I, i.e., the molecules of the first layer are not kept frozen. A glance to the previous figure indicates that the binding energies of these molecules are smaller than the ones located in the first layer by at least a factor of two. This is true for both liquid (full circles) and triangular incommensurate (full squares) phases. The lines on top of each set of symbols are least-squares fits to the expression

\[ E/N = (E/N)_0 + a(\rho - \rho_0)^2 + b(\rho - \rho_0)^3, \]

where \( \rho \) is the hydrogen density in that second layer and \( \rho_0 \) stands for the density at which the energy per H\(_2\) molecule has a minimum \( (E/N)_0 \). The parameters obtained for the liquid and solid phases, together with their first layer (taken from Ref. 4) and pure two-dimensional (from Ref. 29) counterparts, are shown in Table I. Those results indicate that a pure two-dimensional system of H\(_2\) molecules is a worse description for the second than for the first layer of H\(_2\) on graphene, and that the 2D equation of state is worse for a liquid phase than for a solid one. For instance, the energy differences, once subtracted the infinite-dilution energy \( (E/N)_\infty \), between a 2D system and a first layer of H\(_2\) on graphene are \( \sim 6.5\% \) for a liquid, and \( \sim 5\% \) for a solid. In comparison, those same differences between a 2D and a second layer system are \( \sim 71\% \) and \( 22\% \), respectively. The same, but to a lesser extend, can be said of the differences between the values of \( \rho_0 \): \( \sim 2.3\% \) for the 2D and first layer solids difference, gap that increases up to \( \sim 4.2\% \) if instead of a first layer we have a second one. Those same percentages grow to \( \sim 6.5\% \) and \( 15\% \) for the same comparisons for liquid phases. Moreover, Table I gives us another interesting piece of information: the upper solid is more stable with respect to a liquid arrangement than in a flat structure or a first layer sheet. This is because the energy difference at the respective equilibrium densities is highest in the second layer case. The corresponding values are 6.8 K (second layer), versus 2.023 K (2D), and 1.8 K (first layer). We can also see that the density minimum for the second-layer solid phase (0.0646 \( \pm \) 0.0009 \( \text{Å}^{-2} \)) is comparable to the equilibrium density mentioned above (0.060 \( \pm \) 0.002 \( \text{Å}^{-2} \)), obtained from the Maxwell construction including the whole system.

### Table I: Density and energy per molecule for the phases displayed in Fig. 2 from fits using Eq. (5). The results for a pure two dimensional system (2D, Ref. 29) and a first layer of H\(_2\) on top of graphene (gr, Ref. 4) are also shown for comparison. \( E_\infty \) is the energy per H\(_2\) molecule in the infinite dilution limit for each system, \( (E/N)_0 \) is the minimum energy per H\(_2\) molecule and \( \rho_0 \) is the density that corresponds to that energy.

|       | Liquid | Solid |
|-------|--------|-------|
|       |        |       |
| \( E_\infty \) (K) | -21.43 \( \pm \) 0.02 | -23.453 \( \pm \) 0.003 |
| \( (E/N)_0 \) (K) | -451.88 \( \pm \) 0.03 | -454.1 \( \pm \) 0.3 |
| \( (E/N)_0 - E_\infty \) (K) | -20.09 \( \pm \) 0.07 | -22.3 \( \pm \) 0.3 |
| \( \rho_0 \) (\( \text{Å}^{-2} \)) | 0.0633 \( \pm \) 0.0003 | 0.0673 \( \pm \) 0.0002 |
|       |        |       |
| \( E_\infty \) (K) | -431.79 \( \pm \) 0.06 |
| \( (E/N)_0 \) (K) | -51.098 \( \pm \) 0.03 |
| \( (E/N)_0 - E_\infty \) (K) | -20.09 \( \pm \) 0.07 |
| \( \rho_0 \) (\( \text{Å}^{-2} \)) | 0.06948 \( \pm \) 0.00005 |

{FIG. 2: (Color online) Energy per H\(_2\) for molecules located on the second layer. Full circles are the results of a liquid phase, and full squares, for a triangular incommensurate solid one. The error bars are similar in both cases and for simplicity they are only shown in the liquid case. Lines are three-order polynomial fits to the respective data.}

### B. D\(_2\)

This subsection will closely mirror the previous one since we studied the second layer of D\(_2\) on graphene following the same steps. Our results are summarized in Fig. 3, where all the symbols and lines have a similar meaning to those of Fig. I. The only difference is that the structures with the minimum energy per deuterium molecule are those whose first layer density is 0.105 \( \text{Å}^{-2} \) for both the second layer solid and liquid phases. To be sure of that, densities in the range 0.095 to 0.110 \( \text{Å}^{-2} \) were tested. The first layer results (full squares) are now taken from Ref. 5 and the triangles represent the same registered phases suggested above for H\(_2\). As one can see, these commensurate phases are still unstable with respect to a set of two incommensurate solid deuterium layers. As in H\(_2\), we have also that the energies per D\(_2\) molecule
for a second layer liquid. This means that the phases in equilibrium are again a first layer incommensurate triangular solid of density 0.100±0.002 Å⁻² and a second layer incommensurate solid whose total density is 0.175±0.002 Å⁻². Both results are again in excellent agreement with the calorimetric data of Ref. 23 on graphite: 0.099 Å⁻² for second layer promotion, and 0.178 Å⁻² for the minimum density at which the double solid is stable. Since in this last case, the density of the lower layer is 0.105 Å⁻², we can state that there is also a compression of the first layer of around a 5% upon the adsorption of a second layer of D₂ on top of D₂. As in the case of H₂, our simulation results allow us to calculate the chemical potential of D₂ at the monolayer-bilayer transition. The result was -180±2 K, a healthy 30% larger than the corresponding to bulk D₂. We found also that both the oblique incommensurate structure and the 4/7 one were unstable with respect to the double incommensurate arrangement, as can be seen in Fig. 3. The values of their energies per D₂ molecule were -326.1±0.1 K and -352.7±0.1 K, versus -341.1±0.1 K and -358.0±0.1 K for the double incommensurate for the same densities (0.186 and 0.164 Å⁻², respectively).

We can also study the second layer by itself and compare the results to those of a first layer of D₂ adsorbed on graphene, and to a pure 2D system. That can be done with the help of Fig. 4 and Table II. The conclusions we can draw from this set of information are similar to those already described in the H₂ case: the equilibrium densities are essentially compatible with each other, even more so in the solid case, and the energy per D₂ molecule differences between liquid and a solid phases are larger in the case of a second layer solid than for a single deuterium sheet. The equilibrium density of the second layer solid is also comparable to the one extracted from the Maxwell construction for the entire system (0.078 versus 0.070 Å⁻²).

**TABLE II: Density and energy per D₂ molecule for the phases displayed in Fig. 4, obtained by the same means as those of H₂. The results for a pure two-dimensional system (2D, Ref. 24 with only data for the solid phase) and a first layer of D₂ on top of graphite (gr, Ref. 2 for the liquid and this work of the solid) are also shown for comparison.**

| Density (Å⁻²) | Liquid | Solid |
|--------------|--------|-------|
| 2D           |        |       |
| (E/N)₀ (K)   | -326.1 | -358.0 |
| (E/N)₀ - E₀ (K) | -326.1 | -358.0 |
| ρ₀ (Å⁻²)     | 0.0785 | 0.0700 |
|        |        |       |
| gr          |        |       |
| (E/N)₀ (K)   | -464.87 | -495.0 |
| (E/N)₀ - E₀ (K) | -464.87 | -495.0 |
| ρ₀ (Å⁻²)     | 0.0785 | 0.0700 |

**FIG. 3:** (Color online) Same as in Fig. 1 but for D₂ instead of H₂. The lower limit for the surface density is fixed as the inverse of the experimental density for a promotion of D₂ to a third layer (5.8 Å², Ref. 23).

**FIG. 4:** (Color online) Same as in Fig. 2 but for D₂ instead of H₂.
IV. CONCLUSIONS

In this work, we have studied the complete phase diagram of the second layer of both $\text{H}_2$ and $\text{D}_2$ adsorbed on top of a single graphene layer. Since we used a set diffusion Monte Carlo calculations, the results correspond to the zero-temperature ground state of the system. To obtain the real stability limits, we had to perform Maxwell constructions between phases that comprised one and two hydrogen layers, but we found that if we used only the data corresponding to a second one, the description is good enough to reproduce the solid equilibrium of the complete two-sheet system. Our results for the promotion density to the second layer and the minimum density in this second layer agree satisfactorily with available calorimetric results for graphite, pointing to the accuracy of both the method used and the interaction potentials entering in the Hamiltonian. That our results on graphene are comparable to the experimental ones on graphite also means that both surfaces are basically equivalent as absorbents, being the only possible difference the binding energy of the hydrogen molecules to the carbon surface.

However, calorimetric measurements only give the total density for the onset of a solid structure at $T \to 0$. From Ref. 23, those densities appear to be 0.165 ($\text{H}_2$) and 0.178 ($\text{D}_2$) Å$^{-2}$. Both are lower than the ones assigned in the same work to the oblique structures: 0.173 ($\text{H}_2$) and 0.186 ($\text{D}_2$) Å$^{-2}$, respectively. The fact that the former densities are compatible with our simulation results supports our suggestion of a double incommensurate solid as the structure for the inferred solid phase. Moreover, a comparison of the energies per molecule for the oblique solid and the two sets of triangular layers at the same densities indicates that, at least for $T = 0$, the incommensurate arrangement is not stable. One could speculate that the disagreement between calorimetric data, in one side, and LEED and neutron diffraction, on the other, could be originated by the tiny difference between the diffraction patterns of a double incommensurate structure and a double oblique layer. In fact, the oblique phase was proposed in the past as the ground state of a second layer of $^4\text{He}$ on graphite in neutron diffraction studies, but further calorimetric measurements concluded that triangular structures were preferred. This could be also the case for hydrogen. On the other hand, our results rely on empirical potentials that have been used to reproduce reasonably well the equation of state of the first layer of both $\text{H}_2$ and $\text{D}_2$ on graphite, but we cannot exclude that, in the future, more elaborate interactions could change our predictions for the second layers.

On a different note, from the behavior of this second layer, we can say that when the number of adsorbed layers grows, the entire arrangement becomes more "solid-like" than "liquid-like". The reason is that, even though the binding energy in the second layer is lower than in the first one, the incommensurate solid structure is much more stabilized with respect to a liquid than in the first layer case. This is probably the reason why the experimental critical points for the liquid-vapor coexistence regions of $\text{H}_2$ and $\text{D}_2$ adsorbed on graphite approach those of the bulk solids as the number of adsorbed layers grows, making impossible to obtain a stable liquid by reducing the dimensionality of the system.

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