Theoretical investigation of methane under pressure

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Abstract

We present computer simulations of liquid and solid phases of condensed methane at pressures below 25 GPa, between 150 and 300 K, where no appreciable molecular dissociation occurs. We used molecular dynamics (MD) and metadynamics techniques, and empirical potentials in the rigid molecule approximation, whose validity was confirmed a posteriori by carrying out \textit{ab initio} MD simulations for selected pressure and temperature conditions. Our results for the melting line are in satisfactory agreement with existing measurements. We find that the fcc crystal transforms into a hcp structure with 4 molecules per unit cell (B phase) at about 10 GPa and 150 K, and that the B phase transforms into a monoclinic high pressure phase above 20 GPa. Our results for solid/solid phase transitions are consistent with those of Raman studies but the phase boundaries estimated in our calculations are at higher pressure than those inferred from spectroscopic data.
I. INTRODUCTION

Methane is the most abundant organic molecule in the universe, and the phase diagram of its condensed forms is of great relevance in several branches of science, including planetary physics and petrology. For example, spectroscopic data have revealed the presence of solid methane on the surface of different planets, e.g. Pluto, and the properties of methane at moderate pressures are crucial for modeling and understanding the formation and stability of hydrocarbons in the Earth’s mantle and crust. The phase diagram of methane is rather complex and poorly understood: at low pressure it exhibits several solid phases differing both for the positions of the carbon atoms and for the orientation of the molecules in the unit cell.

At room temperature and $P = 1.6$ GPa methane crystallizes in the so called phase I, with C atoms occupying fcc lattice sites while H atoms are free to rotate; phase I has one molecule per unit cell. By isothermal compression at room temperature starting at 1.6 GPa, transformations to different phases have been observed by X-Ray diffraction, IR and Raman studies; these transitions occur approximately at 5GPa (phase I to phase A), 12 – 18 GPa (A to B) and 25 GPa (B to a so called high-pressure phase-HP). Bini et al., based on IR and Raman data, proposed a tetragonal crystal structure for the phase A, while Umemoto and Hirai proposed instead a cubic structure for phase B. All experiments found that the A-B transition is very sluggish, suggesting a possible, complex structural rearrangement taking place between the two phases. Hirai et al. also suggested the existence of a phase intermediate between A and B, which they called pre-B, with a diffraction pattern totally different from the one of phase A. At pressure above 25 GPa, a so called high pressure (HP) phase appears, and the B-HP transition is not accompanied by a change in the spatial arrangement of the carbon atoms. Finally the melting line of methane has been measured between 100 K and 450 K in three different experiments.

In this paper we present molecular dynamics (MD) simulations based on empirical po-
potentials, aimed at investigating both the melting line of methane and its solid phases below 25 GPa and between 150K and 300K, i.e. in a regime where CH$_4$ molecules do not dissociate. We used a metadynamics algorithm$^{16,17,18}$ to investigate solid/solid transitions and compared the results of several techniques in the case of the solid/liquid transition. In addition we performed ab initio MD simulations using Density Functional Theory (DFT) to validate a posteriori the empirical model utilized in our classical MD simulations. We do not consider $T < 150K$, where quantum effects are expected to be important for the description of nuclear motion.

The paper is organized as follows. In Section II we briefly describe some technical aspects of both classical and ab initio simulations. In Section III we present calculations of the melting line of methane, comparing results obtained by thermodynamic integration$^{19}$ with those of a two-phase coexistence method$^{20,21}$. In section IV we discuss metadynamics results for solid/solid phase transitions and we justify a posteriori the use of the selected forcefield, by carrying out ab initio MD simulations for several (P,T) points. Finally in section V we draw our conclusions.

II. COMPUTATIONAL TECHNIQUES

We performed molecular dynamics (MD) simulations with the DLPOLY code$^{22}$, using the empirical force field TraPPE-EH from Ref.$^{23}$, which was parametrized to describe vapor-liquid equilibria in alkanes. CH$_4$ molecules are treated as rigid bodies and the intermolecular interactions are described by Lennard Jones (LJ) sites located on the carbon atoms, and on the centers of carbon-hydrogen bonds$^{23}$. As we discuss in detail below, ab initio calculations confirmed a posteriori the validity of the rigid molecule approximation. The MD equations of motions were integrated using the leap-frog Verlet algorithm with a time step of 0.5 fs.

We also carried out Born-Oppenheimer (BO) ab initio simulations in the NPT ensemble with the Qbox code$^{24}$, using a generalized gradient corrected approximation, PBE$^{25}$, for the exchange and correlation functional. We used norm conserving pseudopotentials and a plane-wave expansion of the Kohn-Sham orbitals with a kinetic-energy cut-off of 60Ry. For all the systems investigated here we used a Γ point sampling of the Brillouin zone. In our ab initio MD we substituted hydrogen with deuterium for computational convenience, so as to use a larger time step (5 a.u.) when integrating the Newton equations of motion.
III. CALCULATION OF METHANE MELTING LINE

Hysteresis effects near a solid liquid transition hamper an accurate estimate of melting temperatures from heat-until-melt MD runs. We carried out the calculation of the methane melting line by adopting two complementary techniques. One is based on the direct simulation of coexisting liquid and solid phases at given thermodynamic conditions, and we call it two-phase coexistence method. The second approach consists of calculating the free energies of the solid and the liquid, and in obtaining the temperature of coexistence of the two phases by determining when their free energies are equal. Once a point on the coexistence line is known, it is then possible to find the whole coexistence curve by integrating the Clapeyron equation, i.e. by varying the pressure and temperature in such a way as to maintain equal the chemical potentials of the liquid and the solid: $\mu_l = \mu_s$.

The two-phase simulations were performed using periodic boundary conditions, with supercells containing 3456 molecules, half of them prepared in the solid $fcc$ phase and half of them prepared in the liquid phase, at pressures of 0.01, 0.5 and 1. GPa. The liquid and solid phases were initially equilibrated separately in NPT ensemble runs (constant number of particles, pressure and temperature) with fixed cell shape and then the two systems were merged together in a single supercell with a solid-liquid interface (we call the direction perpendicular to the interface $z$ axis).

We performed MD simulations of the coupled solid/liquid systems in the NPH ensemble (constant number of particles, pressure and enthalpy). During the simulation the pressure is applied isotropically while the cell angles cannot deform. If the initial pressure and temperature are sufficiently close to the conditions at which the solid and the liquid coexist, the temperature equilibrates to the value on the coexistence line, e.g. to the melting temperature, otherwise one phase prevails over the other. This approach differs from the one used in ref, where coexisting phases are simulated in the NPT ensemble at different values of $(P,T)$, and for each run one determines the phase that is most stable under those conditions. When carrying out simulations in the NPH ensemble, if a reasonable choice of the initial $P − T$ conditions can be made, a single MD run suffices to determine the melting temperature. However, MD cells much longer than those adopted in NPT simulations are usually required. As an example, we report a plot of the temperature and enthalpy at 0.01 GPa in Fig. [1].
FIG. 1: Variation of enthalpy (upper panel) and temperature (lower panel) as a function of simulation time for a solid-liquid interface simulated in the (NPH) ensemble at 0.01 GPa. The solid lines are the running averages over a 0.5ps time interval.

In order to estimate finite size effects on our results, we carried out simulations with different cell sizes. The size of the simulated system may affect the values of the computed melting temperature in two ways: by the extension of the solid-liquid interface and, perhaps most importantly, by the size of the system in the direction orthogonal to the interface (the $z$ direction in our case). During the equilibration process in the NPH ensemble, solidification of the part of the system originally prepared as a liquid, and melting of the part originally prepared as a solid may occur, until a solid-liquid interface is stabilized. One of the two phases may take over, if the system size in the $z$ direction is not large enough, making it impossible to stabilize a two-phase system. In table I we summarize the simulation results obtained with different supercell sizes. In our calculations, we first fixed the length of the cell in the direction orthogonal to the interface, and verified that changes in the extension of the solid-liquid interface does not substantially affect the computed value of the melting temperature: simulations performed with the three supercells $6 \times 6 \times 24$, $5 \times 5 \times 24$, $4 \times 4 \times 24$ give the same melting temperature with an error of less than 1 K. As expected, size effects along the $z$ direction are more dramatic. We considered the same geometry along the interface as indicated above (e.g. $6 \times 6$ and $5 \times 5$ ) and we halved the length along the $z$ direction ($1726$ molecules with $6 \times 6 \times 12$ supercell and $1200$ molecules with $5 \times 5 \times 12$ supercell ): in this case the system completely transforms into one phase and no phase coexistence can be reached.
As a validation of the results obtained with the two-phase method, we also computed the solid-liquid boundary by a direct calculation of free energies. Using thermodynamic integration\textsuperscript{19}, we can compute the Gibbs free energy difference ($\mathcal{G}$) of a system with Hamiltonians $H^I$, and that of a reference system with Hamiltonian $H^{II}$, for which the melting curve is known by:

$$\mathcal{G}^I - \mathcal{G}^{II} = \int_0^1 d\lambda \langle H^I - H^{II} \rangle_\lambda.$$  \hspace{1cm} (1)

The ensemble average in the integrand of Eq. (1) is evaluated by performing MD simulations in the constant pressure and constant temperature (NPT) ensemble, for a system with Hamiltonian $H(\lambda) = \lambda H^I - (1 - \lambda) H^{II}$, at different values of $\lambda$. As a reference for our thermodynamic integration we chose a Lennard-Jones system ($\varepsilon_{LJ} = 0.010323577$ eV and $\sigma_{LJ} = 3.405$), which has the same equilibrium crystalline phase as methane, and for which the melting curve was accurately determined in previous calculations\textsuperscript{28}. Free energy differences $\Delta \mathcal{G}$ were computed for the coexistence conditions of the reference system, both for the crystalline and the liquid phase. In particular we performed NPT runs of about 50 ps, for supercells containing 108 CH$_4$ molecules, and we considered 20 values of $\lambda$ for each phase. Size effects were tested by repeating simulations for a system containing 500 CH$_4$ molecules; the results for the melting temperature differed from those obtained with 108 molecules by less than 1%. Once two $\Delta \mathcal{G}$ values were obtained, we performed simulations for each phase, using the reversible scaling method\textsuperscript{29,30} to determine the coexistence temperature, at a given pressure.

This procedure has been applied to find the melting temperature at $P=0$, and $P=0.5$.
GPa, yielding $T_c=113$ and 243 K, respectively, in excellent agreement with phase-coexistence calculations (see Fig. 2).

We can now use one of the points on the melting line calculated with the previous methods, as the initial condition for integrating the Clapeyron equation. We start from the Gibbs-Duhem equation,

$$d(\beta \mu) = h d\beta + \beta v dP,$$

(2)

where $\beta = 1/kT$, $h$ and $v$ are the molar enthalpy and volume respectively, $P$ the pressure. We then write the Clapeyron condition $\mu_s = \mu_l$ as a first-order ordinary differential equation

$$\frac{dP}{d\beta} = -\frac{\Delta h}{\beta \Delta v},$$

(3)

where $\Delta h = h_l - h_s$ is the difference in molar enthalpies between the coexisting phases, while $\Delta v = v_l - v_s$ is the difference in volume; the derivative is evaluated along the coexistence line $\sigma$.

We solved Eq. (3) using a predictor-corrector formula; simultaneously performing two NPT simulations for the liquid and the solid and then evaluating $\Delta h = h_l - h_s$ and $\Delta v = v_l - v_s$. We verified possible size effect using two different supercells of 108 and 500 molecules. First we checked that the solution of Eq. (3) is independent on the initial conditions, i.e. the choice of different points on the melting line as initial states does not affect the results obtained for the melting line. We also verified possible size effect using two different supercells of 108 and 500. For temperature $T < 350$ K size effects observed when solving Eq. (3) are less than 1% and the supercells with 108 and 500 molecules give the same results. For $T > 350$ the error on the melting temperature is about 5%, when using 108 molecules.

All the results obtained for the methane melting line are summarized in Fig. 2 together with the experimental values from $^{13,14,15}$. The melting line obtained solving the Clapeyron equation is almost identical to that given by the phase-coexistence method and by free energy calculations.

The overall agreement between computed and measured melting temperatures is satisfactory, with a rigid shift of approximately 28 K between the experimental and theoretical curves below 200K. Above 200 K the slope of the theoretical and experimental melting curves differ by about a factor of 4, making the agreement qualitative. The overall good agreement indicates that the forcefield used in our simulations is reasonably accurate in the
FIG. 2: Experimental and calculated melting line for the fcc-liquid phase transition. The experimental value are extracted from 13 (cross), 14 (stars), 15 (plus). Calculated melting points are obtained with phase coexistence method (triangle) and integration of the Clapeyron equation (open circles and open squares respectively). We did not include the points calculated with free energy integration, because they superimpose to the ones given by the phase coexistence method.

description of methane-methane interactions at moderate pressures. We use the same force field in the next section to investigate solid-solid phase transitions.

IV. SOLID-SOLID PHASE TRANSITION IN METHANE AT MODERATE PRESSURES

In order to predict the crystal structure of the higher pressure crystalline phase of methane, we used the metadynamics technique. Metadynamics 16 is a method for exploring free energy surfaces as a function of selected collective coordinates, e.g. simulation cell edges 17,18,31. This approach has been applied successfully to several systems including, the crystal structure prediction of organic molecules 32,33. Our simulations were performed with supercells of 108 and 168 molecules.

We first considered the compression of a fcc crystal (108 molecules) at 10 GPa, and 150 K and 300 K. For both temperatures a transition to a hcp structure with four molecule per unit cell was observed, in which the molecules freely rotate in their final configuration.
FIG. 3: Phase diagram of the solid methane. Experimental phase boundaries (solid line) are from 4. Squares correspond to the points where the system does not evolve from an fcc during a metadynamics simulation. Circles represent points where an fcc-hcp transition occurs during metadynamics run. A transition from hcp to a monoclinic crystal was observed with Parrinello-Rahman simulations (triangles) on the hcp structure. The dashed line is a tentative phase boundary between the hcp and the monoclinic phase.

We did not find any evidence of intermediate stable configurations, nor we observed sizable jumps in the volume or in enthalpy of the system during the transition. As discussed in 31,34, the analysis of the eigenvectors and eigenvalues of the Hessian matrix (see table (II)) allows for a detailed description of the dominant crystal deformations occurring during the fcc-hcp transition.

The projection of the eigenvectors of the Hessian matrix on the six independent components of the cell edge matrix $h$ (arranged in a six-component vector $(h_{11}, h_{22}, h_{33}, h_{12}, h_{13}, h_{23})$), are related to the elastic modes of the initial crystalline phase and the eigenvalues are related to the elastic moduli, i.e. the curvature of the free energy along the corresponding eigenvectors. For example, the largest eigenvalue corresponds to the eigenvector $s_6$ in Tab. (II) that points along the (111000) direction, and this is associated to a hydrostatic volume contraction/expansion. In this case $s_1$ and $s_2$ are two equivalent orthorhombic deformations, which involve a compression/dilatation along pair of axes, while $s_3$, $s_4$ and $s_5$ are shear modes involving changes of the off-diagonal components of the Hessian matrix. In Fig. (III) we report the evolution of the different modes during a metadynamics simulation: the only inactive mode is $S_3$, confirming that a complex, concerted rearrangement
of the atoms take place during the fcc-hcp transition.

Metadynamics simulations were performed also at 5 GPa and 8 GPa, but no additional stable crystal structures were found. Based on a possible similarity between CH$_4$ and CF$_4$, Hirai et al. suggested that a phase A exists, with a cubic structure with 21 molecule per cell, similar to a structure found for carbon fluoride. In order to test this hypothesis, we performed a metadynamics simulation at 8 GPa and 300 K with 168 molecules/supercell, that is commensurate with 21 molecules per unit cell. In this case we observed a transition to a defective hcp structure and no stable structure corresponding to that of CF$_4$.

In order to explore the validity of the rigid molecule approximation used here, we performed BOMD simulations to test a posteriori our approximation. In particular, we carried out BOMD simulations starting from a hcp phase at 10 GPa and 20 GPa. After a short simulation of $\sim$1 ps, at both pressure we did not observe any significant distortions in the shape of the charge densities localized on the molecules, and each CH$_4$ does behave as a rigid body. In Fig. (5) we report the distribution of the intra-molecular angle between C-H bond. The angle distribution has the same profile both at 10 GPa and 20 GPa and is centered around the fixed value $\theta = 1.9102$ rad used in our classical MD simulations.

Starting from the hcp structure, we performed a series of constant pressure Parinello-Rahman simulations for different values of the pressure. At the temperature of 150K the
FIG. 4: Evolution of the metadynamics collective variable during the transition at 150 K and 10 GPa from fcc to hcp

FIG. 5: Distribution of the intramolecular C-H, C-H angle calculated from ab initio molecular dynamics simulations at 10 GPa (dashed dotted line), 20 GPa (solid line) and 150 K.

hcp structure with freely rotating molecules is stable up to 20GPa, when deformation of one of the cell angle occurs (Fig. 6). The hcp cell with four molecules per unit cell transforms into a monoclinic structure (P21/c) with four molecule per unit cell. At 20GPa the molecules stop rotating and a denser crystal structure is stabilized. The “freezing” of the hydrogen atoms is accompanied by a deformation of one of the cell angles as clearly shown in Fig. 6,
FIG. 6: Cell angle as a function of the simulation time. The Parrinello-Rahman simulation was performed at 20 GPa and 150 K, starting from an hcp crystal structure with four molecules per unit cell, in an orthorhombic supercell containing 108 molecules. Inset: a snapshot of the final carbon position from the [100] and [001] directions.

where the angles are plotted as a function of the simulation time. A snapshot of the carbon positions in the final configuration at 20GPa, is shown in the inset of Fig. 6 along the [100] and [001] directions.

V. CONCLUSION

We have presented results on the methane melting line, in satisfactory agreement with existing measurements and discussed in detail the comparison between several different computational techniques. These results also validate the empirical potential in the thermodynamic regime considered in this work.

In addition we propose two crystal structures for methane at moderate pressure and we provide insight into the transition mechanisms between such crystalline phases. The results from metadynamics calculations and Parrinello-Rahman simulations are summarized in Fig. 6 together with the experimental boundary lines obtained from 4. In our simulations we find that under pressure, at temperatures between 150 and 300 K, the fcc crystals transforms into a hcp structure with 4 molecules per unit cell (B-phase) and then into a monoclinic (4
molecules per cell) high pressure phase. These results are consistent with those of IR and Raman investigations by Bini et al., although we did not find any A phase, intermediate between the fcc and B phases. In addition, the estimated solid/solid phase boundaries are at higher pressure in our calculations than those experimentally inferred from spectroscopic experiments. The absence of an A phase is not entirely surprising as the A-B transition has been found to be very sluggish in all experiments reported to date, and it may be accompanied by a complex atomic rearrangements not commensurate with the MD cells used in our simulations. We did not find any evidence of a phase with 21 molecules per unit cell proposed in Ref. by analogy with the CF₄ crystal and our results do not seem to agree with the interpretation of the experimental findings of Refs. While additional work is required to understand in detail the solid phases of methane, our simulations represent a useful set of qualitative results, consistent with available experimental spectroscopic evidence, on which more refined \textit{ab initio} MD simulations can be based upon.

Further work is in progress to study the methane phase diagram in a regime where molecules dissociate, using first principle techniques.

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