We present an ab – initio study of boron nitride as well as carbon allotropes. Their relative thermodynamic stabilities and structural phase transitions from low- to high-density phases are investigated. Pressure-temperature phase diagrams are calculated and compared to experimental findings. The calculations are performed using quantum chemical wavefunction based as well as density functional theories. Our findings reveal that predicted energy differences often depend significantly on the choice of the employed method. Comparison between calculated and experimental results allows for benchmarking the accuracy of various levels of theory. The produced results show that quantum chemical wavefunction based theories allow for achieving systematically improvable estimates. We find that on the level of coupled cluster theories the low- and high-density phases of boron nitride become thermodynamically degenerate at 0 K. This is in agreement with recent experimental findings, indicating that cubic boron nitride is not the thermodynamically stable allotrope at ambient conditions. Furthermore we employ the calculated results to assess transition probabilities from graphitic low-density to diamond-like high-density phases in an approximate manner. We conclude that the stacking order of the parent graphitic material is crucial for the possible formation of meta-stable wurtzite boron nitride and hexagonal carbon diamond also known as lonsdaleite.

I. INTRODUCTION

The pressure-temperature phase diagrams of carbon and boron nitride reflect a delicate balance between weak and strong interatomic interactions. Although covalent bonds are the main source of their large cohesive energies, the accumulation of weak van der Waals interactions contributes significantly to the relative stability of their low- and high-density phases. Furthermore vibrational effects play a crucial role in the temperature dependence of the equilibrium phase boundary. Altogether this makes the prediction of phase diagrams and structural phase transition pathways a challenging task for modern electronic structure theories. In this work we seek to investigate boron nitride as well as carbon allotropes using various approximate electronic structure theories and compare theoretical with experimental findings. The aim is to benchmark their accuracy and help interpreting experimental results better if possible. To this end we employ a range of approximate density functional theories (DFT) and quantum chemical wavefunction based methods.

During the last decades approximate exchange and correlation (XC) density functionals have made significant progress in becoming more accurate and predictive for the description of interatomic interactions while keeping a high level of computational efficiency that allows for studying systems containing several hundreds of atoms routinely. The so-called Jacob’s ladder describes a ladder of approximations for the XC energy using increasingly complex as well as in general more accurate methods [1].

These rungs include functionals based on the local density approximation (LDA) [2, 3], the generalized gradient approximation (GGA) [4], the meta generalized gradient approximation (mGGA) [5] and hybrid functionals [6–9]. The latter include a fraction of (screened) exact exchange energies and have a computational cost that is comparable to Hartree–Fock theory. However, all the functionals mentioned above suffer from shortcomings that are despite many efforts difficult to remedy [10]. In the context of the present work a significant shortcoming is the inaccurate description of long range van der Waals interactions. In order to describe van der Waals and related interatomic interactions more accurately in the framework of approximate XC density functionals, a wide variety of dispersion corrections has been developed. As a consequence of the large number of available density functionals and corrections, there are numerous ground state energy functionals that could be considered in the present work, of which we have only chosen a small selection.

As a complement to the treatment of exchange and correlation on the level approximate density functionals, the computationally significantly more expensive quantum chemical wavefunction based theories are becoming more popular for the study of periodic systems [11–29]. This can partly be attributed to the increase in their computational efficiency, due to methodological developments, and to their ability to predict exchange and correlation energies in a systematically improvable manner. Quantum chemical methods constitute a hierarchy, which starting from the one-particle Hartree–Fock (HF) approximation, allows for a systematic treatment of the quantum many-body effects. The simplest form of such correlated methods is the second-order Møller–Plesset
(MP2) perturbation theory [30]. The next level of theory that achieves a significantly improved trade-off between accuracy and computational cost is based on the coupled cluster ansatz for the many-electron wavefunction [31]. Coupled cluster singles and doubles theory provides a compelling framework of infinite-order approximations in the form of an exponential of cluster operators [32]. The coupled-cluster singles and doubles (CCSD) method where the triples are treated in a perturbative way, termed as CCSD(T), achieves chemical accuracy in the description of many molecular properties and is sometimes referred to as the gold standard method [33].

In this work we seek to investigate the accuracy of the electronic structure theories mentioned above for carbon and boron nitride allotropes. To this end we compare predicted ground state energy differences as well as calculated phase diagrams to experimental findings. Experimentally several phases have been synthesized as single crystals [34–38] or as powder [39, 40]. Single crystals are usually synthesized in a closed chamber over a longer time period crystallizing from a solution. For synthesizing metastable structures the samples are put under static pressure and heated electrically or by laser [41–45]. Another way of transforming samples into metastable phases is shock wave synthesis [46, 47]. With these materials thermodynamic characterization can be performed [48–53] by determining relative enthalpy, entropy and heat capacity. These properties can be used to compare with the calculated energy differences and construct phase diagrams. Furthermore the phase diagrams can also be obtained by observing phase transitions directly [54–61]. In the present work we also investigate pressure-driven concerted phase transition pathways. In particular we study activation barrier heights for the transformation from low-density to high-density systems considering small unit cells that contain a few atoms at most. These models are far from realistic conditions under which phase transitions occur in experiment. Temperature-driven kinetic effects and catalysts are needed in practice to observe phase transitions close to the equilibrium phase boundary [62–65]. Hot liquid metals can be used to dissolve graphite and diamond will precipitate at the cooler region. However, the aim of the current work is to explore the accuracy of various electronic structure theories for transition states occurring in these phase transitions and provide a qualitative description of the various possible phase transition mechanisms. We believe that the small supercells considered are sufficient to describe these effects qualitatively correct.

This paper is organized as follows. Section II provides a description of the considered stable and metastable structures followed by an overview of the considered phase transition pathways. The employed structures can also be found in the Supplementary informations [66]. The results section IV summarizes the calculated ground state energy differences of the (meta-)stable structures and their activation energies at 0 K for the investigated phase transition pathways. By calculating the Gibbs energies, pressure-temperature phase diagrams are predicted and compared to experiment. Furthermore we assess the temperature and pressure dependence of the activation energies. Based on these results the experimentally observed phase transitions will be reviewed. Furthermore the existence of the wurtzite structure of carbon and boron nitride will be discussed. In the course of the discussion of these results we will assess the accuracy of the various approximate electronic structure theories.

II. CRYSTAL STRUCTURES AND PHASE TRANSITION PATHWAYS FOR CARBON AND BORON NITRIDE

A. (Meta-)stable structures

![View along c-axis with different stacking order. A' is rotated by 60° compared to A. Translating A by the red arrows creates the B, C and D layers. Blue arrow translates the D to a B layer. Panel (a) shows the lattice vectors of the 8 atomic orthorhombic unit cell employed in Tab. I. The crystal structures for carbon can easily be derived from boron nitride by substituting all B- and N-atoms by C-atoms.](image_url)
In this work we consider the most abundant crystal structures of carbon and boron nitride: the graphitic and diamond-like phases. Fig. 1 illustrates the corresponding structures. The low-density phases are graphitic with all atoms being $sp^2$ bonded and arranged in the planar honeycomb lattice with different stackings: AA, AB and ABC as depicted in Fig. 1. The G-AB and G-ABC have been observed experimentally for carbon and can be transformed into each other by translation of the layers. For boron nitride the stable low-density phase is (hexagonal) $h$-BN. $h$-BN exhibits an AA stacking order, indicating the other atom types for lattice sites on top of each other in the direction of stacking as shown in Fig. 1a. We note that the AA (G-AA) stacking is unstable for carbon.

The high-density phases of carbon and boron nitride are diamond-like. All atoms in the considered diamond-like phases can be assigned to chair or boat conformations of six-membered rings and two different stacking orders. For carbon and boron nitride the most stable high-pressure phases are cubic diamond ($c$-D) and zinc blende ($c$-BN), respectively. $c$-D and $c$-BN consist of six-membered rings of $sp^3$ bonded atoms in the chair conformation. There is a second high-density phase that is referred to as wurtzite for boron nitride and hexagonal diamond ($h$-D) for carbon with AA stacking order, exhibiting chair and boat conformations as illustrated in Fig. 1f. The mirror image of A is A, with the mirror parallel to the layer. The chair and boat conformations are parallel and perpendicular to the stacking, respectively. $h$-D is also known as lonsdaleite and serves as marker for shock impact events.

The crystal structures for C can easily be derived from BN, by substituting all B- and N-atoms with C-atoms. This makes (rhombohedral) $r$-BN (Fig. 1c), $c$-BN (Fig. 1e) and $w$-BN (Fig. 1f) equivalent to G-ABC, $c$-D and $h$-D, respectively.

Experimentally single crystals have been reported for $h$-BN [34, 35], $c$-BN [36], G-AB [37] and for $c$-D [38]. There exist no single crystals for $w$-BN, but the XRD measurements show no sign of $c$-BN in the samples and a small amount (<2%) of the starting material $h$-BN [39]. The synthesis of $r$-BN results in fibrous micro-crystals, but show no mixture with $h$-BN [40]. The $h$-D phase has been investigated in a number of theoretical as well as experimental studies in the past [54, 67–69]. However, recent experimental studies indicate that the previously believed samples of $h$-D are in fact $c$-D crystals that contain a large number of twins and stacking faults, creating X-ray diffraction patterns similar to the hypothetical $h$-D [70].

**B. Structural phase transition pathways**

We now discuss the investigated structural phase transition pathways. For the present study we keep the computational cost of the coupled cluster theory calculations low by restricting ourselves to transition state geometries that contain at most four atoms in the unit cell. Similar transition states have already been investigated in Refs. [67–69, 71–74]. To drive a transition from the low-density graphitic phases to the high-density diamond-like phases the application of pressure is needed. Under pressure the c-axis of the graphitic phase experiences a much larger compression than the other axes and will therefore be referred to as the compression axis. At high pressures the planar structure of graphite splits. Fig. 2 depicts two basic mechanisms by which the splitting of the planar six-membered rings present in the honeycomb lattice occurs. The mechanisms are referred to as buckling or puckering. Buckling and puckering creates the boat and chair conformation of six-membered rings, respectively. We employ the following naming convention for transition states. The first letter refers to the puckering (p) or buckling (b) mechanism and the second letter refers to the cubic (c) or wurtzite (w) structure corresponding to the final state of the considered transition.

pc-TS is the transition state in the G-ABC to $c$-D transition via the puckering mechanism. We note that the cubic phase contains only six-membered rings in the chair conformation. Therefore it is reasonable to consider this
FIG. 4. (a) \( h \rightarrow BN \) (b) \( pw-TS \) (c) \( BN; [72]; C; [68] \). Red arrows indicate the atomic displacements and support together with the atom numbers the assignment during the phase transition. Dotted lines show new bonds to be formed and red lines represent strong interaction during the transition state.

FIG. 5. (a) \( BN-AB \) (b) \( bw-TS \) (c) \( w-BN \). Red arrows indicate the atomic displacements and support together with the atom numbers the assignment during the phase transition. Dotted lines show new bonds to be formed and red lines represent strong interaction during the transition state.

We also consider the transition from \( w-BN \) to \( c-BN \), which occurs in a stepwise layer-to-layer rearrangement through 4H intermediate structures and will be referred to as \( l-pc-TS \) in this work [43]. During this transformation the boat conformation along the c-axis of the \( w-BN \) structure transforms into the chair conformation, while the six-membered rings perpendicular to the c-axis break apart (e.g. atoms 3+8 bottom layers) and rebond differently (e.g. atom 4+7 bottom layers) (Fig. 6a \( \rightarrow 6c \) with plane \((001)_w \parallel (111)_c \) and direction \([10\overline{1}0]_w \parallel [112]_c \).

In total we consider four different transition states including pc-TS, pw-TS, bw-TS and l-pc-TS for carbon and boron nitride allotropes. These parameters have been optimized using DFT in the LDA. This is necessary because forces are not yet implemented in the employed coupled cluster theory code. We believe that the LDA provides sufficiently accurate structures compared to experiment that allow for an unbiased comparison between the employed electronic structure theories and to experiment. We note that the LDA lattice parameters deviate by about 1\% only from experiment even for the lattice vector \( |c| \) parallel to the

C. Crystal lattice parameters

Tab. I summarizes the lattice parameters of the employed geometries for the low- and high-density phases and transition states for carbon and boron nitride allotropes. These parameters have been optimized using DFT in the LDA. This is necessary because forces are not yet implemented in the employed coupled cluster theory code. We believe that the LDA provides sufficiently accurate structures compared to experiment that allow for an unbiased comparison between the employed electronic structure theories and to experiment. We note that the LDA lattice parameters deviate by about 1\% only from experiment even for the lattice vector \(|c| \) parallel to the
compression axis as summarized in Tab. II. The only exception is h-BN, where the deviation is slightly larger. To allow for a direct comparison between the lattice parameters of the (meta-)stable structures as well as transition states we consider an eight atomic unit cell with monoclinic lattice vectors $\vec{a}$, $\vec{b}$ and $\vec{c}$ such that $|\vec{a}| \neq |\vec{b}| \neq |\vec{c}|$, $\beta = \gamma = 90^\circ$ and $\alpha$ can also be $90^\circ$. The $\vec{a}\vec{b}$ plane can be seen in Fig. 1d. The length of $\vec{a}$ corresponds to the width of a honeycomb ring and the vector points out of plane in Fig. 3 – 6. $\vec{b}$ points from left to right and $\vec{c}$ from bottom to top and spans across two layers. The ratio $|\vec{a}|:|\vec{b}|$ is 1:1:3 for all structures except for bw-TS and l-pc-TS. The ratio of R is larger by up to 3%. We point out that in l-pc-TS R is much larger compared to the other transition states. However, R refers to all bonds between the layers in the other transition states, whereas this not the case for l-pc-TS. In l-pc-TS additional interlayer bonds exist with a bond length of 1.654 Å (1+6 or 2+5 in Fig. 6b). For carbon (BN) the average of these two bond lengths is 2.063 Å (2.050 Å), respectively and comparable to R of the other transition states. The employed structures can be found in the Supplementary informations [66].

### III. METHODS

A. Density functional and Hartree–Fock theory

All electronic structure calculations have been performed using the projector augmented wave (PAW) method [79] as implemented in the Vienna ab – initio simulation package (VASP) [80, 81]. We present results obtained using some of the most widely-used $ab – initio$ methods to approximate the exchange and correlation energy in the framework of DFT. These methods include the LDA functional as parametrized by Perdew-Zunger [2, 3], the GGA functional as parametrized by Perdew-Burke-Ernzerhof (PBE) [4], the dispersion corrected PBE functional using the many-body dispersion energy method (PBE+MBD) [82, 83], the meta-GGA as parametrized for the SCAN functional [5] and the hybrid density functionals PBE0 [6], Becke-3-parameter-Lee-Yang-Parr (B3LYP) [7] and Heyd-Scuseria-Ernzerhof (HSE06) [8, 9]. We note that we have chosen only a small selection of functionals that could be considered.

The B 2s$^2$2p$^1$, N 2s$^2$2p$^3$ and C 2s$^2$2p$^4$ states have been treated as valence states in all calculations.

The geometries have been relaxed until the forces on all atoms are smaller than 10$^{-5}$ eV Å$^{-1}$. The total energies have been converged using the self-consistent field approach to within 10$^{-8}$ eV. For all DFT and HF calculations we employed a 16 atom supercell and a $4 \times 4$ Monkhorst-Pack $k$-point mesh. The corresponding supercell structures are summarized in the supplementary informations. The kinetic energy cutoff for the plane

| TABLE I. Structural parameters of the eight atomic monoclinic type unit cell ($\beta = \gamma = 90^\circ$) including interlayer distance $d$, the forming bond distance $R$ marked red in the transition state of Fig. 3 – 6 and its corresponding bond length in the other structures, atomic volume $V$ and density $\rho$. See text for further details. |
|---|---|---|---|---|---|---|---|
| Carbon | G-ABC | G-AB | G-AA | pc-TS | pw-TS | bw-TS | l-pc-TS | c-D | h-D |
| $|\vec{a}|$ [Å] | 2.446 | 2.446 | 2.445 | 2.465 | 2.461 | 2.471 | 2.456 | 2.498 | 2.484 |
| $|\vec{b}|$ [Å] | 4.236 | 4.236 | 4.236 | 4.270 | 4.263 | 4.164 | 4.219 | 4.329 | 4.302 |
| $|\vec{c}|$ [Å] | 6.735 | 6.590 | 7.171 | 4.899 | 4.780 | 4.835 | 4.705 | 4.326 | 4.137 |
| $\alpha$ | 77.9 | 90.0 | 90.0 | 73.1 | 90.0 | 90.0 | 90.0 | 70.5 | 90.0 |
| $d$ [Å] | 3.292 | 3.295 | 3.585 | 2.344 | 2.390 | 2.418 | 2.319 | 2.039 | 2.069 |
| $R$ [Å] | 3.292 | 3.295 | 3.585 | 2.058 | 2.108 | 2.100 | 2.472 | 1.530 | 1.549 |
| $V$ [Å$^3$ atom$^{-1}$] | 8.528 | 8.535 | 9.285 | 6.167 | 6.268 | 6.220 | 6.021 | 5.510 | 5.526 |
| $\rho$ [g cm$^{-3}$] | 2.417 | 2.415 | 2.220 | 3.343 | 3.289 | 3.315 | 3.424 | 3.742 | 3.731 |

| TABLE II. Relative difference to experimental values [39, 75–78] of lattice vectors $|\vec{a}|$ and $|\vec{c}|$ for stable structures of carbon an BN. In the case of w-BN, h-BN, h-D and G-AB we compare to a range of experimental values. |
|---|---|---|---|---|---|---|
| c-BN | w-BN | h-BN | c-D | h-D | G-AB |
| $|\vec{a}|$ [%] | 0.94 | 1.03–1.04 | 0.47–0.63 | 0.97 | 1.04–0.96 | 0.58–0.62 |
| $|\vec{c}|$ [%] | 0.63–1.21 | 2.53–2.55 | –0.62–1.10 | 0.78–1.75 |
wave basis set was set to 1000 eV. We note that smaller kinetic energy cutoffs would have sufficed but these calculations do not constitute a computational bottle neck compared to the more expensive coupled cluster theory calculations.

The phonon calculations have been performed using the PHONOPY code [84], creating the displacements within a $2 \times 2 \times 2$ supercell of the 16 atom cell. The forces are calculated using VASP and the LDA. These calculations employed a kinetic energy cutoff of 800 eV and a $2 \times 2 \times 2$ $k$-mesh.

B. Quantum chemical wavefunction theories

Results obtained using post-Hartree–Fock methods have been converged with respect to several computational parameters including energy cutoffs defining the plane wave basis sets, the number of virtual orbitals and the $k$-mesh. We have employed kinetic energy cutoffs of 500 eV for defining the orbital plane wave basis set and 300 eV for defining an auxiliary basis set that is used in the calculation of electron repulsion integrals required in post-HF methods. For the twist averaging technique we have employed a $4 \times 4 \times 4$ $k$-mesh. Furthermore 14 unoccupied orbitals have been used per atom. These parameters ensure a convergence of the energy difference between graphite and diamond to within a few meV per atom. The same parameters have been employed in Ref. [85]. For the virtual orbital space we employ MP2 natural orbitals that are obtained using a procedure outlined in Ref. [14]. Our estimates of the remaining basis set incompleteness error indicate that the energy difference between carbon diamond and graphite should be converged to within approximately 4 meV atom$^{-1}$ as shown in Fig.2 of Ref. [85]. A similar level of accuracy is expected for the other energy differences.

1. Finite size errors

We stress that the convergence of ground state energies obtained using post-HF methods such as MP2 theory with respect to the employed $k$-mesh or supercell size is slower than for their DFT counterparts. Wavefunction based methods account for non-local electronic correlation effects explicitly and therefore the observed interatomic interactions such as van der Waals forces lead to a slower rate of convergence with respect to the employed $k$-mesh. Recently we have introduced a finite size correction scheme that allows for accelerating the rate of convergence for periodic systems. We refer to results obtained using the finite size correction scheme by employing the following naming convention. Corrected MP2 and CCSD results are referred to as MP2-TA-FS and CCSD-TA-FS, respectively. TA and FS stand for twist averaging and an interpolation method, respectively. For the perturbative triples (T) correction on top of CCSD-TA-FS, we employ the twist averaging technique only. As such CCSD(T)-TA refers to CCSD-TA-FS plus the (T)-TA contribution. The improved $k$-mesh and supercell size convergence of CCSD-TA-FS was demonstrated and discussed in Ref. [85]. If not stated otherwise, all MP2, CCSD and CCSD(T) results in this work include the finite size corrections.

In the present work we employ box plots to depict finite size errors in Figs. 8, 9, 10, 11, 12 and 13. The box plots show the distribution of the obtained results for a set of different $k$-meshes. We stress that finite size errors must not be confused with stochastic errors. However, small error bars indicate that results are not affected significantly by the size of the employed $k$-mesh and can therefore be considered converged with respect to the $k$-mesh density. The employed box plots mark the largest and lowest value, the second and third quartile as well as the mean value. The bars in Figs. 8, 9, 10, 11, 12 and 13 mark the value with the largest $k$-point mesh. Some results for certain $k$-meshes differ significantly from the other $k$-meshes and were marked as outlier with a “+”. We find that very anisotropic Brillouin zone sampling using $k$-meshes with one $k$-point along one direction only yields results that are considered outliers.

For the results depicted in Figs. 8 and 11 we employ two-atomic unit cells and the following $k$-meshes: $2 \times 2 \times 2$, $2 \times 2 \times 4$, $3 \times 3 \times 2$ and $3 \times 3 \times 3$.

For the calculations of the barrier heights shown in Figs. 9, 10 and Figs. 12, 13 we have employed four and eight atomic unit cells. The following $k$-meshes were employed to sample the Brillouin zone of the four atomic unit cells G-ABC/h-BN, pc-TS, pw-TS (bw-TS and l-pc-TS in parentheses): $2 \times 2 \times 2$ ($1 \times 2 \times 4$, $3 \times 3 \times 1$ ($1 \times 3 \times 3$) and $3 \times 3 \times 2$ ($2 \times 3 \times 3$). We note that two transition states (bw-TS and l-pc-TS) need four atoms per layer to be described correctly and have only one layer. Therefore the $c$-axis is just half as long and the $a$-axis is doubled compared to the other cells. Consequently the $k$-mesh was adjusted appropriately. For the eight atomic unit cell we have employed a $1 \times 3 \times 2$ $k$-mesh.

C. Thermodynamic properties

For the calculation of pressure-temperature phase diagrams we need to compute the Gibbs energies ($G$) of all phases. $G$ is defined as the sum of the ground state energy, as obtained from DFT or a similar approach ($E_{\text{tot}}$), all entropic contributions and the $pV$ term. The vibrational ($F_{\text{vib}}$) contribution is the largest entropy related contribution. Using the finite displacement method a phonon density of state ($D(\omega)$) has been obtained for the frequency range ($\omega$). This phonon density of state contains the vibrational information for a specific volume ($V$) and can be used to calculate $F_{\text{vib}}$ at any temperature ($T$) with the Planck ($h$) and Boltzmann constant.
peratures mostly arises from the explicit dependence of the behavior of the transition probabilities at low temperatures and a selection of transition states. We note that transition probabilities and activation energies for carbon have been published in Refs. [87, 88].

The universal equation of state (EOS) [86] has been used to find the minimum of the Gibbs energy is calculated for at least five different volumes. The universal equation of state (EOS) [86] has been used to find the minimum of $G$ with respect to the volume for a given temperature and pressure. For all graphite-like phases and transition states anisotropic expansion along the c-axis has been included. One parameter changes the unit cell isotropically and a second one changes only the c-axis. A fourth order polynomial fit is used to interpolate the EOS between the sampling points. This increases the number of data points and also the accuracy. Using the procedure described above we have calculated the Gibbs energies employing the LDA for a wide range of pressures and temperatures. However, due to the computational cost involved it is currently not possible to perform the same calculations on the level of CCSD(T). The CCSD(T) Gibbs energies have therefore been approximated using the following expression

$$G_{\text{CCSD(T)}}(V,T) \approx G_{\text{LDA}}(V,T) - E_{\text{tot}}^{\text{LDA}}(V_0) + E_{\text{tot}}^{\text{CCSD(T)}}(V_0),$$

where $V_0$ corresponds to the LDA (equilibrium) volume of the (meta-)stable allotropes and the transition state geometries. As such the volume and temperature dependence of the CCSD(T) Gibbs energy is approximated using LDA, which achieves sufficiently accurate descriptions of the phonons and bulk moduli for the purpose of the present study.

### D. Phase transition probability

In this work we will approximate the probability that a phase transition occurs using the activation energy only and disregard kinetic effects. The activation energy ($\Delta G$) is the difference between the Gibbs energies of the transition and initial states. The Gibbs energy can be calculated as described in the previous subsection. Close to the equilibrium phase boundary the back and forward reaction has to be taken into account and the probability $f$ depends exponentially on $\Delta G$

$$f = \exp\left(-\frac{\Delta G_{\text{forward}}}{RT}\right) - \exp\left(-\frac{\Delta G_{\text{back}}}{RT}\right)$$

with $R$ as the gas constant. A comparable ansatz was published in Refs. [87, 88].

Fig. 7a to 7f shows the calculated CCSD(T) phase transition probabilities and activation energies for carbon and a selection of transition states. We note that the behavior of the transition probabilities at low temperatures mostly arises from the explicit dependence of $f$ on the temperature in the exponent rather than the temperature dependence of $\Delta G_{\text{forward}}$ or $\Delta G_{\text{back}}$. Sec. IV D provides a more detailed discussion of the obtained results.

### IV. RESULTS AND DISCUSSION

We now turn to the discussion of the obtained DFT, HF and post-HF results. The following section is organized as follows. We first summarize the energy differences obtained using different methods for carbon (sec. IV A) and boron nitride (sec. IV B) allotropes, respectively. Subsequently a comparison between results obtained for carbon and boron nitride will be drawn in sec. IV C. Sec. IV D employs the calculated ground state energies on the level of CCSD(T) theory and the DFT results to predict the pressure-temperature phase diagrams of carbon and boron nitride. In sec. IV E we review experimentally observed phase transitions and compare to the produced theoretical results. Section IV F focuses on the hexagonal form of diamond.

#### A. Carbon allotropes

Fig. 8 depicts the electronic ground state energy differences ($\Delta E = E_{c-D} - E_{G-ABC}$) between carbon diamond (c-D) and graphite (G-ABC) obtained using a range of DFT and quantum chemical wavefunction based theories. Experimentally G-AB is the most stable form of graphite. However, G-ABC and G-AB are degenerate to within a few meV per atom. Positive and negative energy differences in Fig. 8 indicate the thermodynamic stability of graphite and diamond, respectively. We stress that these calculations employ the DFT-LDA relaxed structures and that further relaxation effects of the respective functionals are not taken into account. Further relaxation effects can be significant for functionals that fail to describe the interlayer binding in graphite. However, this section will focus on benchmarking the accuracy of the employed functionals for a fixed geometry only. In passing we note, however, that for comparison we have repeated the DFT calculations summarized in Fig. 8 using geometries relaxed on the level of the PBE+MBD functional. The corresponding energy differences did not change by more than five percent as a result of the small changes in the employed geometries.

The grey bar in Fig. 8 corresponds to the experimental estimate of the ground state energy difference corrected for zero-point vibrational energies (ZPVEs). The experimental value of the difference in the Gibbs energy between graphite and diamond is 25 meV/atom$^{-1}$ and has been obtained from the heat of combustion and extrapolation to 0 K using the heat capacity [48]. Therefore the latter value includes ZPVEs that stabilize graphite compared to diamond. To allow for a direct comparison between experiment and theory we have re-
B3LYP hybrid functional does not follow this trend and a few 10 meV atom$^{-1}$, overestimating the stability of graphite compared to experiment by further improvement compared to SCAN, overestimating energy differences of 52 meV atom$^{-1}$ by including dispersion effects on the level of MBD or by switching to the SCAN functional, yielding reduced by including dispersion effects on the level of perturbative MBD (MBD) and 94 meV atom$^{-1}$ (SCAN), respectively. Furthermore hybrid functionals such as PBE0 or HSE06 constitute a further improvement compared to SCAN, overestimating the stability of graphite compared to experiment by a few 10 meV atom$^{-1}$ only. However, we note that the B3LYP hybrid functional does not follow this trend and gives the worst agreement with experiment out of all theories considered in the present study. Therefore a systematic improvability of the employed XC functionals with respect to their rung and computational cost can not be achieved in the present case. Furthermore, one conclusion of the above findings is that non-van der Waals corrected higher-level functionals (PBE, SCAN, PBE0 and HSE06) predict the graphitic phase to be more stable than diamond, whereas the inclusion of van der Waals corrections (MBD) can reverse their ordering. Indeed we note in passing that in contrast to HSE06, HSE06+MBD predicts c-D being more stable than G-ABC by 30 meV atom$^{-1}$.

We now turn to the discussion of the results obtained using wavefunction based theories as depicted in Fig. 8. HF predicts graphite to be more stable than diamond by approximately 100 meV atom$^{-1}$, albeit neglecting dispersion effects that play an important role in the interlayer binding of graphite. We note that due to the neglect
of these contributions, HF would predict the isolated graphene sheets to be more stable than graphite. Second-order Møller-Plesset (MP2) perturbation theory corresponds to the next level of wavefunction based method and predicts diamond to be slightly more stable than graphite by approximately 19 meV atom\(^{-1}\). However, the finite size errors of the obtained MP2 results are significant as indicated by the box plot, which is described in Sec. III B 1. The \(k\)-point mesh convergence using CCSD-TA-FS theory is much faster compared to MP2-TA-FS theory as indicated by the smaller error bar. We find that CCSD-TA-FS theory predicts diamond to be more stable than graphite by 31 meV atom\(^{-1}\). Including the perturbative triples contribution to CCSD-TA-FS theory yields an even better agreement with experiment albeit predicting diamond to be slightly more stable than graphite by 14 meV atom\(^{-1}\). In passing we note that DMC has been used in Ref. [89] to predict an energy difference in almost perfect agreement with experiment, whereas the random-phase approximation (RPA) predicts both allotropes to be exactly degenerate [90]. The good agreement between DMC and experiment is partly fortuitous due to remaining errors from the stochastic sampling, the fixed-node approximation and the employed pseudopotentials. However, the agreement between the high-level methods such as wavefunction based theories DMC and CCSD(T), the RPA results and experiment to within a few ten meV atom\(^{-1}\) is encouraging. The remaining finite size and basis set errors in CCSD(T) theory calculations do not allow for predicting which carbon allotrope is more stable, although we can conclude that they are expected to be degenerate to about 10–20 meV atom\(^{-1}\) including ZPVE. Our findings indicate that quantum chemical wavefunction based theories allow for a systematic improbability of the predicted energy differences as one increases the level of theory ranging from HF, MP2, CCSD to CCSD(T).

Having demonstrated that CCSD(T) theory is expected to yield accurate energy differences for the thermodynamically most stable carbon allotropes we now seek to investigate the pressure-driven transition pathways introduced in Sec. II B. To this end we focus on the activation barrier height that is defined as the difference in the electronic ground state energy between graphite and the corresponding transition state \(E_A = E_{\text{TS}} - E_{G-ABC}\). The considered transition states are referred to as pc-TS, pw-TS, bw-TS and l-pc-TS. The activation barriers can not be compared to experimental observations directly but serve as theoretical benchmark systems and qualitative models for realistic phase transitions. Fig. 9 depicts the difference of calculated activation barrier heights between various methods and CCSD(T) (including finite size corrections); for example, \(\Delta E_A^{\text{LDA}} = E_A^{\text{LDA}} - E_A^{\text{CCSD(T)-TA}}\). The depicted results confirm well-known trends for the accuracy of DFT methods, assuming that CCSD(T) theory can be considered an accurate benchmarking reference for the activation barrier height. LDA underestimates the activation barrier heights for all investigated transition states by 50–100 meV atom\(^{-1}\), showing that this level of theory suffers from larger errors in the description of XC energies for transition states compared to initial and final states (G-ABC and c-D). Including gradient corrections on the level of the PBE functional improves the agreement with CCSD(T) theory noticeably, yielding overestimated activation energies with errors smaller than 50 meV atom\(^{-1}\) for all four transition states. This is in contrast to PBE results for molecular activation barrier heights in the gas phase that are in general underestimated [10]. However, we believe that the overestimation of the PBE barriers for the studied solids is caused by the neglect of interatomic van der Waals forces, which play an important role for the present systems. We stress that the inclusion of dispersion effects to PBE on the level of PBE+MBD theory yields again underestimated activation energies that agree with CCSD(T) to within 50 meV atom\(^{-1}\). The SCAN functional yields barrier heights that are almost identical to our PBE findings. Furthermore the inclusion of non-local exchange in the PBE0 and HSE06 hybrid functionals yields on average slightly larger barrier heights. We note that adding the MBD effect (from the difference between PBE and PBE+MBD calculations) to these hybrid functionals would yield barrier heights in almost perfect agreement with CCSD(T) theory. On the other hand, we find that the B3LYP hybrid functional yields overestimated barrier heights, exhibiting errors on a scale of more than 100 meV atom\(^{-1}\).

We now turn to the discussion of activation barrier heights calculated using wavefunction based theories starting with the HF method. Our findings are depicted in Fig. 9 and show that HF yields strongly overestimated barrier heights with errors on the scale of almost 300 meV atom\(^{-1}\) compared to CCSD(T). This trend is known from molecular quantum chemistry and can be explained by the fact that HF neglects electronic correlation effects, which are in general larger in the tran-

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**FIG. 8.** Energy difference between c-D and G-ABC (\(\Delta E = E_{c-D} - E_{G-ABC}\)) compared with DMC [89] and experimental values with a freely chosen error of ±10 meV atom\(^{-1}\) (gray bar) [48] excluding 9 meV atom\(^{-1}\) ZPVE. See text for further details.
10

FIG. 9. Energy difference between G-ABC and transition state (TS) compared to CCSD(T)-TA referred as 0 meV atom$^{-1}$. Outlier marked with a “+” and dotted line connects energies from $k$-meshes with only one $k$-point along one direction. See text for further details.

sition state compared to the initial and final state of most chemical reactions. Accounting for electronic correlation effects on the level of MP2 theory corrects for this tendency although it yields underestimated barriers by about 100–200 meV atom$^{-1}$ for all transition states. We note that the box plots of the MP2 results in Fig. 9 also indicate that the remaining finite size errors for these estimates are significant. We attribute this to the observation that some transition states exhibit a metallic character in DFT calculations (l-pc-TS and bw-TS) and that MP2 theory suffers from severe shortcomings in metals such as $k$-point mesh divergence [91]. CCSD results for the barrier heights constitute a substantial improvement over MP2 findings, exhibiting errors compared to CCSD(T) that are smaller than 50 meV atom$^{-1}$. Furthermore we note that the box plot for CCSD-TA-FS results is significantly smaller, indicating that the remaining finite size errors are below a few 10 meV atom$^{-1}$. From these findings we conclude that quantum chemical wavefunction based theories including MP2 and CC theories have the potential of achieving results for activation barrier heights in solid-solid phase transitions with systematically improvable accuracy.

Having discussed the accuracy of DFT and wavefunction based methods for predicting the activation barrier heights, we now seek to address the question: which transition states are energetically the most favorable? This is an important question because it affects through which transition state a pressure-driven phase transition proceeds and which (meta-)stable carbon allotrope will be the outcome. Fig. 10 depicts the energy differences of the activation barrier heights with respect to the pc-TS for the respective electronic structure theories. Unequivocally all theories predict the pc-TS to be the energetically most favorable transition state, implying that the puckering mechanism is expected to play the most important role in pressure-driven graphite to diamond transitions. In passing we note that our LDA results are comparable to previous work [68, 69] and that the energy difference between the boat and chair conformation of graphane is 55 meV atom$^{-1}$, favoring the chair conformation [92]. As regards the ordering of the remaining transition states (bw-TS, pw-TS, and l-pc-TS), we find that the DFT methods shown in Fig. 10 predict all very similar orderings. bw-TS is energetically the least favorable transition state, whereas pw-TS and l-pc-TS agree to within a few meV per atom, except for the PBE, SCAN and B3LYP functionals that predict the pw-TS to be slightly more favorable in energy than the l-pc-TS. In the case of results obtained using wavefunction based methods depicted in Fig. 10 we find that the bw-TS corresponds to the largest activation barrier height and that pw-TS and l-pc-TS agree to within the remaining finite size errors. However, MP2 theory deviates from this trend by predicting equally large activation barrier heights for the bw-TS and l-pc-TS, making the pw-TS the second most favorable transition state. However, we stress that MP2 results are perhaps not meaningful due to the metallic character of some transition states. From the above re-
Positive and negative energy differences in Fig. 11 indicate that the low- ($h$-BN / $r$-BN) and high-density ($c$-BN) phase is predicted to be more stable, respectively. Grey bars show experimental findings. Solozhenko et al. predict $c$-BN to be more stable than $h$-BN by 78 meV atom$^{-1}$ at 0 K (including ZPVE) [51]. This result was obtained from fluorine combustion and extrapolation to 0 K using the heat capacity. Other recent experiments obtain the equilibrium phase boundary directly from catalytic transitions with X-ray diffraction analysis [61], finding that $h$-BN is more stable even at 0 K. To allow for a comparison between experiment and theory we have removed 4 meV atom$^{-1}$ ZPVE from the experimental estimates in Fig. 11 estimated using LDA.

We find that DFT-LDA calculations predict $c$-BN to be more stable than $r$-BN with an energy difference of $-56$ meV atom$^{-1}$ in good agreement with results from Ref. [96]. Including gradient corrections on the level of the PBE functional reverses their order and yields an energy difference of 64 meV atom$^{-1}$. However, it is known that van der Waals interactions have to be taken into account for an accurate description of electronic correlation effects especially in layered compounds. Inclusion of MBD on top of the PBE functional allows for capturing such correlation effects. Compared to PBE, PBE+MBD reverses the order between both allotropes again, yielding an energy difference of $-13$ meV atom$^{-1}$. The SCAN functional goes beyond the GGA and is expected to perform better than PBE [5]. Our findings shown in Fig. 11 reveal that SCAN reduces the energy difference between $c$-BN and $r$-BN compared to PBE from 64 meV atom$^{-1}$ to 47 meV atom$^{-1}$. The hybrid functionals PBE0 and HSE06 continue this trend and predict $r$-BN to be more stable than $c$-BN with an even smaller energy difference of 30 meV atom$^{-1}$ and 31 meV atom$^{-1}$, respectively. However, the B3LYP functional significantly overestimates the stability of $r$-BN, predicting a difference of 160 meV atom$^{-1}$. The different results from the various DFT methods make it difficult to provide a firm conclusion on the true energy difference. However, we believe that the results allow for a similar conclusion as for the case of carbon allotropes: higher-level functionals (PBE, SCAN, PBE0 and HSE06) predict the graphitic phase to be more stable than the diamond-like phase but the inclusion of van der Waals corrections can reverse their ordering.

We now turn to the discussion of results for the energy difference between $c$-BN and $r$-BN obtained using wavefunction based methods as depicted in Fig. 11. HF theory, disregarding electronic correlation effects, substantially overestimates the stability of $r$-BN compared to $c$-BN. The HF energy difference is the largest of all considered methods (222 meV atom$^{-1}$). The simplest treatment of electronic correlation effects on the level of wavefunction based methods is achieved using MP2 theory, predicting $c$-BN to be more stable than $r$-BN by 58 meV atom$^{-1}$. However, as indicated by the large error bars we find that the remaining finite size errors
on the level of MP2 theory are on the scale of several 10 meV atom$^{-1}$. CCSD-TA-FS predicts r-BN to be more stable than c-BN by 14 meV atom$^{-1}$ and is well converged with respect to the employed $k$-mesh. The inclusion of perturbative triples yields an energy difference of 2 meV atom$^{-1}$ only. From these findings we conclude that the series of wavefunction based theories ranging from HF, MP2, CCSD to CCSD(T) predicts an oscillating but convergent energy difference between r-BN and c-BN that is close to zero on the level of CCSD(T) theory. Due to the remaining finite size errors that are estimated to be on the scale of approximately 10 meV atom$^{-1}$, we conclude that r-BN and c-BN are degenerate to within 10 meV atom$^{-1}$ on the level of CCSD(T).

We now seek to investigate the pressure-driven transition pathways introduced in Sec. II B for boron nitride. The discussion and results are analogue to our findings for carbon. We investigate again the activation barrier height that is defined as the difference in the electronic ground state energy between h-BN and the corresponding transition state $E_A = E_{TS} - E_{h-BN}$. Fig. 12 depicts the difference in activation barrier heights between various methods and CCSD(T) (including finite size corrections); for example, $\Delta E_{A,FDA} = E_{A,FDA} - E_{A,CCSD(T)-TA}$. LDA underestimates all activation barrier heights by approximately 50 meV atom$^{-1}$. Including the effect of gradient corrections on the level of the PBE functional improves the agreement with CCSD(T) theory slightly, yielding overestimated activation energies with average errors of roughly 25 meV atom$^{-1}$ for all four transition states. This is in contrast to PBE results for molecular activation barrier heights in the gas phase that are in general underestimated. However, we believe that the overestimation of the PBE barriers for the studied solids is caused by the neglect of interatomic van der Waals forces in the same manner as for carbon. We stress that the inclusion of dispersion effects to PBE on the level of PBE+MBD theory yields again underestimated activation energies that agree with CCSD(T) to within approximately 25 meV atom$^{-1}$. Moving to the next level of theory, we find that the SCAN functional yields barrier height results that are slightly better than our PBE findings. The PBE0 and HSE06 hybrid functionals yield barrier heights similar to SCAN. However, the B3LYP hybrid functional overestimates barrier heights substantially, exhibiting errors on a scale of more than 100 meV atom$^{-1}$.

We now turn to the discussion of activation barrier heights calculated on the level of wavefunction based theories, starting with the HF method. Our findings are depicted in Fig. 12 and show that HF yields significantly overestimated barrier heights with errors ranging from 150 meV atom$^{-1}$ to 200 meV atom$^{-1}$ compared to CCSD(T). Accounting for electronic correlation effects on the level of MP2 theory corrects for this tendency partly despite yielding significantly underestimated barriers on the scale of 50 meV atom$^{-1}$ and suffering from non-negligible finite size errors as indicated by the error bars. CCSD results for the barrier heights constitute a substantial improvement over MP2 findings, overestimating the barriers by about 25 meV atom$^{-1}$. From these findings we conclude again that quantum chemical wavefunction based theories including MP2 and CC theories have the potential of achieving results for activation barrier heights in solid-solid phase transitions with systematically improvable accuracy. However, their finite size errors are a dominant source of error in our present calculations.

Having discussed the accuracy of DFT and wavefunction based methods for predicting the activation barrier heights, we now seek to address the question which transition states are energetically the most favorable for boron nitride allotropes. As for carbon this is an important question because it affects through which transition a pressure-driven phase transition proceeds and which (meta-)stable boron nitride allotrope will be the outcome. Fig. 13 depicts the energy differences of the activation barrier heights with respect to the pc-TS for all employed electronic structure theories. Unequivocally all theories predict the pc-TS to be the energetically most favorable transition state in the same manner as for carbon. As regards the ordering of the remaining transition states (bw-TS, pw-TS, and l-pc-TS), we find that all methods predict pw-TS to be the second most favorable transition state, whereas bw-TS and l-pc-TS have the largest barrier heights. An important conclusion from the above results is that all levels of theory agree qualitatively and predict similar energy differences between the barrier heights. However, we note in passing that our MP2 and CCSD results exhibit very large finite size errors for the bw-TS.

All the energies discussed in this section are summarized in Tab. IV. Furthermore we note that all employed methods predict w-BN to be less stable than c-BN by

| System                  | LDA  | PBE  | PBE+MBD | SCAN | PBE0 | B3LYP | HSE06 | HF    | MP2-TA-FS | CCSD-TA-FS | CCSD(T)-TA |
|-------------------------|------|------|---------|------|------|-------|-------|-------|-----------|------------|------------|
| c-D – G-ABC            | −12  | 121  | 52      | 94   | 45   | 206   | 49   | 110   | −19 ± 29  | −19 ± 29   | −31 ± 5    | −14        |
| h-D – c-D              | 25   | 24   | 25      | 29   | 27   | 30    | 27   | 37    | 30 ± 14   | 30 ± 14    | 35 ± 10    | 33         |
| pc-TS – G-ABC          | 329  | 423  | 372     | 422  | 437  | 530   | 435  | 671   | 210 ± 46  | 426 ± 1    | 385        |
| bw-TS – G-ABC          | 395  | 488  | 438     | 495  | 512  | 604   | 509  | 772   | 334 ± 49  | 529 ± 20   | 481        |
| pw-TS – G-ABC          | 362  | 452  | 404     | 452  | 474  | 560   | 470  | 710   | 238 ± 61  | 469 ± 14   | 423        |
| l-pc-TS – G-ABC        | 307  | 470  | 413     | 473  | 473  | 581   | 470  | 685   | 350 ± 41  | 469 ± 16   | 441        |
FIG. 12. Energy difference between h-BN and transition state (TS) compared to CCSD(T)-TA referred as 0 meV atom$^{-1}$. Outlier marked with a "+' and dotted line connects energies with the smallest k-meshes. See text for further details.

TABLE IV. Energy differences in meV per atom for boron nitride structures as obtained by various levels of theory.

| System          | LDA | PBE  | PBE+MBD | SCAN | PBE0 | B3LYP | HSE06 | HF   | MP2-TA-FS | CCSD-TA-FS | CCSD(T)-TA |
|-----------------|-----|------|---------|------|------|-------|-------|------|-----------|-------------|-------------|
| c-BN – r-BN     | −56 | 64   | −13     | 47   | 30   | 160   | 31    | 222  | −58 ± 21  | 14 ± 11     | 2           |
| w-BN – c-BN     | 18  | 17   | 18      | 20   | 19   | 20    | 19    | 23   | 23 ± 7    | 24 ± 4      | 22          |
| pc-TS – h-BN    | 169 | 250  | 195     | 240  | 249  | 329   | 249   | 409  | 173 ± 17  | 243 ± 10    | 221         |
| bw-TS – h-BN    | 214 | 292  | 240     | 286  | 294  | 372   | 294   | 454  | 233 ± 42  | 299 ± 30    | 281         |
| pw-TS – h-BN    | 179 | 259  | 207     | 251  | 258  | 338   | 258   | 412  | 188 ± 18  | 256 ± 11    | 234         |
| l-pc-TS – h-BN  | 202 | 293  | 230     | 288  | 286  | 381   | 287   | 461  | 213 ± 32  | 283 ± 19    | 259         |

FIG. 13. Energy difference between pc-TS and other transition states. See text for further details.

about 20 meV atom$^{-1}$.

C. Comparing the carbon and boron nitride systems

As discussed in the previous sections and as summarized in Tab. III and IV, the produced results for carbon and boron nitride systems are very similar. Overall the employed electronic structure theories exhibit the same trends for the prediction of energy differences between (meta-)stable allotropes and barrier heights. However, one interesting exception is the energy difference between the high- (c-BN/c-D) and low-density (r-BN/G-ABC) phases. Comparing this difference for DFT based methods between boron nitride and carbon systems reveals that DFT tends to stabilize the high-density phases of boron nitride more than in the case of carbon. The opposite trend can be observed for wavefunction based methods with the exception of MP2 theory. This trend is most evident when comparing LDA (BN: c-BN− r-BN= −56 meV atom$^{-1}$; C: c-D−G-ABC= −12 meV atom$^{-1}$) to HF (BN: c-BN− r-BN= 222 meV atom$^{-1}$; C: c-D−G-ABC= 110 meV atom$^{-1}$). Improving upon the respective rung of theory makes this trend less pronounced as can be seen for HSE06 (BN: c-BN− r-BN= 31 meV atom$^{-1}$; C: c-D−G-ABC= 49 meV atom$^{-1}$) and CCSD(T) (BN: c-BN− r-BN= 2 meV atom$^{-1}$; C: c-D−G-ABC= −14 meV atom$^{-1}$). This shows that the true energy differences between low- and high-density phases of carbon and boron nitride systems are perhaps on the scale of ten meV per atom, indicating that their respective phase diagrams are very similar.

Moreover we note that the barrier heights in carbon are approximately 180 meV atom$^{-1}$ larger than for boron nitride which can be attributed to the stronger covalent bond in carbon allotropes.
D. Phase diagrams and transition probabilities

We now turn to the discussion of pressure-temperature phase diagrams for carbon and boron nitride. The calculated phase diagrams are obtained from the Gibbs energies \( G \) as defined in Eq. 1. Furthermore the Gibbs energy of the transition states \( G_{TS}(T, P) \) allows for determining approximate phase transformation probabilities as defined by Eq. 4.

Fig. 14 depicts the pressure-temperature phase diagrams for carbon and boron nitride. The experimental equilibrium phase boundaries are shown by dotted lines and separate the low-density graphitic phases (\( h\)-BN and G-AB) at lower pressures and higher temperatures from the high-density diamond-like phases (\( c\)-BN and \( c\)-D) at higher pressures and lower temperatures. In the case of carbon, the two experimental phase boundaries deviate from each other only at temperatures higher than 2000 K \cite{87, 93}. Furthermore both experimental results predict graphite being the stable carbon allotrope at ambient conditions. This is in contrast to boron nitride where the experimental equilibrium phase boundaries disagree by several GPa at lower temperatures \cite{51, 61}. The experimental findings by Solozhenko et al. \cite{51} and Fukunaga et al. \cite{61} even differ in their prediction of the thermodynamically most stable allotrope at ambient conditions. Solozhenko et al. predict the zinc blende phase, whereas Fukunaga et al. find the \( h\)-BN phase being more stable.

Fig. 14 also shows the calculated equilibrium phase boundaries obtained using DFT on the level of the LDA. In the case of carbon, the LDA results agree with both experimental findings to within a few GPa at low temperatures. At temperatures higher than 2000 K the LDA boundary is closer to the Berman-Simons line. However, we stress that LDA results are less accurate for higher temperatures. At temperatures higher than 2000 K \cite{87, 93}.

Furthermore both experimental results predict graphite being the stable carbon allotrope at ambient conditions. This is in contrast to boron nitride where the experimental equilibrium phase boundaries disagree by several GPa at lower temperatures \cite{51, 61}. The experimental findings by Solozhenko et al. \cite{51} and Fukunaga et al. \cite{61} even differ in their prediction of the thermodynamically most stable allotrope at ambient conditions. Solozhenko et al. predict the zinc blende phase, whereas Fukunaga et al. find the \( h\)-BN phase being more stable.

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Experimental phase diagram of BN is at 1500 K and that of carbon at 2000 K [51, 54]. The contour lines reveal an important problem in the synthesis of cubic diamond-like phases. When starting from a graphite-like phase a diamond-like phase can be obtained by applying a minimum pressure, which increases strongly at lower temperatures. However, once the metastable wurtzite phase is created or stacking faults and twins are present the kinetics is very different and much higher minimum temperatures are required to transform into the cubic phase or heal stacking faults. This explains the almost rectangular region in the experimental phase diagram for the cubic phase.

E. Revision of observed experimental phase transitions

We now summarize experimental findings of observed phase transitions in more detail. As an overview all ground and transition states are depicted in Fig. 15. The first row shows the graphite-like phases (except for w-BN in the last column), the transition states are in the middle row and the high-density phases in the bottom row. The green and black arrows correspond to the experimentally observed transitions of boron nitride and carbon, respectively. A detailed discussion is provided in the following subsections.

1. Transitions between low and high density carbon phases

Since graphite occurs in the AB stacking there is no direct transition pathway to c-D without introducing shear stress. This is illustrated by the yellow lines of Fig. 15b and 15i, which are not parallel to each other. Shear stress can be reduced by twin formation or stacking faults parallel to (111)\textsubscript{\text{c}}, as can be observed by high resolution TEM [101]. The stacking faults can be interpreted as a so-called “wave-like buckling and slipping” mechanism [102]. This could change the stacking order from AB to ABC and would make the pc-TS accessible, reducing the activation energy since the stacking order from AB to ABC would make the c-axis of graphite, inducing a so-called “wave-like buckling and slipping” mechanism [102].

Experimental observations of the transformation from G-AB to c-D suggest a structural relationship with plane (001)\textsubscript{G} \parallel (112)\textsubscript{c} and direction [10\overline{1}0]\textsubscript{G} \parallel [111]\textsubscript{c} [46]. The same orientation has been found in an MD simulation [99]. This orientation fits exactly to the combination of Fig. 15b (G-AB) $\xrightarrow{15f}$ 15j/15d $\xrightarrow{15h}$ 15i (c-D). If graphite is compressed to over 20 GPa at room temperature the resistance increases, especially perpendicular to the c-axis and returns to its original value after pressure is released [103, 104]. For temperatures between 800 °C and 1600 °C a mixture of h-D and c-D is formed [76]. With a minimum pressure of 12 GPa and temperatures beyond 1800 °C the portion of h-D decreases and only c-D is left [54]. Beyond 1800 °C the temperature is high enough to reduce shear stress and to break bonds to heal...
FIG. 15. Structures for different phase transition pathways: (a) \( r\text{-BN} \xrightarrow{(e) pc\text{-TS}} (i) \text{(BN: [72]; C: [67])} \), (b) \( \text{BN-AB} \xrightarrow{(f) bw\text{-TS}} (j) \text{(BN: [44]; C: [69])} \), (c) \( h\text{-BN} \xrightarrow{(g) pw\text{-TS}} (k) \text{(BN: [72]; C: [68])} \), (d) \( w\text{-BN} \xrightarrow{(h) l-pc\text{-TS}} (l) \text{(C: [69])} \). Red arrows indicate the atomic displacements and support together with the atom numbers the assignment during the phase transition. Dotted lines show new bonds to be formed and red lines represent strong interaction during the transition state. Black (carbon) and green (BN) arrows represents possible transitions (See text for further information). The crystal structures for C can easily be derived from BN, by substituting all B- and N-atoms with C-atoms. This makes \( r\text{-BN} \), \( \text{BN-AB} \), \( h\text{-BN} \), \( c\text{-BN} \) and \( w\text{-BN} \) equivalent to G-ABC, G-AB, G-AA, c-D and h-D, respectively.

without stacking faults. This is in good agreement with the calculated transformation probability contour lines depicted in Fig. 14. h-D can be considered an intermediate structure and can be left out in the conversion from G-AB to c-D by slightly different carbon displacements G-AB \( l-pc\text{-TS} \) (Fig. 15b) \( \xrightarrow{(h) l-pc\text{-TS}} \) c-D. The bw-TS is higher in energy than the l-pc-TS and the latter one would be preferred. This means the h-D would not be created from the energetic point of view. However, the latter mechanism is only probable if one accounts for the induced shear stress in some manner.

G-AB (Fig. 15b) can transform into h-D without introducing shear stress. The structure relation is the one via bw-TS (Fig. 15f) as proposed by Bundy and Kasper [103] and confirmed many times [42, 46, 76].

The activation energies for the carbon system are higher than for boron nitride and therefore require higher pressures and larger temperatures to observe. For the catalytic transition at the phase boundary the activation energy in solution is about 1.3–1.7 eV [54]. How-
ever, this is not comparable with the calculated solid-solid phase transitions from this work since the carbon dissolves within the liquid catalyst.

2. Transitions between low and high density BN phases

For the boron nitride system the direct transitions via pc-TS and pw-TS are the most probable because the stable graphite-like structures exhibit an ABC (r-BN) and AA’ (h-BN) stacking and these transition states have the lowest activation energies of those calculated in the present work. Experimentally direct conversion of h-BN→w-BN and r-BN→c-BN have been observed. Experimental observations of structural orientation relationships of initial and final states can be used to infer which transition states are possible. Experimentally the transitions r-BN→c-BN (Fig. 15a) with plane (001)_r∥(111)_c and direction [210]_r∥[110]_c) and h-BN→w-BN (Fig. 15c) with plane (001)_w∥(001)_w and direction [100]_w∥[101]_w has been observed during shock wave synthesis [47]. Due to the different stacking sequence of G-AB and h-BN the structure relationship to h-D and w-BN is different [105]. The 90° rotation can be seen by comparing G-AB (Fig. 15b) with h-BN (Fig. 15j) with plane (001)_w∥(001)_w and direction [101]_w∥[101]_w has been observed during shock wave synthesis [47]. The value for shock wave synthesis is even higher: 8.7 eV atom⁻¹ [56]. Since all these values for the w-BN→c-BN reaction are obtained from non equilibrium conditions, they can not be compared with the calculated ones. Obtained activation energies are influenced by kinetic effects, which arise from gran size, defects or other structural distortions [58]. Therefore these values are rather upper bounds than real activation energies. If these were real activation energies, the backward reaction would have a similar activation energy. However, this is not true as can be seen by comparing with the w-BN→h-BN reaction with an experimental activation energy of 0.22 eV atom⁻¹ [60]. The latter value is comparable with the calculated one at the same experimental conditions (∼0.17 eV atom⁻¹ at 900°C).

3. Transition between low density graphite-like BN phases

The r-BN has an ABC stacking and could be converted to AB or AA trough translation because the layers are only shifted towards each other (Fig. 1b and 1c). h-BN has an AA’ stacking, where each layer is rotated by 60° towards their neighboring layers and a translation would never create an AA, AB or ABC stacking (Fig. 1). To reach bw-TS and pw-TS pressure has to be applied. The transformation from r-BN→h-BN occurs only at elevated pressure and by applying shear stress [45], which could be a sign, that the layers have first to be changed into the AD stacking order and buckle before they can transform into each other. One possible transition path could be similar to Fig. 15a with 15b by changing the stacking sequence from ABC to AD and result in the bw-TS or close to it. It can switch to the pw-TS (15g) by changing some bond lengths. This changes the orientation of the c-axis and the new c-axis has an AA’ stacking order and after de-puckering and elongation along this new c-axis the h-BN structure is formed.
F. Hexagonal diamond (Lonsdaleite)

While the synthesis of high quality wurtzite boron nitride crystals is possible, the existence of the corresponding carbon polymorph known as lonsdaleite has recently been called into question [70]. A number of studies have investigated the wurtzite structure of carbon and shown the main XRD peaks of the cubic phase are also part of the XRD spectra of the wurtzite phase, making it difficult to distinguish these phases. Ignoring the relative intensities of the peaks maybe due to textures effects, makes it impossible to determine the exact amount of the cubic phase within the wurtzite phase. The latter problem occurred in older publications about diamond, where the detected lines only have been published without the XRD spectra [42, 103]. The largest peak of the h-D spectra which is not part of the c-D spectra has a d-spacing of 2.18 Å and is just a shoulder of the main peak with a d-spacing of 2.06 Å. In a recent publication of XRD spectra for a natural and a synthetic sample the h-D peaks are just shoulders and the sample is mainly c-D [70]. The authors point out that the peaks are due to stacking faults of basal planes and twinning, which are supported by STEM images. The defects create new planes with different d-spacing compared to cubic single crystals and have the same spacing like the ones in h-D because these planes are in the wurtzite structure. Therefore the amount of h-D can be assigned to the amount of defects. Transforming G-AB $\xrightarrow{pc-TS} c$-D implies to produce shear stress due to changing the stacking order from AB to ABC, as already pointed out by Tateyama [69]. By forming a (111) twin the stacking order inverts and the stress is reduced. The formation of one type of twins can be seen in an MD simulation [99]. The twin planes where the stacking order inverts corresponds to the h-D structure. This is a clear example why h-D can not by synthesized as a single crystal easily. Yoshiasa et al. observed a higher ratio of h-D to c-D for X-ray diffraction profiles perpendicular to the c-axis of the parent graphite [76]. This supports that h-D stacking is not produced by the pw-TS, but via bw-TS because of the orientation of the c-axis. The pressure and temperature region observed by Bundy and Kasper [103] for the formation of h-D has been investigated by other groups and not all were able to synthesize a detectable amount of h-D [41], which was attributed it to the different experimental conditions.

The same type of defects as described above were also observed in the BN system [44]. However, in contrast to h-D, w-BN can be synthesized relatively pure [39]. We attribute this to the existence of a stable graphite-like (h-BN) structure together with a relatively low transition state (pw-TS) that forms directly the wurtzite phase without inducing shear stress. In the carbon system the graphite-like phase with the same stacking order as h-D (G-AA) does not exist. G-AB needs to shift individual sheets to reach the bw-TS or 1-pc-TS. This is very unlikely to happen and would also require larger temperatures. Since h-D and w-BN are meta-stable structures an increase in temperature and pressure will always lead to a transformation into the thermodynamically stable c-D and c-BN structures.

V. CONCLUSION

In this work we have investigated (meta-)stable boron nitride as well as carbon allotropes for a range of pressures and temperatures. Furthermore corresponding concerted transition pathways have been explored. The calculations were performed using a selection of approximate exchange and correlation density functionals and quantum chemical wavefunction based theories including the coupled cluster method. A comparison between the theoretical and experimental findings reveals that highly accurate predictions for equilibrium phase boundaries constitute a true challenge for state of the art electronic structure theories.

We have investigated the energy differences between low- and high-density phases of carbon and boron nitride. Due to the variation in the results obtained using LDA, GGA, mGGA and hybrid functionals a firm conclusion and accurate estimate of the energy differences can not be achieved. Furthermore the explicit inclusion of van der Waals interactions on the level of MBD is found to be significant and might change the order of the predicted stability depending on the employed parent XC functional. We stress that considering other approximations to the van der Waals interactions or additional XC functionals would not allow for achieving more reliable results. On the other hand we find that quantum chemical wavefunction based theories allow for a systematic improvability of the obtained results. HF, MP2, CCSD and CCSD(T) methods yield an oscillating but convergent estimate of the calculated energy differences. We note in passing that such a systematic behaviour was recently also reported for calculated transition pressures in LiH crystals [108]. The CCSD(T) method predicts that the corresponding low- and high-density phases of boron nitride as well as carbon are degenerate to about 10–20 meV atom$^{-1}$ including ZPVEs. We stress that the remaining uncertainty of coupled cluster theory results is dominated by finite size effects that can possibly be further reduced in future by studying larger systems. We also note that finite size errors are significantly larger for results obtained using second-order Møller-Plesset perturbation theory (MP2) in particular for small gap systems, where MP2 is considered less accurate. The present coupled cluster theory results for the energy difference between carbon diamond and graphite are in agreement with experimental measurements and quantum monte carlo calculations from literature to within about 20–30 meV atom$^{-1}$, which corresponds to the accuracy that is typically ascribed to CCSD(T) theory. The same conclusion can not be drawn for boron nitride due to a larger spread in the available experimental findings. However,
we hope that this work will motivate further calculations using quantum monte carlo methods and experimental studies to help providing more accurate estimates of the corresponding equilibrium phase boundaries.

The obtained coupled cluster theory results for the activation barrier heights in the graphitic to diamond-like transitions of boron nitride as well as carbon also allow for benchmarking different levels of approximate exchange and correlation density functionals. We conclude that the accuracy of the employed LDA, GGA and mGGA functionals follows roughly the same trends as for activation barrier heights in molecular gas phase reactions: LDA, GGA and mGGA functionals underestimate the barrier heights if the effect of van der Waals interactions is taken into account. Furthermore the results for hybrid functionals indicate a strong dependence on the choice of parametrization. We find that PBE0 and HSE06 yield significantly more accurate results than B3LYP, confirming previous findings for a wide range of solids [109]. Furthermore we note that the investigated transition states are not very strongly correlated as indicated by the good agreement of a few 10 meV/atom−1 between CCSD and CCSD(T) theory. The observed finite size effects are larger for the predicted coupled cluster barrier heights than for the energy differences of the (meta-)stable allotropes. An important conclusion for the investigated transition states is that their ordering and relative stabilities is mostly independent from the employed electronic structure theory. All employed theories predict unequivocally that the puckering mechanism as present in the pc-TS is energetically the most favorable transition mechanism for boron nitride as well as carbon.

The prediction of pressure-temperature phase diagrams requires the calculation of Gibbs energies. We have shown that approximating the CCSD(T) Gibbs energy using the CCSD(T) energies of the (meta-)stable and transition states only and the LDA for its temperature and pressure dependence yields reliable pressure-temperature phase diagrams. The obtained phase boundaries agree with experimental results of carbon to within about one GPa at temperatures below 2000 K. In the case of boron nitride we find a similarly good agreement with a recently obtained experimental result of Fukunaga et al. in Ref. [61]. Furthermore we have provided estimates of approximate phase transition probabilities in a similar manner. The calculated phase transition probabilities confirm trends in the measured pressure and temperature dependence of experimentally observed phase transitions.

Finally we have addressed the conversion of graphite to hexagonal diamond also known as lonsdaleite using the obtained results for transition and (meta-)stable states. In the context of the present work it is reasonable to ask the question: why can the wurtzite phase of boron nitride be synthesized as an almost pure powder whereas the existence of single crystals of lonsdaleite is still under debate? We conclude that the puckering mechanism for the corresponding phase transitions is always the most probable due to its energetically more favorable transition state. However, the stacking of the parent graphitic phase that is put under pressure has a significant influence on the kinetics of the phase transition. We note that cubic diamond and wurtzite structures exhibit an ABC and AA′ stacking, respectively. Experimentally w-BN is formed only when applying pressure to h-BN, which exhibits also an AA′ stacking. In the case of carbon the corresponding G-AA phase is not stable, making a transformation from G-AB or G-ABC to lonsdaleite only possible by introducing stacking faults or similar defects. This conclusion is in agreement with recent experimental work.

ACKNOWLEDGMENTS

This project has received funding from the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation program (grant agreement No 715594). The computational results presented have been achieved in part using the Vienna Scientific Cluster (VSC). Helpful discussions with Ali Alavi are gratefully acknowledged.
