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## Understanding vapor phase growth of hexagonal boron nitride†

Anja Sutorius, <sup>a</sup> René Weiβing, <sup>a</sup> Carina Rindtorff Pérez, <sup>a</sup> Thomas Fischer, <sup>a</sup> Fabian Hartl, <sup>a</sup> Nilanjan Basu, <sup>b</sup> Hyeon Suk Shin <sup>b,c</sup> and Sanjay Mathur <sup>a\*</sup>

Hexagonal boron nitride (*h*BN), with its atomically flat structure, excellent chemical stability, and large band gap energy (~6 eV), serves as an exemplary 2D insulator in electronics. Additionally, it offers exceptional attributes for the growth and encapsulation of semiconductor transition metal dichalcogenides (TMDCs). Current methodologies for producing *h*BN thin films primarily involve exfoliating multi-layer or bulk crystals and thin film growth *via* chemical vapor deposition (CVD), which entails the thermal decomposition and surface reaction of molecular precursors like ammonia boranes ( $\text{NH}_3\text{BH}_3$ ) and borazine ( $\text{B}_3\text{N}_3\text{H}_6$ ). These molecular precursors contain pre-existing B–N bonds, thus promoting the nucleation of BN. However, the quality and phase purity of resulting BN films are greatly influenced by the film preparation and deposition process conditions that remain a substantial concern. This study aims to comprehensively investigate the impact of varied CVD systems, parameters, and precursor chemistry on the synthesis of high-quality, large scale *h*BN on both catalytic and non-catalytic substrates. The comparative analysis provided new insights into most effective approaches concerning both quality and scalability of vapor phase grown *h*BN films.

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### Introduction

Even two decades after the discovery of graphene monolayers, van der Waals (vdW) materials continue to attract significant interest due to their unique two-dimensional structure and remarkable electronic characteristics.<sup>1–7</sup> Hexagonal boron nitride (*h*BN), with a band gap of approximately 6 eV, distinguishes itself among 2D materials due to its structural resemblance to graphene and exceptional physical and chemical attributes.<sup>8–13</sup> Comprising alternating B and N atoms in its honeycomb lattice held together by in-plane  $\sigma$ -bonds of  $\text{sp}^2$  hybridized boron and nitrogen orbitals.<sup>14</sup> In its bulk form, *h*BN demonstrates weak inter-layer interactions making it amenable to exfoliation using chemical<sup>15,16</sup> or physical methods.<sup>17</sup> However, owing to the difference in the electronegativity of boron (2.04) and nitrogen (3.04),<sup>18</sup> electron density tends to localize around nitrogen centers<sup>19</sup> favoring

thermodynamically preferred AA' stacking mode, which accommodates in-plane dipolar interactions.

Given its superior mechanical strength, chemical and thermal stability as well as optical transparency, *h*BN is versatile for numerous applications<sup>20–22</sup> Consequently, current challenges in *h*BN synthesis include direct deposition techniques that yield homogeneous films with uniform thickness, large domain size and high crystallinity. The objective is to avoid transfer processes known to induce mechanical defects.<sup>23</sup> While mechanical exfoliation and chemical transfer procedures produce *h*BN flakes in the tens of micrometers, they often contain notable interfacial impurities. Achieving precise thickness control in CVD grown thin films in the Ångström region remains elusive.

In recent years research has predominantly focused on several methods for producing hexagonal boron nitride synthesis (Fig. 1), notably using multi-source approaches ( $\text{NH}_3$  and  $\text{BCl}_3$ ),<sup>24,25</sup> atmospheric-pressure CVD using ammonia borane or borazine (AB,  $\text{NH}_3\text{BH}_3$ )<sup>26–28</sup> and low-pressure CVD using borazine ( $\text{B}_3\text{N}_3\text{H}_6$ ).<sup>29,30</sup> While convenient, the use of multi-source systems poses challenges in vapor phase growth due to increased contamination through undesired elements and secondary phases. Thus, the use of single source precursors containing both B and N in single molecular compounds are promising for better control over phase purity, morphology, and structure of deposited thin films.<sup>31–35</sup> Ammonia borane represents a suitable single source precursor for *h*BN synthesis<sup>36</sup> due to its pre-existing B–N bonds, non-toxicity and

<sup>a</sup>Institute of Inorganic and Materials Chemistry, Department of Chemistry, University of Cologne, Greinstraße 6, 50939 Cologne, Germany.

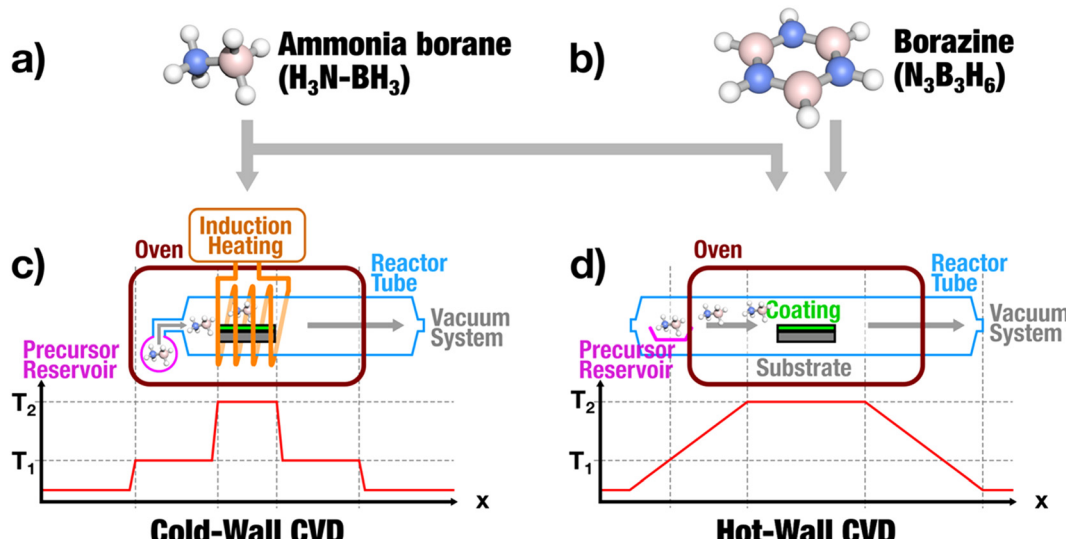
E-mail: [sanjay.mathur@uni-koeln.de](mailto:sanjay.mathur@uni-koeln.de)

<sup>b</sup>Center for 2D Quantum Heterostructures, Institute for Basic Science (IBS), Sungkyunkwan University (SKKU), Suwon 16419, Republic of Korea

<sup>c</sup>Department of Energy Science, Sungkyunkwan University (SKKU), Suwon 16419, Republic of Korea

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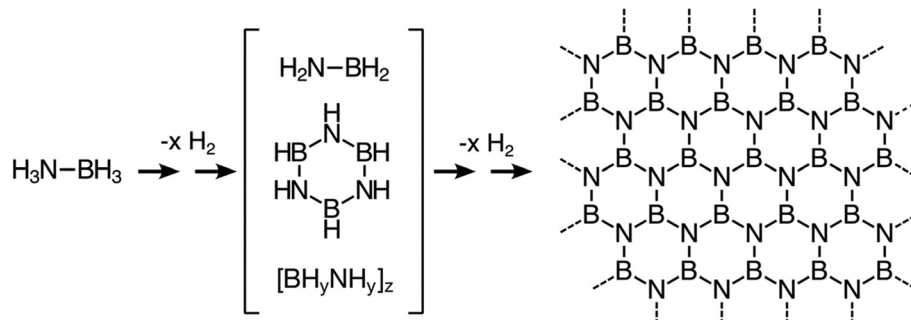
**Fig. 1** Schematic overview of different precursors: (a) ammonia borane ( $\text{NH}_3\text{BH}_3$ ) and (b) borazine ( $\text{N}_3\text{B}_3\text{H}_6$ ) and deposition systems where these precursors were used. Hot-wall CVD (c) and inductively heated cold-wall CVD (d) with different temperature profiles during the deposition of hexagonal boron nitride ( $T_1$ : sublimation temperature;  $T_2$ : decomposition temperature).

decomposition into benign by-products. Additionally, its cost-effectiveness, solid form and air stability makes it a favourable starting material.<sup>37–39</sup> However, the decomposition of ammonia borane is essentially a two-step process initially forming borazine before generating *h*BN (Fig. 2).

Moreover, the decomposition of ammonia borane yields not only borazine but also polymeric structures like polyamine borane ( $[\text{BH}_2\text{NH}_2]_x$ ) and polyimino borane ( $[\text{BHNH}]_x$ ), which makes the detection of borazine in the gas phase somewhat unpredictable. Interestingly, some calculations propose that the formation of polyimino borane is energetically more favourable than that of borazine.<sup>40</sup> Nonetheless, both byproducts contribute toward *h*BN formation. Borazine, also called “inorganic benzene” ( $\text{B}_3\text{N}_3\text{H}_6$ ), serves as another promising single source precursor for *h*BN growth due to pre-existing B–N bonds and possessing “ $\text{B}_3\text{N}_3$ ” hexagons that can exert a templating effect in the synthesis of BN monolayer. However, its utilization is constrained by several factors such as high costs,

toxicity, accessibility limitations that vary by country, and the requirement of specialized equipment for handling.

In most reports, catalytic substrates like copper,<sup>27,41</sup> nickel,<sup>42</sup> gold<sup>43</sup> and platinum,<sup>44</sup> have been employed. However, these substrates inevitably require transfer steps (bubble or etching transfer) for using *h*BN in electronic devices. The major hurdle in the use of *h*BN is related to lack of synthetic protocols enabling high-quality *h*BN on non-catalytic substrates, which would eliminate the need for additional transfer processes. Recent research demonstrates initial efforts toward direct deposition of *h*BN on sapphire and silicon substrates.<sup>45</sup> Singhal and co-workers showed promising outcomes using ammonia borane in a low-pressure CVD set-up by direct deposition of *h*BN on silicon (100) and sapphire wafer.<sup>46</sup> Despite achieving high-quality *h*BN compositions, the challenge persists in forming a continuous, ultra-thin film with a thickness in the range of a few nanometers or less as the thin film thickness can influence the functional properties.<sup>47</sup>



**Fig. 2** Schematic representation of the thermal decomposition pathway of ammonia borane into amino borane, borazine and polymeric compounds, subsequently yielding hexagonal boron nitride under  $\text{H}_2$  elimination.

In this study, our objective was to conduct a comprehensive analysis of the influence of precursor chemistry, deposition parameters and substrate nature on the quality and phase purity of deposited hexagonal boron nitride. Initially, we examined the thermal deposition of ammonia borane on copper substrates in an atmospheric pressure set-up. Drawing from this experience, similar experiments were conducted on non-catalytic substrates, specifically silicon dioxide and sapphire. Lastly, we demonstrate successful deposition of few layer *hBN* achieved through the use of borazine in a low pressure hot-wall CVD reactor.

## Results and discussion

Thin films of hexagonal boron nitride were grown in a hot-wall CVD set-up, as depicted in Fig. 1. To prevent potential clogging of exhaust pipelines due to precursor condensation, ammonia borane was placed inside the reactor tube. However, this arrangement was found to be challenging in controlling the thin film growth particularly due to limited control over the precursor flux during the deposition process.

A series of experimental investigations was conducted to examine the correlation between substrate positioning and *hBN* coverage on the Cu substrate, akin to previously reported studies.<sup>28</sup> A copper foil (length  $\times$  width = 15 cm  $\times$  1 cm) was placed in a vertical tube furnace at atmospheric pressure and a homogeneous deposition temperature of 1050 °C throughout the whole substrate length (Fig. 3a). The quality and composition of the CVD deposits were analysed using vibrational spectroscopy (FT-IR) and microscopy (SEM). In contrast to previously reported observations,<sup>28</sup> we

found that positioning the substrate closer to the precursor inlet yielded better quality films rather than placing it towards the reactor outlet, which resulted in minimal coverage and lower quality *hBN* deposits. These variations may be possibly attributed to subtle differences in experimental set-ups and process parameters, such as varying tube diameters or gas flow rates.

The typical *hBN* signals associated with B-N-B out-of-plane (805 cm<sup>-1</sup>) and B-N in-plane vibration (1350 cm<sup>-1</sup>)<sup>43</sup> were found to be the most intense peaks in the IR spectra (Fig. 3b and ESI-Fig. 2†). No vibration signals associated with B-O (770 cm<sup>-1</sup>, 500 cm<sup>-1</sup>)<sup>32,33</sup> or cBN (1068 cm<sup>-1</sup>) frequencies<sup>30,34</sup> were detected. However, IR spectra recorded for substrate regions closer to the reactor outlet (sample position: 8.5 cm) displayed an additional vibration signal at 1573 cm<sup>-1</sup>, typically attributed to  $-\text{NH}_2$  groups, which might be due to limited lateral mobility and diffusion of precursor fragments and surface-bound species.<sup>35-37</sup> This signal could possibly originate from undecomposed precursor groups or from N-terminated *hBN* flakes. Recent reports suggest the most stable forms of *hBN* involve N-termination, where open rings are terminated by NH<sub>2</sub>-groups.<sup>38</sup> Measurements close to the precursor outlet (sample position: 12.5 cm) showed only B-N-B out-of-plane stretching vibrations with low intensity, as well as a peak at 2983 cm<sup>-1</sup> (marked with \*) specific to the ATR-FT-IR measurement set-up. Corresponding SEM images (Fig. 3c) show a declining substrate coverage, where the front part facing the precursor inlet (sample positions: 0.5 cm and 4.5 cm) exhibits a high coating homogeneity, while at the back part (sample positions: 8.5 cm and 12.5 cm) a poor coverage and sample quality is observed. The results obtained with optimal substrate position were chosen for further analysis

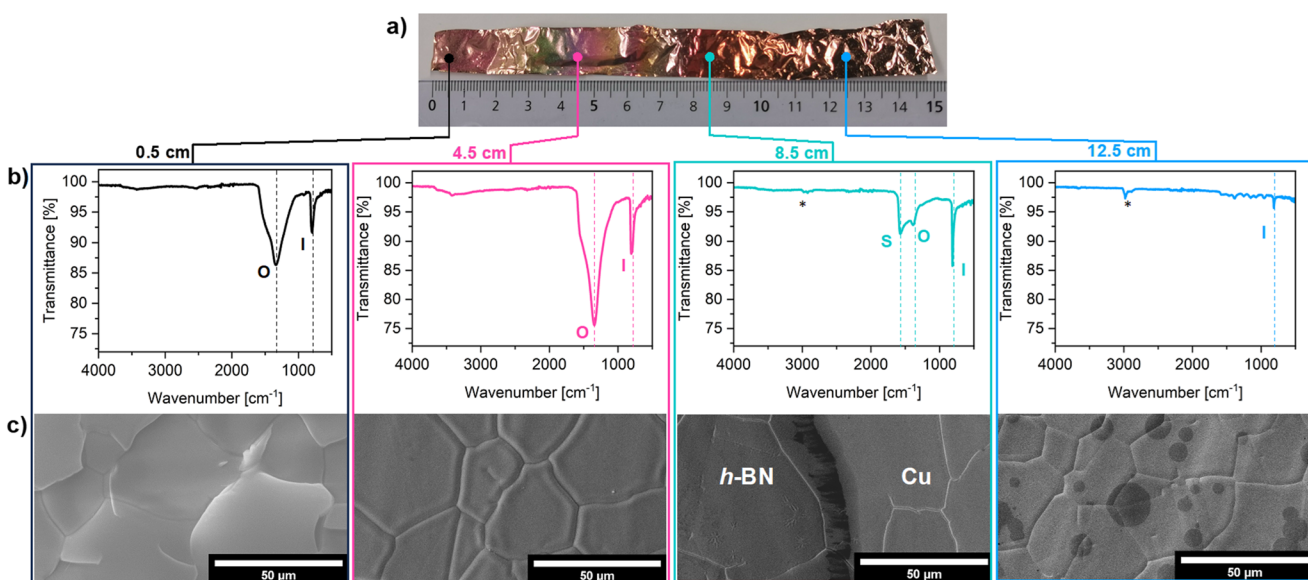


Fig. 3 (a) Photograph of a coated copper substrate (length  $\times$  width = 15 cm  $\times$  1 cm) with lateral changes of (b) spectroscopic (ATR-FT-IR) and (c) morphological (SEM) features at different sample positions (0.5 cm, 4.5 cm, 8.5 cm and 12.5 cm). The assignment of spectroscopic features follows the specific B-N-B out-of-plane (O; 792–811 cm<sup>-1</sup>), B-N in-plane (I; 1334–1382 cm<sup>-1</sup>) and N-H stretching (S; 1573 cm<sup>-1</sup>) modes. Signals inherent to the ATR-FT-IR measurement setup are marked with an asterisk (\*).



and for optimizing the deposition parameter to achieve homogeneous film growth and uniform coverage of the substrates. Studies for optimizing the deposition temperature were conducted in the temperature range 800–1050 °C with steps of 25–50 °C, as the melting point of copper (1085 °C) defines the upper temperature limit and temperatures below 800 °C were already investigated to yield only nanocrystalline BN.<sup>48</sup> Especially for ammonia borane, with its intrinsic disproportionation characteristics and decomposition reactions, which are affected by different solubilities of boron and nitrogen in copper, a precise control of the substrate temperature is essential for defined material deposition.<sup>40,49,50</sup> ATR-FT-IR analysis confirmed the optimal deposition temperature of 1050 °C with highest quality of *h*BN and absence of noticeable NH<sub>2</sub> vibrations (ESI-Fig. 1 and ESI-Table 1†).

Additional analysis confirmed the formation of *h*BN at 1050 °C from ammonia borane in a hot wall reactor: Similar to graphene, the D- and G-bands vibration in the Raman spectra with the wavenumber 1336 cm<sup>-1</sup> found to be overlapping with the prominent *h*BN signals at 1336 cm<sup>-1</sup> and 1592 cm<sup>-1</sup> (Fig. 4d and ESI-Fig. 5a†).<sup>51</sup> The XPS spectra confirmed the presence of boron nitride with a small amount of oxygen (B:N:O = 1.00 : 0.91 : 0.16) probably formed by the oxidation of unreacted precursor species (Fig. 4a–c). The morphology of the gained thin films (Fig. 4f) underly the further assumption of a heterogeneous CVD process and not dominated by reac-

tions in the reaction zone of the gas phase similar to observations reported by Ismach *et al.*<sup>48</sup> Only few particle-like structures were observed that could have formed in the gas phase due to predominant homogeneous nucleation instead of heterogeneous nucleation on the substrate surface.

For AFM and UV-vis analyses of the CVD deposits, the samples were transferred on a sapphire substrate that also enabled determination of film thickness and analyzing the bending and stretching vibrations in the material. The surface of the transferred film was found to be smooth with an average roughness of 9 nm measured over an area of 10 μm<sup>2</sup> whereas the overall thickness of the film was determined to be *ca.* 210 nm (ESI-Fig. 7a, c†). The high average roughness compared to RMS values of around 3–4 nm<sup>52,53</sup> is possibly due to the buckling of the film during the cool down phase. The UV-vis spectra of films transferred on sapphire showed characteristic UV absorption (maximum at 250 nm) corresponding to the large band gap of *h*BN (~6 eV) (Fig. 4e). Additional absorption peaks between 300–800 nm indicated non-uniform thickness and partial oxidation or contamination due to surface adsorbents, caused during the transfer process.

As-deposited thin-films were transferred onto TEM grids allowing both selected area electron diffraction (SAED) studies as well as high resolution imaging (Fig. 5). The (100) and (110) *h*BN lattice planes are clearly visible in the polycrystalline sample (Fig. 5a). Moreover, the bright spots indicate a pre-

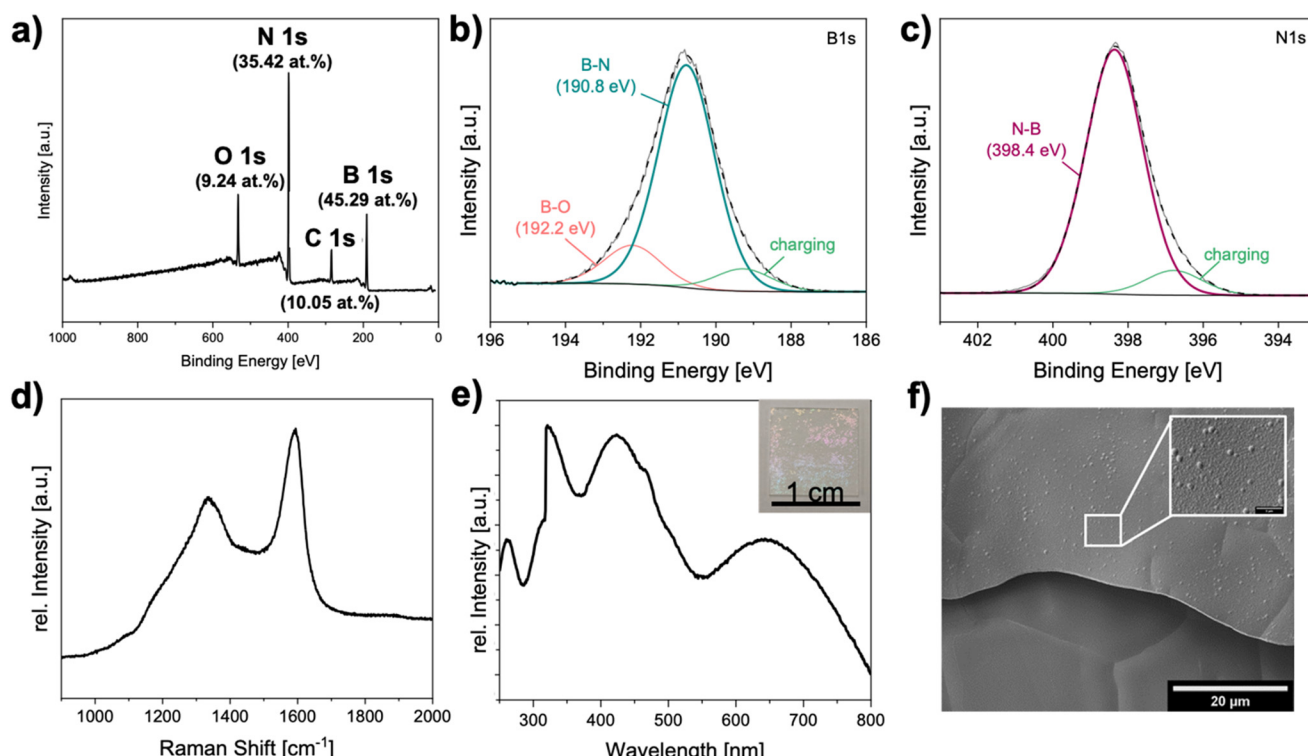
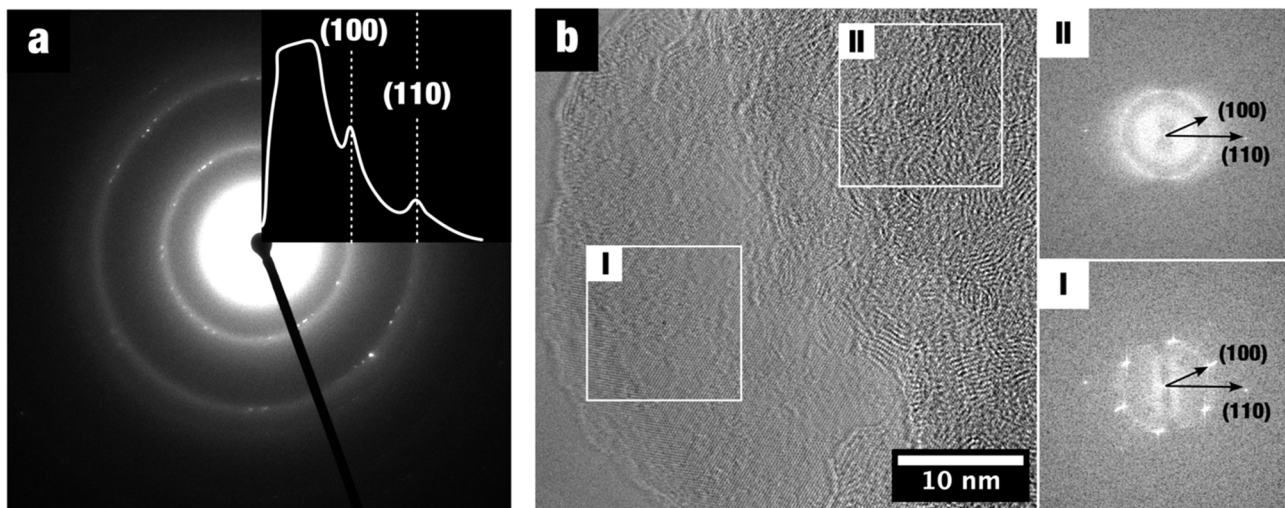


Fig. 4 *h*BN deposition on copper foil using ammonia borane in a hot-wall CVD reactor: (a) Survey XPS spectra indicating a B : N-ratio of 1 : 0.9. High resolution XPS spectra of B 1s (b) and N 1s (c) core levels support the formation of BN with some surface oxidation present. Characteristic D- and G-band vibration visible in the Raman spectrum of as deposited thin films (d) and UV-Vis absorption (e) of transferred thin films (inset) on sapphire substrate. Homogeneous morphology of as deposited thin films visible in SEM (f).



**Fig. 5** TEM analysis of hBN films transferred on TEM grids (Quantifoil): (a) SAED diffraction pattern of hBN films with a radial profile (inset) to highlight the (100) and (110) lattice planes in a polycrystalline sample region with preferred orientation (bright spots). (b) Crystalline few layer hBN supported by FFT of selected ROIs clearly show the (100) and (110) lattice planes.

ferred orientation and extended long range order, which is confirmed by high resolution image (Fig. 5b) also showing the same lattice planes in the FFT of two different regions of interest (ROI). The interplanar spacing was measured to be  $3.6\text{ \AA}$ , in accordance with literature values reported for hBN.<sup>31,40</sup> The differences in the structural features in two selected regions (inset I and inset II) are possibly caused by the transfer process of the hBN thin film in conjunction with different film thicknesses of the hBN flakes, that overlap in some parts.

In summary, layers of hBN with disordered topography and regional inhomogeneity in the film thickness were produced on a catalytic substrate (copper foil) by using ammonia borane in a hot-wall CVD reactor set-up.

In addition, hBN growth was evaluated using the same process conditions in a hot-wall CVD reactor enabling direct growth on non-catalytic substrates, with copper substrates acting as the reference. Single crystalline sapphire ( $\alpha\text{-Al}_2\text{O}_3$ ; *c*-plane (0001)) and silicon substrates with thermally grown  $\text{SiO}_2$  layer ( $\text{SiO}_2@\text{Si}(100)$ ) were chosen, due to their common usage in electronic device manufacturing. The IR-spectroscopic features of (I) N-H stretching ( $1575\text{ cm}^{-1}$ ), (II) B-N in-plane ( $1350\text{ cm}^{-1}$ ) and (III) B-N-B out-of-plane ( $805\text{ cm}^{-1}$ ), which are present on the Cu reference sample, are also visible – with significantly varying absolute intensities and intensity ratios – on the  $\alpha\text{-Al}_2\text{O}_3$  and  $\text{SiO}_2@\text{Si}(100)$  substrates, respectively (Fig. 6a). Especially on the sapphire substrate ( $\alpha\text{-Al}_2\text{O}_3$ ) pronounced vibrational bands of the desired hBN thin films are obtained.

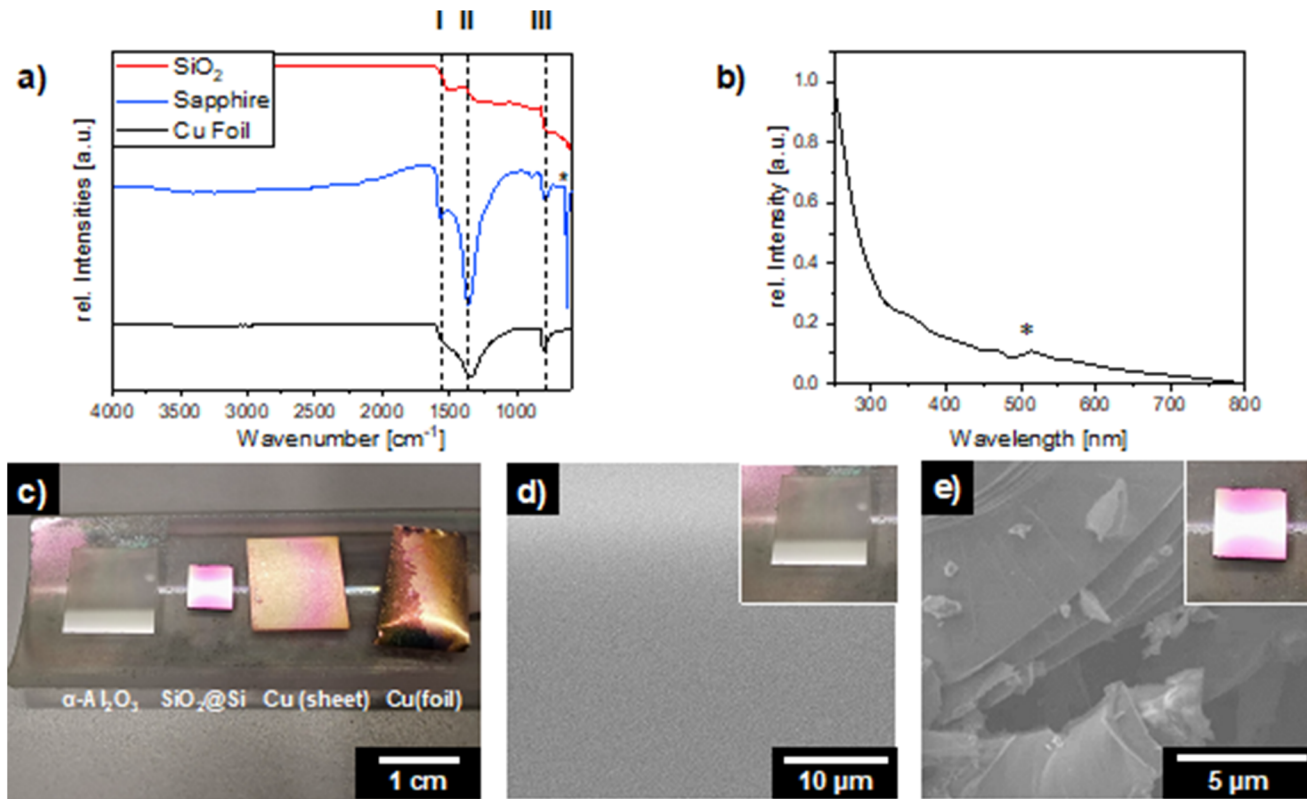
The UV-vis spectra recorded on sapphire (Fig. 6b) supports this assumption exhibited an absorption in the UV region with a maximum below  $250\text{ nm}$  which can be unambiguously attributed to the deposited material and corresponds to presumably hexagonal boron nitride. The accompanying SEM images of the films showed homogeneous morphology (Fig. 6d and e) with minimal roughness. Conversely, the  $\text{SiO}_2$  sample

revealed multi-layered structures. This difference in morphology, in contrast to the deposition on copper foil, is presumably due to epitaxial growth on sapphire,<sup>54,55</sup> whereas rougher surface in case of the silicon sample leads to non-uniform surface morphology.

In contrast to hBN growth on catalytic substrates (Cu), depositions on non-catalytic substrates such as sapphire highlight the need of higher deposition temperatures to facilitate sufficient mass transport and diffusion. So far, only few examples are known, where hBN films with sufficient film quality were obtained on non-catalytic substrates at the expense of significantly harsher process conditions (*e.g.*  $T > 1100\text{ }^\circ\text{C}$ , low pressure, pre-treatment).<sup>46,56–58</sup> The correlation of low-pressure process conditions with increased growth rates using ammonia borane on copper substrates was shown<sup>26</sup> and might be transferred to future thin film growth of hBN on non-catalytic substrates. Moreover, different growth mechanisms for hBN formation on non-catalytic sapphire substrates were proposed with an initial B-O bond formation, which can however significantly affect the films characteristics.<sup>59</sup>

In contrast to hot-wall reactors with a continuous temperature profile (Fig. 1c), cold wall reactors offer a segmented temperature profile, which allows to separate evaporation and decomposition processes (Fig. 1d) and therefore provide a more controlled environment for the precursor decomposition. Moreover, the deposition process is carried out at low pressure ( $p = 10^{-4}\text{ mbar}$ ), thus effectively excluding intermolecular collisions of the precursor molecules, by extending the mean free path length from few nanometers (*e.g.* atmospheric pressure conditions in hot wall reactor) to several meters (low pressure cold wall reactor) and avoiding the necessity of reactive carrier gases (*e.g.* 5%  $\text{H}_2$  in Ar). The cold-wall reactor in its current configuration used in this study is limited to a maximum substrate temperature of  $T_{\text{max}} = 950\text{ }^\circ\text{C}$ .

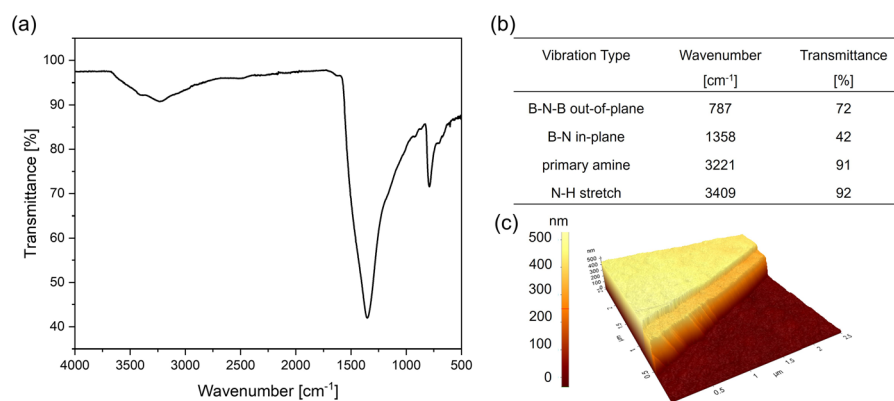




**Fig. 6** Hot-wall CVD of ammonia borane on sapphire ( $\alpha$ -Al<sub>2</sub>O<sub>3</sub>), SiO<sub>2</sub>@Si(100), and copper (sheet, foil): (a) ATR-FT-IR spectra of the samples on different substrates; the asterisk marks a spectroscopic feature of the underlying sapphire substrate (b) UV-vis spectrum of as-deposited hBN on sapphire; the asterisk highlights an inherent spectrometer artifact caused by an internal lamp change. (c) Photograph of the as deposited hBN thin films on different substrates. SEM image with inset picture of the samples of hBN thin films on (d)  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> and (e) SiO<sub>2</sub>@Si(100).

The thin films resulting from the decomposition of ammonia borane in a low-pressure cold wall reactor exhibited strong N-H vibration bands due to residual amine groups alongside the B-N vibrations (Fig. 7a). The broad signal with peaks at 3227 and 3413 cm<sup>-1</sup>, indicated that under chosen experimental conditions, complete decomposition of ammonia borane was not achieved. Transferred films on sapphire (Fig. 7c and ESI-Fig. 7b, d†) showed a film thickness of 458 nm using AFM measurements, however given the limitations of attainable temperatures further depositions using this set-up were discontinued.

In analogy to amino borane, borazine is also a single-source precursor with hexagonally pre-organized B-N units that structurally resemble with the 2D structure of the desired



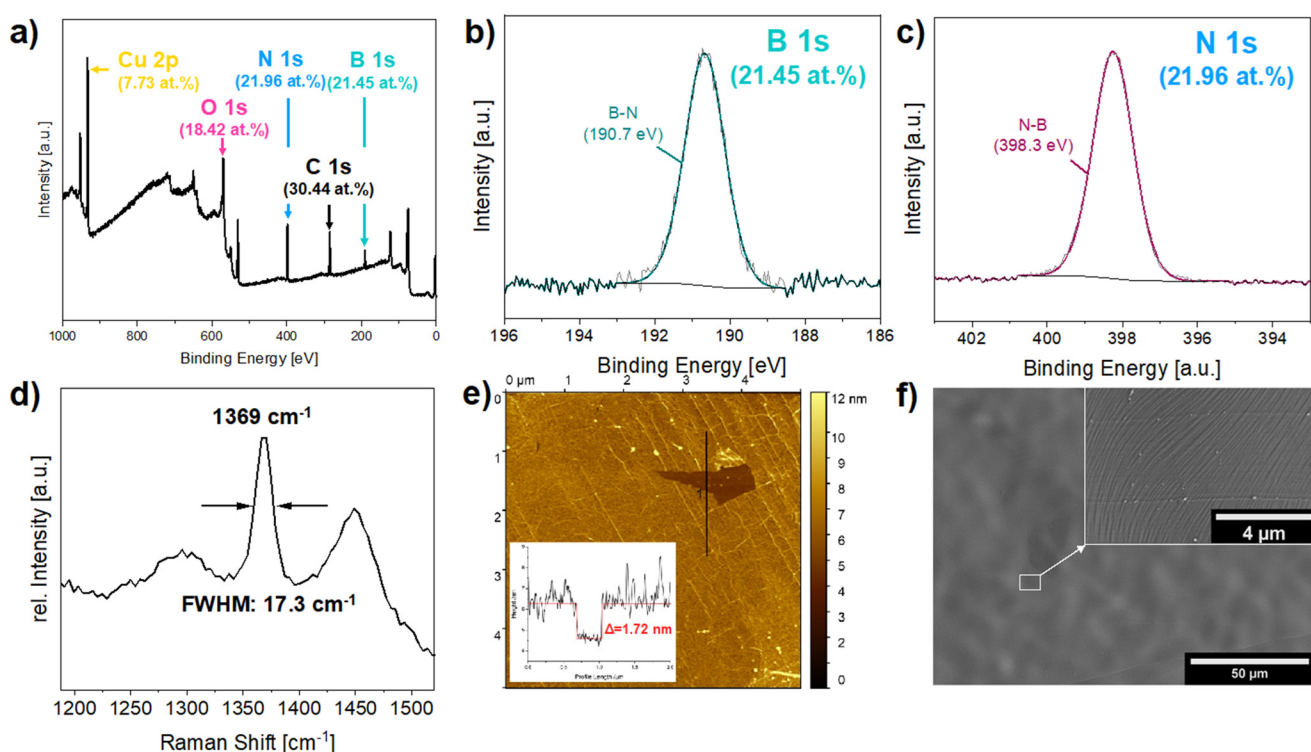
**Fig. 7** Decomposition of ammonia borane in a low-pressure cold-wall CVD reactor on copper foil. (a) ATR-FT-IR spectrum of the films on copper with (b) associated stretching frequencies and peak intensities and (c) an AFM image of the thin film transferred on a sapphire substrate.



*hBN* material. Interestingly, borazine is formed as an intermediate compound in the thermal decomposition of amino borane (Fig. 2) to *hBN*, which exhibits a significantly higher reactivity for forming *hBN* compared to amino borane and highlights the fact that initial precursor configuration is subject to bond-forming and bond-breaking phenomena that are intrinsic to the transformation of the molecular precursor into solid-state materials. Therefore, and due to the increased volatility of this compound, the precursor reservoir has to be maintained at  $-10\text{ }^{\circ}\text{C}$  to ensure a constant precursor flow during the decompositions in a hot wall reactor. As confirmed by XPS analysis, the films have a 1:1 B-N ratio (Fig. 8a) and no oxidation of the films is observed (missing of B-O contributions in the high-resolution B 1s XPS spectrum; Fig. 8b), thus confirming that borazine leads to a more defined chemical composition in the deposited *hBN* films without any side products or secondary phases. Due to the low of films thickness, signals from the underlying copper substrate, as well as a relatively high oxygen content are visible in the survey spectrum (Fig. 8a). The Raman signal detected at  $1369\text{ cm}^{-1}$  with a full width at half maximum (FWHM) of  $17.3\text{ cm}^{-1}$  (Fig. 8d) is in accordance with reported characteristics of high-quality *hBN* films.<sup>11,60,61</sup> The additional signals detected to the left and right of the *hBN* vibration are attributed to the underlying  $\text{SiO}_2$  substrate. AFM analysis (Fig. 8e and ESI-Fig. 8†) showed a step height of  $1.7\text{ nm}$  which is equivalent to four-layer structured *hBN* as measurements of 2D films on  $\text{SiO}_2$  are often

thicker than theoretical values of multiplying the interplanar distance under ambient conditions.<sup>11,62,63</sup> The SEM micrographs of the as-deposited films confirm the uniform coverage of the copper substrate (Fig. 8f). The textured surface is characteristic for *hBN* monolayers as a result of substrate shrinkage during the cooling phase.<sup>64</sup> This texturing effect is predominant in thicker films; however, it is no longer observable in the transferred films, apparently due to the relaxation of in-plane strain.

For application purposes, the wetting behaviour of 2D materials is an important criterion to fabricate electronic devices since the contact angle can be influenced by the morphology, thickness, and functional end groups of the thin film as well as defects and the underlying substrate.<sup>65</sup> Moreover, it provides information whether a high or low energy surface is obtained. For this purpose, the contact angles of three different solvents (water, diiodomethane and ethylene glycol (EG)) were measured on the *hBN* films produced by hot-wall CVD using AB and borazine as well as by cold-wall CVD using AB and bare copper as reference, respectively (ESI-Fig. 9†). In a first approach all surfaces can be classified as low energy surfaces (Fig. 9),<sup>66</sup> which show some deviation from the uncoated Cu reference substrate with a low surface energy of  $25.4 \pm 1.3\text{ mJ m}^{-2}$ , supporting the existence of an oxidized copper surface.<sup>67</sup> It is worth mentioning that, despite their different process conditions with ammonia borane deposited in a cold-wall reactor and borazine deposited in a hot-wall reactor, these



**Fig. 8** *hBN* films grown on copper substrate in hot-wall CVD reactor using borazine as precursor: (a) the XPS survey spectrum suggests an equimolar B : N ratio together with high resolution spectra of (b) B 1s and (c) N 1s region. (d) Raman spectrum and (e) AFM image of transferred *hBN* on a  $\text{SiO}_2$  substrate with inset of the height profile. (f) SEM image with magnification of the as-grown *hBN* thin film.



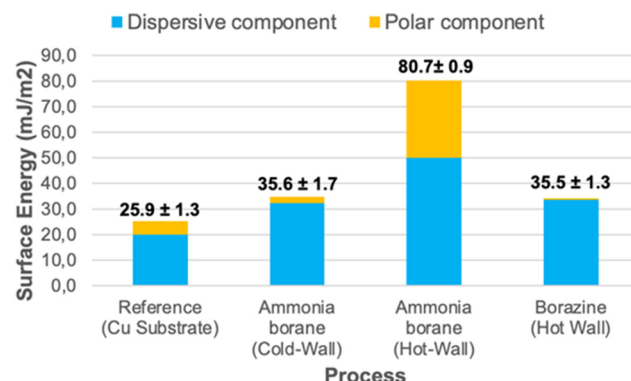


Fig. 9 Surface energies as calculated by the Fowkes relationship of disperse and polar components through contact angle measurements of  $\text{H}_2\text{O}$  and  $\text{CH}_2\text{I}_2$  (data available in ESI-Table 2†).

surfaces yielded the lowest surface energy of similar value ( $35.6 \pm 1.7 \text{ mJ m}^{-2}$  and  $35.5 \pm 1.3 \text{ mJ m}^{-2}$ , respectively), with minimal contributions from polar components. On the other hand, ammonia borane deposited in a hot-wall reactor yields the highest surface energy ( $80.7 \pm 0.9 \text{ mJ m}^{-2}$ ) with the largest absolute ( $30.3 \text{ mJ m}^{-2}$ ) and relative (37%) polar contributions of all *h*BN coatings, which is in line with the observed N–H moieties still present in the sample. Due to the fast spreading of the solvent on the *h*BN surface originating from the decomposition of ammonia borane in a hot-wall reactor low contact angle readings ( $\ll 10^\circ$ ) are afflicted with reasonable error. Nonetheless, the overall error of the surface energy can be estimated to  $80.7 \pm 0.9 \text{ mJ m}^{-2}$ .

## Conclusion

Gas phase depositions of hexagonal boron nitride were carried out using various chemical vapor deposition set-ups, including both hot and cold-wall configurations, and different precursor chemistries. The aim was to comparatively analyze the composition and structural quality of as-deposited boron nitride films. The collective analytical findings showed significant disparities in *h*BN quality depending contingent on the chosen CVD system and precursor composition. Although both ammonia borane and borazine proved to be suitable single-source precursors for producing phase-pure, high-quality *h*BN films, the comparison highlighted substantial differences depending on the chosen CVD configuration and parameters. Despite the advantages of ammonia borane in terms of ease of handling and low toxicity, the results obtained within the CVD

set-up used in this work did not yield few or monolayer *h*BN of high quality and size. Nevertheless, these conditions can reproducibly produce insulating layers in the range of a few hundred nanometers. The most promising findings were achieved from decomposing borazine on a copper substrate in a hot-wall CVD system using an Ar/H<sub>2</sub> mixture as the reactive carrier gas. In summary, this study presents the intricate inter-dependencies of precursor composition and CVD reactor configurations, as known in other coating systems<sup>68</sup>, in growing phase pure *h*BN thin films.

## Experimental section

### Cleaning of the samples

The copper foils were sonicated in iso-propanol, acetone, and water for 10 min each and dried in air. Afterwards, they were treated with an etchant solution (Nickel etchant from Alfa Aesar, 044586.LV) for 2 min and washed with ethanol.

### Deposition of hBN using ammonia borane in hot-wall CVD reactor

The cleaned samples were placed in the middle of the reactor on a quartz piece. The deposition tube was flushed with an Ar/H<sub>2</sub> mixture (95 : 5) for 45 min to remove the oxygen from the system. The reactor was then heated to the desired deposition temperature ( $T_d = 1025^\circ\text{C}$ ) with a heating rate of  $10^\circ\text{C min}^{-1}$  and holding time of 1 h. Subsequently, ammonia borane ( $m = 100 \text{ mg}$ , Sigma Aldrich, 95%) was gradually heated (70–140 °C) to transfer it in the gas phase (Table 1). After deposition, the substrate was annealed at the deposition temperature and with a continuous gas flow for 3 h.

### Deposition of hBN using ammonia borane in Cold-wall CVD reactor

The clean substrates were placed on the substrate holder. Within a quartz glass tube and the set-up was evacuated to  $10^{-5} \text{ mbar}$ . Ammonia borane (50 mg, Sigma Aldrich, 95%) heated to  $T_p = 85^\circ\text{C}$  was introduced in the chamber and decomposed on the substrates heated to  $T_d = 950^\circ\text{C}$ . The CVD process was run for a deposition time of 15 min (Table 1) and subsequently the substrates were cooled down.

### Deposition of hBN using borazine in Hot-wall CVD reactor

The cleaned copper foil was placed inside the reactor tube and the system was evacuated to  $10^{-2} \text{ mbar}$  for 20 min. Afterwards, the system was flushed with hydrogen (12 sccm) while maintaining the oven to  $1025^\circ\text{C}$  for 1 h. After 20 min of annealing

Table 1 Overview of the deposition parameters of the CVD processes

Process	Substrate	Precursor	$T_{\text{precursor}}$ [°C]	$T_{\text{substrate}}$ [°C]	$p_{\text{reactor}}$ [mbar]	Carrier gas
Hot-wall CVD	Cu, sapphire, $\text{SiO}_2@\text{Si}(100)$	AB	70–140	1025	1000	H <sub>2</sub> /Ar
Cold-wall CVD	Cu	AB	85	950	$10^{-5}$	—
Hot-wall CVD	Cu	Borazine	–10	1025	$10^{-2}$	H <sub>2</sub> /Ar

the substrate at the growth temperature, 2 min of film growth time was chosen using borazine (0.25 sccm, JSI Silicone) as precursor (Table 1).

The precursor was maintained at  $-10\text{ }^{\circ}\text{C}$  to prevent borazine polymerisation using a bath circulator from JioTech with a mixture of ethyl glycol and DI water (6 : 4) as cooling medium.

### Transfer of hBN thin films

For Raman analysis the as-deposited thin films were transferred to  $\text{SiO}_2@\text{Si}(100)$  substrates. Firstly, the samples were spin coated with PMMA (MW  $\sim 350\text{ g mol}^{-1}$ , 1 M in chlorobenzene, 4000 rpm, 60 s). The layer was detached from the copper *via* bubble transfer using a 1 M NaOH solution. Following the transfer step, the hBN/PMMA film were washed with water and transferred on  $\text{SiO}_2@\text{Si}(100)$  substrate. After drying the PMMA was removed by washing the samples several times with acetone.

### Characterization methods

The FT-IR spectra were recorded using a Spectrum 400 (PerkinElmer) in the range of  $500\text{--}4000\text{ cm}^{-1}$  with four scans per sample, using a data interval of 1 nm and a scan speed of  $266.75\text{ nm min}^{-1}$ . The SEM measurements were performed on a 300 VP RISE field emission scanning microscope (FESEM) from Zeiss Sigma. The Raman spectra were recorded using a Lab Ram HR 800 spectrometer (Horiba Ltd) using a green line  $\text{Ar}^+$  ion laser (514.5 nm) in the range of  $900\text{--}2000\text{ cm}^{-1}$ . The XPS spectra were measured with an ESCA M-Probe spectrometer (Surface Science Instruments). A monochromatic Al  $\text{K}_{\alpha}$  X-ray source (1486.6 eV,  $\lambda = 8.33\text{ \AA}$ ) was used and the spectra were recorded in the range of 0–1000 eV at ultra-high vacuum ( $10^{-9}\text{ mbar}$ ). The survey spectra were recorded using the following parameters: detector pass energy: 158.3 eV; step size: 0.5 eV; dwell time: 125 ms; data accumulation: seven scans per sample. High resolution spectra were recorded using the following parameters: detector pass energy: 55.2 eV; step size: 0.05 eV; dwell time: 175 ms; data accumulation: ten scans per sample (carbon) and 25 (other elements) were chosen, respectively. The data was evaluated using CASA XPS software suite (Casa Software Ltd). The UV-vis spectra were recorded using a LAMBDA 950 spectrometer (PerkinElmer). The samples were measured in the transmission mode compared to a reference consisting of the same substrate material. The AFM measurement of the sample thickness and surface structure of the BN layers were carried out on a XE-100 (Park System) atomic force microscope in a non-contact mode. High resolution transmission electron microscopy (HR-TEM) was carried out using a JEOL JEM 2200 FS equipped with UHR pole piece at 200 kV and an JEOL EDX detector. The thin films were transferred onto Quantifoil® Multi A10 copper grids (200 mesh). Data analysis was carried out using ImageJ software suite and dedicated macros. The contact angles were measured with the drop shape analysis system DSA 100 (KRÜSS GmbH). A defined amount (1  $\mu\text{L}$ ) of pure solvent (water, diiodomethane or ethylene glycol) is placed on the surface and the angle of the tangent between the baseline (substrate surface) and the

solvent droplet is measured; each measurement is repeated on 3 independent positions and the resulting mean value is reported.

## Data availability

The data supporting this article have been included as part of the ESI.<sup>†</sup>

## Conflicts of interest

The authors declare no competing conflict of interest.

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## References

- 1 K. S. Novoselov, A. K. Geim, S. V. Morozov, D.-e. Jiang, Y. Zhang, S. V. Dubonos, I. V. Grigorieva and A. A. Firsov, *Science*, 2004, **306**, 666–669.
- 2 A. K. Geim and K. S. Novoselov, *Nat. Mater.*, 2007, **6**, 183–191.
- 3 S. Moon, J. Kim, J. Park, S. Im, J. Kim, I. Hwang and J. K. Kim, *Adv. Mater.*, 2023, **35**, 2204161.
- 4 S. Vaidya, X. Gao, S. Dikshit, I. Aharonovich and T. Li, *Adv. Phys.: X*, 2023, **8**, 2206049.
- 5 V. Brune, N. Raydan, A. Sutorius, F. Hartl, B. Purohit, S. Gahlot, P. Bargiela, L. Burel, M. Wilhelm, C. Hegemann, U. Atamtürk, S. Mathur and S. Mishra, *Dalton Trans.*, 2021, **50**, 17346–17360.
- 6 V. Brune, C. Hegemann and S. Mathur, *Inorg. Chem.*, 2019, **58**, 9922–9934.
- 7 L. Wang, S. Yang, F. Zhou, Y. Gao, Y. Duo, R. Chen, J. Yang, J. Yan, J. Wang and J. Li, *Small*, 2023, 2306132.
- 8 S. Fukamachi, P. Solís-Fernández, K. Kawahara, D. Tanaka, T. Otake, Y.-C. Lin, K. Suenaga and H. Ago, *Nat. Electron.*, 2023, **6**, 126–136.



9 P. Ajayan, P. Kim and K. Banerjee, *Phys. Today*, 2016, **69**, 9–38.

10 R. Geick, C. Perry and G. Rupprecht, *Phys. Rev.*, 1966, **146**, 543.

11 T.-A. Chen, C.-P. Chuu, C.-C. Tseng, C.-K. Wen, H.-S. P. Wong, S. Pan, R. Li, T.-A. Chao, W.-C. Chueh and Y. Zhang, *Nature*, 2020, **579**, 219–223.

12 L. Wang, X. Xu, L. Zhang, R. Qiao, M. Wu, Z. Wang, S. Zhang, J. Liang, Z. Zhang and Z. Zhang, *Nature*, 2019, **570**, 91–95.

13 F. Zeng, R. Wang, W. Wei, Z. Feng, Q. Guo, Y. Ren, G. Cui, D. Zou, Z. Zhang and S. Liu, *Nat. Commun.*, 2023, **14**, 6421.

14 S. Roy, X. Zhang, A. B. Puthirath, A. Meiyazhagan, S. Bhattacharyya, M. M. Rahman, G. Babu, S. Susarla, S. K. Saju and M. K. Tran, *Adv. Mater.*, 2021, **33**, 2101589.

15 K. Kinoshita, R. Moriya, M. Onodera, Y. Wakafuji, S. Masubuchi, K. Watanabe, T. Taniguchi and T. Machida, *npj 2D Mater. Appl.*, 2019, **3**, 22.

16 F. Yuan, W. Jiao, F. Yang, W. Liu, J. Liu, Z. Xu and R. Wang, *J. Mater. Chem. C*, 2017, **5**, 6359–6368.

17 D. Gonzalez Ortiz, C. Pochat-Bohatier, J. Cambedouzou, M. Bechelany and P. Miele, *Nanomaterials*, 2018, **8**, 716.

18 L. Pauling, *J. Am. Chem. Soc.*, 1931, **53**, 3225–3237.

19 A. J. L. Catão and A. López-Castillo, *J. Mol. Model.*, 2017, **23**, 1–12.

20 C.-L. Lo, M. Catalano, K. K. Smithe, L. Wang, S. Zhang, E. Pop, M. J. Kim and Z. Chen, *npj 2D Mater. Appl.*, 2017, **1**, 42.

21 J. Wang, F. Ma, W. Liang and M. Sun, *Mater. Today Phys.*, 2017, **2**, 6–34.

22 J. Eichler and C. Lesniak, *J. Eur. Ceram. Soc.*, 2008, **28**, 1105–1109.

23 J. Lee, A. V. Ravichandran, J. Mohan, L. Cheng, A. T. Lucero, H. Zhu, Z. Che, M. Catalano, M. J. Kim and R. M. Wallace, *ACS Appl. Mater. Interfaces*, 2020, **12**, 36688–36694.

24 S. Le Gallet, G. Chollon, F. Rebillat, A. Guette, X. Bourrat, R. Naslain, M. Couzi and J.-L. Bruneel, *J. Eur. Ceram. Soc.*, 2004, **24**, 33–44.

25 X. Chen, C. Tan, X. Liu, K. Luan, Y. Guan, X. Liu, J. Zhao, L. Hou, Y. Gao and Z. Chen, *J. Mater. Sci.: Mater. Electron.*, 2021, **32**, 3713–3719.

26 W.-H. Lin, V. W. Brar, D. Jariwala, M. C. Sherrott, W.-S. Tseng, C.-I. Wu, N.-C. Yeh and H. A. Atwater, *Chem. Mater.*, 2017, **29**, 4700–4707.

27 Y. Uchida, T. Iwaizako, S. Mizuno, M. Tsuji and H. Ago, *Phys. Chem. Chem. Phys.*, 2017, **19**, 8230–8235.

28 R. Y. Tay, X. Wang, S. H. Tsang, G. C. Loh, R. S. Singh, H. Li, G. Mallick and E. H. T. Teo, *J. Mater. Chem. C*, 2014, **2**, 1650–1657.

29 Y. Shi, C. Hamsen, X. Jia, K. K. Kim, A. Reina, M. Hofmann, A. L. Hsu, K. Zhang, H. Li, Z.-Y. Juang, M. S. Dresselhaus, L.-J. Li and J. Kong, *Nano Lett.*, 2010, **10**, 4143.

30 J. S. Lee, S. H. Choi, S. J. Yun, Y. I. Kim, S. Boando, J.-H. Park, B. G. Shin, H. Ko, S. H. Lee and Y.-M. Kim, *Science*, 2018, **362**, 817–821.

31 D. Graf, J. Schläfer, S. Garbe, A. Klein and S. Mathur, *Chem. Mater.*, 2017, **29**, 5877–5885.

32 L. Jürgensen, D. Höll, M. Frank, T. Ludwig, D. Graf, A. K. Schmidt-Verma, A. Raauf, I. Gessner and S. Mathur, *Dalton Trans.*, 2020, **49**, 13317–13325.

33 P. Tutacz, D. Stadler, T. Karimpour, T. s. Duchoň, S. Cramm, C. M. Schneider, T. Fischer, D. N. Mueller and S. Mathur, *Chem. Mater.*, 2023, **35**, 8050–8056.

34 U. Atamturk, E. Jung, T. Fischer and S. Mathur, *Chem. Mater.*, 2022, **34**, 7344–7356.

35 S. Mathur, M. Veith, T. Ruegamer, E. Hemmer and H. Shen, *Chem. Mater.*, 2004, **16**, 1304–1312.

36 S. Frueh, R. Kellett, C. Mallery, T. Molter, W. S. Willis, C. King'ondu and S. L. Suib, *Inorg. Chem.*, 2011, **50**, 783–792.

37 P. Wang and X.-d. Kang, *Dalton Trans.*, 2008, 5400–5413.

38 E. Mayer, *Inorg. Chem.*, 1972, **11**, 866–869.

39 E. Mayer, *Inorg. Chem.*, 1973, **12**, 1954–1955.

40 C. R. Miranda and G. Ceder, *J. Chem. Phys.*, 2007, **126**, 184703-1–184703-11.

41 M. Schwarz, A. Riss, M. Garnica, J. Ducke, P. S. Deimel, D. A. Duncan, P. K. Thakur, T.-L. Lee, A. P. Seitsonen and J. V. Barth, *ACS nano*, 2017, **11**, 9151–9161.

42 H. Jeong, D. Y. Kim, J. Kim, S. Moon, N. Han, S. H. Lee, O. F. N. Okello, K. Song, S.-Y. Choi and J. K. Kim, *Sci. Rep.*, 2019, **9**, 1–8.

43 Z. Zhang, X. Ji, J. Shi, X. Zhou, S. Zhang, Y. Hou, Y. Qi, Q. Fang, Q. Ji and Y. Zhang, *ACS Nano*, 2017, **11**, 4328–4336.

44 T. H. Lee, K. Kim, G. Kim, H. J. Park, D. Scullion, L. Shaw, M.-G. Kim, X. Gu, W.-G. Bae and E. J. Santos, *Chem. Mater.*, 2017, **29**, 2341–2347.

45 A. Bansal, M. Hilse, B. Huet, K. Wang, A. Kozhakhmetov, J. H. Kim, S. Bachu, N. Alem, R. Collazo and J. A. Robinson, *ACS Appl. Mater. Interfaces*, 2021, **13**, 54516–54526.

46 R. Singhal, E. Echeverria, D. N. McIlroy and R. N. Singh, *Thin Solid Films*, 2021, **733**, 138812.

47 T. Singh, T. Lehnert, T. Leuning, D. Sahu and S. Mathur, *Appl. Surf. Sci.*, 2014, **289**, 27–32.

48 A. Ismach, H. Chou, D. A. Ferrer, Y. Wu, S. McDonnell, H. C. Floresca, A. Covacevich, C. Pope, R. Piner and M. J. Kim, *ACS Nano*, 2012, **6**, 6378–6385.

49 F. Baitalow, J. Baumann, G. Wolf, K. Jaenicke-Rößler and G. Leitner, *Thermochim. Acta*, 2002, **391**, 159–168.

50 A. Lozovoi and A. Paxton, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2008, **77**, 165413.

51 L. M. Malard, M. A. Pimenta, G. Dresselhaus and M. S. Dresselhaus, *Phys. Rep.*, 2009, **473**, 51–87.

52 J. C. Koepke, J. D. Wood, Y. Chen, S. W. Schmucker, X. Liu, N. N. Chang, L. Nienhaus, J. W. Do, E. A. Carrion and J. Hewaparakrama, *Chem. Mater.*, 2016, **28**, 4169–4179.

53 J. Fernandes, T. Queirós, J. Rodrigues, S. S. Nemala, A. P. LaGrow, E. Placidi, P. Alpuim, J. B. Nieder and A. Capasso, *FlatChem*, 2022, **33**, 100366.



54 Q. Li, Q. Wu, J. Gao, T. Wei, J. Sun, H. Hong, Z. Dou, Z. Zhang, M. H. Rümmeli and P. Gao, *Adv. Mater. Interfaces*, 2018, **5**, 1800662.

55 A.-R. Jang, S. Hong, C. Hyun, S. I. Yoon, G. Kim, H. Y. Jeong, T. J. Shin, S. O. Park, K. Wong and S. K. Kwak, *Nano Lett.*, 2016, **16**, 3360–3366.

56 R. Page, J. Casamento, Y. Cho, S. Rouvimov, H. G. Xing and D. Jena, *Phys. Rev. Mater.*, 2019, **3**, 064001.

57 G. Wang, J. Meng, J. Chen, Y. Cheng, J. Huang, S. Zhang, Z. Yin, J. Jiang, J. Wu and X. Zhang, *Cryst. Growth Des.*, 2022, **22**, 7207–7214.

58 M. Tokarczyk, A. K. Dąbrowska, G. Kowalski, R. Bożek, J. Iwański, J. Binder, R. Stępniewski and A. Wysmołek, *2D Mater.*, 2023, **10**, 025010.

59 R. Chen, Q. Li, Q. Zhang, J. Li, Z. Zhang, W. Fang, L. Wang, F. Yun, T. Wang and Y. Hao, *Cryst. Growth Des.*, 2023, **23**, 8783–8792.

60 G. Kim, A.-R. Jang, H. Y. Jeong, Z. Lee, D. J. Kang and H. S. Shin, *Nano Lett.*, 2013, **13**, 1834–1839.

61 R. V. Gorbachev, I. Riaz, R. R. Nair, R. Jalil, L. Britnell, B. D. Belle, E. W. Hill, K. S. Novoselov, K. Watanabe and T. Taniguchi, *Small*, 2011, **7**, 465–468.

62 D. Wickramaratne, L. Weston and C. G. Van de Walle, *J. Phys. Chem. C*, 2018, **122**, 25524–25529.

63 M. Ishigami, J.-H. Chen, W. G. Cullen, M. S. Fuhrer and E. D. Williams, *Nano Lett.*, 2007, **7**, 1643–1648.

64 J. Lin, R. Y. Tay, H. Li, L. Jing, S. H. Tsang, H. Wang, M. Zhu, D. G. McCulloch and E. H. T. Teo, *Nanoscale*, 2018, **10**, 16243–16251.

65 D. Y. Kwok and A. W. Neumann, *Adv. Colloid Interface Sci.*, 1999, **81**, 167–249.

66 M. Annamalai, K. Gopinadhan, S. A. Han, S. Saha, H. J. Park, E. B. Cho, B. Kumar, A. Patra, S.-W. Kim and T. Venkatesan, *Nanoscale*, 2016, **8**, 5764–5770.

67 A. Ogwu and T. Darma, *Sci. Rep.*, 2021, **11**, 4276.

68 A. Mettenbörger, T. Singh, A. P. Singh, T. T. Järvi, M. Moseler, M. Valldor and S. Mathur, *Int. J. Hydrogen Energy*, 2014, **39**(10), 4828–4835.

