Hydroxylated boron nitride materials: from structures to functional applications

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ABSTRACT

Functionalization of boron nitride (BN) materials with hydroxyls has attracted great attention to accomplish better performances at micro- and nanoscale. BN surface hydroxylation, in fact, induces a change in properties and allows expanding the fields of application. In this review, we have summarized the state-of-the-art in developing hydroxylated bulk and nanoscale BN materials. The different synthesis routes to develop hydroxyl BN have been critically discussed. What emerges is the great variety of possible strategies to achieve BN hydroxylation, which, in turn, represents one of the most suitable methods to improve the solubility of BN nanomaterials. The improved stability of BN solutions creates conditions for producing high-quality nanocomposites. Furthermore, new interesting optical and electronic properties may arise from the functionalization by OH groups as displayed by a wide range of both theoretical and experimental studies. After the presentation of the most significant systems and methodologies, we question of future perspective and important trends of the next generation BN materials as well as the possible areas of advanced research.
Hydroxyl functionalization of boron nitride materials is a key method to control and enhance the properties and design new functional applications.

Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| BN           | Boron nitride |
| h-, c-, r-, w- | Hexagonal, cubic, rhombohedral, wurtzite |
| NS           | Nanosheet   |
| NR           | Nanoribbon  |
| NT           | Nanotube    |
| ICTES        | (3-Isocyanatopropyl) triethoxysilane |
| APTES        | 3-Aminopropyltriethoxysilane |
| PS           | Polystyrene |
| PP           | Polypropylene |
| P2VP         | Poly(2-vinylpyridine) |
| PNBMT        | Poly(5-norbornene-2-methanol) |
| MPTMS        | (3-Mercaptopropyl) trimethoxysilane |
| SPEEK        | Sulfonated poly (ether ether ketone) |
| PFOS         | Perfluoroctanesulfonate |
| PFDA         | Perfluorodecanoate |
| DMSO         | Dimethyl sulfoxide |
| NMP          | N-methyl-2-pyrrolidone |
| PAI          | Polyamide-imide |
| PVA          | Polyvinyl alcohol |
| NLO          | Nonlinear optics |
| OL           | Optical limiting |
| EELS         | Electron energy loss spectroscopy |
| PNIPAM       | Poly(N-isopropylacrylamide) |
| SDS          | Sodium dodecyl sulfate |
| TBP          | Di-tert-butylperoxide |
| PVDF         | Poly(vinylidene fluoride) |
| CVD          | Chemical vapor deposition |
| PC           | Polycarbonate |
| DFT          | Density functional theory |
| MD           | Molecular dynamics |
| GGA          | Generalized gradient approximation |
| DNP          | Double numerical basis sets plus polarization function |
| CASTEP       | Cambridge Serial Total Energy Package |
| DOX          | Doxorubicin hydrochloride |
Introduction

The remarkable properties exhibited by boron nitride-based (BN) nanomaterials have attracted much attention and have risen the expectations of new breakthrough applications. Structurally similar to carbon system, BNs exist in four different polyorphs: graphite-like hexagonal BN (h-BN) and rhombohedral (r-BN), diamond-like cubic BN (c-BN) and wurtzite BN (w-BN) \[ 1, 2\]. Among them, h-BN has shown excellent properties, such as outstanding mechanical strength, high thermal conductivity, and excellent lubrication. 2D h-BN has a layered structure similar to graphene (see Fig. 1a). Graphene is a kind of honeycomb crystals with sp\(^2\) hybridized carbon atoms, and the hexagonal geometry endows it with a very stable structure. An equal number of B and N atoms forms the h-BN layer, where the sp\(^2\) covalent bond links alternating B and N atoms. Weak Van der Waals bonds between different planes keep together the h-BN 2D layers, similarly to graphite; however, the B–N bond has a slight ionic character in comparison with the covalent C–C bond of graphite.

The h-BNs unique structural characteristics are also reflected in the electrical and optical properties \[ 3–6\]. For instance, h-BN presents a wide bandgap (\( E_g \)) value, around 6 eV, confirmed by both theoretical and experimental works \[ 7, 8\]. As in the case of carbon nanomaterials, which includes carbon nanotubes

![Figure 1](image-url) Structural models of h-BN crystal and four typical BN nanomaterials: a h-BN structure and some basic parameters, b BNNSs, c BNNRs, d BNNTs, and e BN fullerenes. The relevant TEM images are shown on the bottom, reproduced from Refs. [11–14] with permissions of Copyright (2010) American Chemical Society, Copyright (2014) American Chemical Society, Copyright (2009) Institute of Physics, and Copyright (2015) Springer.
and graphene quantum dots, the layered structure of h-BN is also flexible enough to form many nanostructures [9, 10], as shown in Fig. 1b–e. In fact, there are BN nanosheets (BNNSs, also indicated as white graphene) [11], BN nanoribbons (BNNRs) [12], BN nanotubes (BNNTs) [13], BN fullerenes and cages [14], etc.

The number of publications on BN-materials has dramatically increased in the past ten years (see Fig. 2a). Figure 2b shows the publishing situation of BN materials in 2019: about 50% regards nanomaterials, with research on both BNNTs and BNNSs accounting for nearly 20%. According to these reports, BN systems are promising candidates as optoelectronic devices [15, 16], quantum emitters [17, 18], semiconductors [19], energy storage [20], thermal conductors [21], cosmetic products [22], water cleaning [23, 24], photocatalytic [25], as well as biomedical science [26], etc.

BN systems for functional applications require advanced synthesis and processing routes [27]. Nanomaterials are strongly dependent on their surface, and the functionalization methods are the key to design the desired properties. This becomes particularly important in 2D nanomaterials whose integration into devices and tailoring of properties are controlled via surface modification. In the case of BNs, some functional strategies have been realized to modify their surface, including hydroxyl (–OH), alkoxy (–OR), amino (–NH₂), and alkyl (–R) groups [1, 28, 29]. Compared with other surface functionalization, hydroxylation is the most important functionalization route for modifications of BN materials. Firstly, –OH groups are usually covalently attached to B sites in BN net-structure, whose introduction is easy and efficient to carry out, such as, by direct thermal oxidation of h-BN in the air [30], and sonication in H₂O [31]. Importantly, hydroxyl groups have been applied as an intermediary bridge to realize further functionalization. For example, –OH on BN surface can react with organically modified alkoxides to obtain organosilane-functionalized BNs [32]. On the other hand, –OH groups in BN materials can react with –COOH of perfluoro butyric acid (CF₃CF₂CF₂COOH) or thioglycolic acid (HSCH₂-COOH) to prepared esterified (–OCOR) BNs [33].

Currently, surface hydroxylation has shown to be a flexible technique to change the chemical characteristics of BNs, which will further affect their physical features and mechanical properties. Hydroxylated BNs exhibit enhanced and innovative properties, either physical or chemical, in comparison with pristine systems. In particular, the introduction of hydroxyl groups allows homogeneous dispersion of BN materials in several solvents, facilitating the incorporation into matrixes, such as polymers and sol–gel systems. This is a key point for integrating BN nanomaterials into a solid-state device.

We have focused this review on the hydroxylated process of BN systems, from structure to applications. The surface hydroxylation has been analyzed both theoretically and experimentally. The current experimental strategies to achieve OH groups on bulk and nanoscale BNs surfaces have been critically revised, focusing on the change of properties caused by the functionalization.

**Hydroxylated BN micron-scale materials**

**Direct hydroxyl-modification of BNs**

The hydroxylation of h-BNs on micron scale is an important process for the development of commercial materials, and several strategies have been developed so far. A direct route to form hydroxyl species on the...
**h-BN surface is the treatment of powders in air at high temperature or in solution with the presence of strong acids, alkali, and oxidants agents (Fig. 3).**

The simplest and most direct way to obtain h-BNs functionalized with –OH is heating the raw material in air, e.g., 1000 °C for 2 h [30, 34, 35]. The hydroxyl groups allow a further BNs functionalization. For instance, BN/epoxy resin nanocomposites can be prepared via a sol–gel method using (3-isocyanato-propyl)triethoxysilane (ICTES) as a cross-linker, which enhances the thermal-oxidative stability [30]. Herein, the reaction between organosilane and –OH groups on h-BN surface permits a uniform dispersion of the particles into an epoxy resin. Another example is the modification of hydroxyl-functional BNs by 3-aminopropyltriethoxysilane (APTES) [35]. A BN-polystyrene (PS) composite has been obtained by in situ polymerization of styrene with the presence of APTES-BNs. By mixing BN-PS, PS and polypropylene (PP), ternary composites have been fabricated as BN-PS/(PP/PS). The thermal conductivity of the ternary composites is twice higher than naked PP and 30% higher than PP/BN composites.

For increasing the hydroxylation, pre-treatment of BN in a hot acidic solution, such as concentrated HNO3 [32], H2SO4 [36], and mixed acids, is a good choice [37, 38]. HO-BNs can be prepared by sonicating BNs in HNO3 and H2SO4 (volume ratio = 3:1) mixed solutions (Fig. 4) [37]. The HO-BNs could form covalent composites with poly(2-vinylpyridine) (P2VP) and poly(5-norbornene-2-methanol) (PNBMT) via hydrogen bonding interactions and etherification reaction, respectively, with –OH groups. Ternary BN/P2VP/PNBMT composites have been obtained by self-assembly of BN/P2VP and BN/PNBMT; they exhibit dense packing, lower surface area, and higher density. These properties are attributed to the participation in cross-linking and the curing reaction by OH on BNs. Additionally, NH2 groups can be introduced through the reaction between HO-BNs and APTES. The product has also been self-assembled with graphene oxide under electrostatic interaction [32].

Another route to hydroxylate h-BN materials is pre-treatment of BN in a basic solution by a combination of stirring and heating [39–43]. In detail, hydroxyl-functional BNs are obtained by grafting organosilane coupling agents, which is a facile way to decrease the interface thermal resistance [39–41]. The organosilane-BNs have been embedded into some polymeric matrixes, such as epoxy [39, 40] and polyurethane acrylate [41], resulting in an improved thermal conductivity and long-term stability. The hydroxyl groups on BNs can further form hydrogen bonding with PVA chains, and, in this way, HO-BN/PVA composite films can be prepared [42]. The composite has shown enhanced thermal conductivity, flame-retardancy capability, and mechanical flexibility.

Oxidants have also been employed to oxidize and functionalize h-BN with OH groups, including H2O2 [44, 45] and KMnO4 [45]. A hydrothermal method with 30% H2O2 was used to modify the surface of BNs, which not only increases hydroxyl percentage on BN but also introduces some boron and oxygen bondings [44]. To better hydroxylate BN, [45] combination of oxidants and acids, including H2SO4, H3PO4, KMnO4, and H2O2, has been employed. After modification by (3-mercaptopropyl)trimethoxysilane (MPTMS), the prepared BN uniformly dispersed in sulfonated poly (ether ether ketone) (SPEEK) to form composite membranes. The membranes show high proton conductivity, low methanol crossover, and high selectivity. These proton exchange membranes can be used to develop direct methanol fuel cells.

Some other approaches have also been used for h-BNs hydroxylation. For example, hydroxyl-functionalized BNs have been obtained by ball milling of h-BN and sucrose [46]. The resulted carbonized BN/lignosulfonate/rubber composites show excellent electrical and mechanical performances, which can be used for producing thermal interfacial management devices.

**Bottom-up ways for hydroxylated BNs**

Bottom-up synthesis routes are also popular for the preparation of hydroxylated BNs. As summarized in Table 1, the resulted products can be divided into two categories: (i) HO-BNs directly obtained via heating temperature in the range of 850–1100 °C [47–51]; (ii)
$h$-BNs hydroxylated via post-processing at the temperature of 1500 °C [52, 53].

In situ hydroxylation has been achieved using H$_3$BO$_3$ and urea as raw materials and applying a thermal treatment at 900 °C in N$_2$ atmosphere [47]. The obtained HO-BNs have been tested to remove perfluorooctanesulfonate (PFOS) and perfluorodecanoate (PFDA). Addition of Ca$^{2+}$ causes a significant increase in the removal effect because of enhanced electrostatic interactions.

H$_3$BO$_3$ has been also combined with melamine to prepare $h$-BN fibers by heating at 1560 °C under NH$_3$ flow, following hydroxy-functionalization by acidic treatment [52]. The resulting HO-BNs exhibit an enhanced removal performance compared with pristine $h$-BNs, that has been tested in methylene blue degradation.
Hydroxylated BNNSs

Top-down routes for producing hydroxylated BNNSs

2D h-BN sheets represent a very important family of BN materials. Top-down routes can be divided into three main types, as shown in Fig. 5: (i) HO-BNNSs are obtained by simultaneous exfoliation and functionalization of bulk h-BN; (ii) h-BNs are firstly hydroxylated and then HO-BNs are exfoliated into nanosheets; (iii) h-BNNSSs are prepared by the exfoliation of BNs, and then functionalized with hydroxyl groups.

One-pot sonication-assisted exfoliation

h-BNs, hydroxylated nanosheets can be directly prepared by thermal oxidation of BNs in air, typically at 1000 °C, followed by treatment in water [54]. Another simple way to obtain HO-BNNSs is to exfoliate h-BNs using water as dispersant solvent by sonication without any additional use of surfactants or organic functionalization [31, 55–59]. Sonication process can provide an intensive longitudinal shear force leading to complete exfoliation of the bulk into h-BNNSs with few layers and reduced lateral sizes, with attached OH groups (Fig. 6) [31]. HO-BNNSs are partially oxidized in the hydrolysis process [57] and show strong affinity toward proteins by Lewis acid–base interactions, confirming that surface-organic-free BNNSs have a high potential in biological applications.

Very recently, our research group has found that HO-BNNSs possess a weak photoluminescent emission in a visible range, centered at ~ 400 nm [58]. Tailored defects can be further introduced through thermal treatment in air, and a maximum emissive intensity is achieved at 300 °C due to the role of hydroxyl groups created by the oxidation process (see Fig. 7). Subsequently, a higher temperature treatment (500 or 700 °C) causes a continuous fluorescent quenching. Notably, only bulk h-BNs exfoliated in water allow introducing –OH groups with the consequent emission. For instance, defect-free BNNSs are obtained when sonication is carried out in dimethyl sulfoxide (DMSO) or N-methyl-2-pyrrolidone (NMP), but they do not exhibit any fluorescence.

Various additives have already been investigated to enhance the hydrolyzation via sonication-assisted exfoliation. Addition of HNO₃ [60], NaOH [61], sodium cholate [62], and plant extracts [63] is one of the most efficient strategies. As-prepared HO-BNNSs obtained by a tip-type sonication process under the assistance of sodium cholate can be easily incorporated into thermoplastic polyamide-imide (PAI) in the presence of polyvinyl alcohol (PVA) [62]. The hydrogen-bonded cross-linking networks between BNNSs and the polymer lead to 3D BN-composites, showing a thermal conductivity enhancement of 409%. More recently, it has been demonstrated the efficacy of extracts of natural plants which work as ecofriendly surfactant agents and improve the exfoliation of h-BN into few-layer nanosheets under sonication. The obtained BNNSs result covered by abundant phytochemicals with OH groups [63].

One-pot ball-milling-assisted exfoliation

Ball-milling method realizes the exfoliation by the mechanical interaction between balls and h-BNs. The collision of the ball with the flank face of BN can make it to wrinkle and exfoliate (Fig. 8). Besides, as seen in the previous section, the introduction of some chemical agents, such as NaOH and KOH [64–67], NaClO [68], urea [69, 70], H₃BO₃ [71], and β-cyclodextrin [72], can lead to a synergetic effect of mechanical process and chemical peeling.

NaOH-assisted ball milling process has been used to prepare HO-BNNSs with a yields of ~ 18% through a simultaneously exfoliation and functionalization process [64]. The NaOH solution is not only used to promote the shear-force exfoliation, but also to functionalize the sheets via reaction of h-BN and OH radicals, leading to a high oxygen content of around 6.4 at%. Liu et al. [67] reported a kind of nanohybrids composed of BNNSs and gold

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Figure 5 Top-down routes to hydroxyl-functionalize BNNSs.
nanoclusters, where the ball-milling technique has been used to prepare BNNSs in the presence of mixed NaOH and KOH. The nanohybrids can prevent aggregation of nanoclusters and exhibit a better catalytic activity for reducing 4-nitrophenol.

$h$-BNs can be functionalized by both –OH and –NH$_2$ by adding urea [69] or H$_3$BO$_3$ into the ball-milling system [71]. A nanocomposite membrane can be fabricated by embedding the obtained $h$-BNNSs into thermally rearranged polyimide for H$_2$ separation applications.

**Other one-pot routes**

As for graphene oxide (GO), Hummer’s-like routes have been specifically designed to synthesize oxidized BNNSs [73–76] by using, for example, a mixture of H$_3$PO$_4$ and H$_2$SO$_4$ acids [73]. The solution has been later treated at 70 °C for 12 h, before adding H$_2$O$_2$ to stop the oxidation process. HO-BNNSs can be also obtained in a solution of H$_2$SO$_4$ and KMnO$_4$ at the mild temperature of 40 °C for 6 h followed by a treatment with H$_2$O$_2$ at 110 °C for 48 h [75].

HO-BNNS with large lateral size, around 2 μm, and a thickness of 4–14 nm has been produced by a three-step exfoliation strategy, involving a series of pretreatment–intercalation–exfoliation processes [77].
In short, pretreatment of h-BNs has been performed by using a mixture of H$_2$SO$_4$ and HNO$_3$ and intercalation has been carried out by stirring in NaOH solution. At last, exfoliation has been obtained via microwave-assisted processing in the presence of NH$_4$HCO$_3$.

Ions-intercalation of Na$^+$ and OH$^-$ can improve the exfoliation yield of h-BN (Fig. 9a) [78]. Interestingly, the obtained HO-BNNSs exhibit nonlinear optical (NLO) response with a positive nonlinear absorption coefficient under excitation at 532 and 1064 nm, suggesting a potential application in optical limiting (OL) (Fig. 9b, c). Intercalation-exfoliation can also be realized using LiCl by means of a hydrothermal treatment [79].

The presence of edge-hydroxylated groups in HO-BNNSs prepared at 850 ºC has been detected via an electron energy loss spectroscopy (EELS) [80]. Due to the large amount of OH groups, the addition of only 0.07% of HO-BNNS in poly(N-isopropylacrylamide) (PNIPAM) contributes to improvement in thermal conductivity of 41%.

Strong shear exfoliation forces can be developed using high-speed dispersion homogenizers [81]. This method allows for an efficient production of HO-BNNSs in water without using high temperature, strongly acidic or alkaline environments. BN precursor in water is shear-exfoliated at room temperature with the support of sodium dodecyl sulfate (SDS). The hydroxylated BNNSs can be easily filtered to remove the SDS by centrifugation.
Two-pots: from BN to BNNS then HO-BNNS

$h$-BNNSs are usually prepared by the sonication of their bulk counterpart. The layers can be further hydroxylate by carrying out a series of post-treatments under acid, alkali, oxidant, etc.[82–88].

Figure 10a shows a strategy to functionalize $h$-BNNSs via two synthesis steps [82]. A peroxide reagent, di-tert-butylperoxide (TBP), is firstly introduced to covalently graft alkoxy groups on B atoms of BNNS, and then, the treatment of $\text{H}_2\text{SO}_4/\text{H}_2\text{O}_2$ hydrolytically converts alk oxy into hydroxyl groups. The functionalization by OH groups increases the solubility of BNNSs in water reaching a remarkable value of $\sim 0.1 \text{ mg mL}^{-1}$ (Fig. 10b). The addition of HO-BNNSs (0.1 wt%) into PVA increases the strain-to-break value up to 358%, improving the performance of the pristine polymer (199%) and the
PVA with BNNSs (223%). It can be concluded that an efficient dispersion of hydroxylated BNNSs into a polymeric matrix is able to produce an enhancement of the mechanical properties.

The cytotoxicity on KB cells of HO-BNNSs, obtained by h-BN sonication and hydroxylation under treatment in H2O2 and H2SO4 has been tested using an MTT colorimetric assay [84]. The results show a good compatibility with more than 90% of living cells at the concentration of 500 μg mL−1 suggesting a potential application for bioimaging.

Hydroxylation has been also achieved in hydrothermally treated h-BNNSs in NaOH solution [86]. A 3D hybrid aerogel of epoxy/graphene nanosheet/HO-BNNS has been prepared in a second step. After a 20-cycles compression measurement, the hybrid aerogel preserves 97% of the original height, demonstrating good compressibility and super-elasticity.

HO-BNNSs obtained by oxidation of BNNSs in HNO3 can enhance the dispersion of BNNSs and empower their interfacial interaction with poly(vinylidene fluoride) (PVDF), resulting in an improvement of the corresponding composite dielectric constant and breakdown strength [87]. Consequently, the energy storage density of PVDF/HO-BNNSs can reach up to 13.1 J cm−3, a great enhancement compared with pristine PVDF (440%, 3.0 J cm−3) and PVDF/BNNSs (166%, 7.9 J cm−3).

Two-pots: from BN to HO-BN then HO-BNNS

Hydroxylated h-BNNSs can also be obtained by exfoliating bulk HO-BNs [89–92]. According to Fu et al. [89], HO-BNNSs can be prepared by the following two steps: firstly, a hydrothermal method is used to hydroxylate BN powders by NaOH and KOH; in a second step, the HO-BNs can be exfoliated by sonication. The authors have also assembled the HO-BNNSs into papers showing a thermal conductivity of 58 W m−1 K−1. It is reported that the hydroxylation enables an efficient phonon transfer between adjacent sheets and produces a significant decrease in the interfacial thermal resistance.

Phosphorylated BNNSs obtained by proceeding a reaction between –OH groups of h-BNNSs and H3PO4 can be coated onto carbon steel Q235 substrate by electrophoretic deposition to increase the corrosion resistance. Thus, a passivation film formed by BNNSs can be applied for metal protection [90].

A simple and efficient method to simultaneously exfoliate and functionalize h-BNs is ball milling in the presence of sucrose. This method produces a high product yield of 87% [91] (Fig. 11). HO-BNNSs have been obtained by hydrolysis of the sucrose-grafted BNNSs. Sucrose/BNNSs can be well dispersed in both water and organic liquids. The BNNS/PVA composite films show an enhancement in tensile strength and thermal dissipation capability compared to PVA films.

Bottom-up routes for hydroxylated BNNSs

Bottom-up techniques allow an effective control of h-BN systems growth. So far, several bottom-up techniques have been developed to prepare hydroxylated BNNSs, i.e., chemical vapor deposition (CVD) [93], high-pressure [94], and high-temperature [95–98] methods.

BNNSs has been grown on a copper foil from borazine by CVD. BNNS coating exhibits multiple advantages and, among the others, it provides a good oxidation and corrosion protection of the foil [93].

Precursors such as boric acid and ammonia (H3BO3 and NH3) are frequently employed in high yield synthesis. For example, Singh et al. [94] synthesized HO-BNNSs with H3BO3 and NH3 as precursors by high-pressure route (700 °C, 10 GPa, 24 h). The resulting non-toxic materials make HO-BNNSs potential candidates for boron neutron capture therapy.

HO-BNNSs are also produced by a combination of H3BO3 and urea thermal treatment (950 °C) in N2 atmosphere and sonication of the bulk BN [95]. The product shows a high fluorescent quenching ability and different affinity toward single-stranded and double-stranded DNA, allowing a sensing platform for complementary DNA fragments.

One of the most efficient preparations is a gram-scale synthesis which reaches 88% of BNNSs yield [97]. The method has been developed as a multi-step process: (i) magnesium borate has been firstly synthesized through a solid-phase process by calcination of H3BO3 and Mg(OH)2; (ii) magnesium borates has been then heated in a 200 sccm NH3 atmosphere to form MgO-BN with core–shell structure; (iii) HO-BNNSs have been finally obtained by ultrasonic cleaning in HCl solution to remove the magnesium oxide (Fig. 12). Coupling the experimental results with theoretical simulations with DFT analysis, the
authors have proposed a vapor–solid–solid mechanism where the reaction between vapor NH$_3$ and solid borates results in the production of BNNSs at the vapor–solid interfaces.

**Theoretical models for hydroxylated BNNSs**

The functionalization of $h$-BN is still at its early stage. Because of the reactivity of boron compounds, theoretical calculations are essential to gain an in-depth insight into the electronic and mechanical properties of layered BN [2]. Density functional theory (DFT) and molecular dynamics (MD) simulations provide valuable predictions and corroborations on the impact of hydroxyl chemical species on the BNNSs surfaces [99–104].

Wang et al. [99] investigated the hydroxyl functionalization of $h$-BNNSs by generalized gradient approximation (GGA) with double numerical basis sets plus polarization function (DNP), taking a $(5 \times 5)$ supercell with $x$–$y$ plane periodic conditions as a model. In this work, different OH radical coverage percentages have been considered and the corresponding electronic properties and structural stability have been evaluated. The simulations have identified the boron atom as the most favorable site for OH adsorption. The OH adsorbed at the boron site produces a deformation of $h$-BN sheet. Correspondingly, the involved boron is moved out of the sheet plane by 1.25 Å, increasing the B–N bond length up to 1.53–1.54 Å (pristine bond is 1.45 Å). In this case, the B hybridization changes from sp$^2$ to sp$^3$.

As shown in Fig. 13a, a covalent B–O bond can form between $-$OH and B site of $h$-BN surface with a length of 1.51 Å, which can elongate to 2.40 Å in the
diffusion transition state. Figure 13b shows that the amount of hydroxyl groups on h-BNNSs will influence their stability. The most stable configuration of a single OH radical presents an adsorption energy of −0.88 eV. When increasing −OH, the adsorption energy per radical changes from −1.84 eV for 20% OH coverage to −1.91 and −1.78 eV for 60, and 100%, respectively. Therefore, h-BNNSs functionalized with 60% −OH radicals possess the most stable structure. Additionally, OH groups can be absorbed from the same side or both sides along a zigzag line or armchair configurations. With four OH radicals (Fig. 13c), the most favorable configuration is a linear adsorption structure along with zigzag chain stitch on both sides of BN.

Grosjean et al. [101] have reached similar conclusions by exploring the chemisorption of OH⁻ ions on the h-BN surface via an implicit solvent scheme. The calculation has confirmed that the strongly favorable adsorption of anionic hydroxide on BN layer is connected to the sp³ rehybridization of the bonding atom, boron anchoring site, which results in a negative surface charge of the layered materials. In particular, the authors demonstrate a superior reactivity of h-BN surfaces in contact with hydroxyl anions in comparison with graphene, whose reactivity to OH radical results less marked.

The mechanical as well as the electromechanical properties of h-BNNSs, before and after hydroxylation, have also been investigated by DFT [102]. Two different hydroxylation configurations have been tested: ordered and amorphous. The latter is expected to reflect the experimental results better. The increase in h-BN surface functionalization promotes a progressive increase in interlayer distance (Fig. 14a). For both configurations, at 60% coverage ratio, the interlayer distance approaches to a maximum of about 5.8 Å, not so far from the measured value of 6.9 Å [102]. In general, Young’s modulus decreases under chemisorption of OH groups. For example, for amorphous coverage, the module decreases from 436 to 284 GPa. However, unlike graphene oxide, the hydroxylation of boron nitride has less influence on the mechanical properties, making it a better material in several engineering applications. Moreover, the presence of −OH can reduce their bandgap and endow h-BNNSs with more electro-conductive ability (Fig. 14b). More precisely, it has been reported that under the application of a strain, the bandgap of h-BNNSs reduces from 1.7 eV down to 0.8 eV by increasing the hydroxylation ratio from 20% up to 60%.

Molecular dynamics simulations have been tested to investigate the fracture mechanisms in HO-BNNSs [104]. Coverage of −OH on h-BNNS has a deteriorating effect on the mechanical properties of BNNSs. As for DFT calculations, the fragility of h-BN sheets grows with the hydroxylation. Nevertheless, h-
BNNSs affected by cracks can be strengthened by suitable OH functionalization. In particular, crack edge passivation with OH shows a 32% improvement of fracture toughness, and BNNSs with edge passivation a 48% increase on tensile strength.

**Hydroxylated BNNTs**

**Theoretical studies of hydroxylated BNNTs**

DFT and MD investigations have been carried out to explore the optimized structure and geometry parameters of hydroxylated BNNTs [105–110]. In contrast with their carbon counterparts, BNNTs present a considerable stability under vacuum conditions and are thought to be good candidates for field-emission devices. Studies on the adsorption of O$_2$ and H$_2$O on an open-ended BNNT have demonstrated that the applications of an electric field increase the stability of adsorbates at the tube tip. The adsorbates contribute to change the field emission properties of the BNNTs with a ionization potential slightly lower for N-rich ended NTs than that B-rich ended [105]. Besides, compared with the isolated BNNT of 4.78 eV energy gap, OH functional groups on BNNTs only cause a small reduction of 0.31 eV, while functionalizing with SH groups promotes a semiconductor-metallic transition. Thus, the semiconductor characteristic is well kept in HO-BNNT systems and, at the same time, the solvation and dispersion properties turn out to be improved by the increase in polarity [106].

Duverger et al. [108] studied the effect of OH chemisorption on the optimized geometry of BNNTs. The interaction of HO-BNNTs with different molecules, including azomethine and Pt(IV) complex (anticancer agent), has also been studied. As shown in Fig. 15, the Pt drug molecule located in a symmetric position along the BNNT principal z-axis shows the most stable configuration inside the tube, demonstrating the feasibility of BNNTs for future drug therapy. HO-BNNTs as the carrier for another anticancer molecule, the ifosfamide [109], has been also explored. According to the calculations, the encapsulation of ifosfamide can also take place at room temperature, preserving the drug from unwanted interactions before reaching the cell sites. All these calculations well support the possibility of an effective use of OH-BNNTs for several applications in electronics and nanomedicine and motivate the great experimental effort dedicated to synthesizing BNNTs.

**Methods for hydroxylated BNNTs**

Besides in situ hydroxylation in the BNNTs forming process [111, 112], OH groups can be introduced by a pre-treatment in the presence of H$_2$O$_2$ [33, 113–116], HNO$_3$ [117], NaOH [118], and other reactants [119, 120] (see Fig. 16). Soares et al. [111] prepared BNNTs with OH groups by a bottom-up process with NH$_4$NO$_3$, amorphous boron, hematite (size < 50 nm) as precursors. Then, BNNTs were functionalized by glycol chitosan. Due to the non-covalent coating of glycol chitosan, the products could be further...
As shown in Fig. 16a, the hydroxylation of BNNTs can be achieved via reactions of BN with $\text{H}_2\text{O}_2$ through a hydrothermal route [33]. The HO-BNNTs could be further esterified by the chemical reaction of COOH and OH. The resulting HO-BNNTs appear more effective for mechanical reinforcement of polymers, e.g., polycarbonate (PC) and polyvinyl butyral (PVB), in comparison with the bare BNNTs.

Figure 16 Hydroxylation of BNNTs under the assist of a $\text{H}_2\text{O}_2$, b $\text{HNO}_3$, c NaOH, d DMSO, and e $\text{Br}_2$. Reproduced from Refs. [33, 117–120] with permissions of Copyright (2009) John Wiley and Sons, Copyright (2012) Elsevier, Copyright (2018) The Royal Society of Chemistry, Copyright (2006) John Wiley and Sons, and Copyright (2018) John Wiley and Sons.
HO-BNNTs have also been tested for applications in biotechnology, such as drug delivery [113] and gene silencing [115], after glutaraldehyde modification. Functionalizing BNNTs with OH groups represents a successful strategy for silanization [117]. OH-activated BNNTs under the strong oxidation of HNO₃ (Fig. 16b) can strongly react with APTES by sonication. APTES-grafting on BNNTs allows covalent functionalization with different biological molecules, with encouraging possible developments in biotechnology. Moreover, it has been demonstrated that the hydrolyzation of BNNTs is a necessary step for a correct dispersion in specific polymers [118]. OH-BNNTs obtained in refluxing NaOH have been grafted by aminopropyltrimethoxy (APS) (see Fig. 16c). The APS-functionalization improves the BNNTs dispersion in epoxy resin, leading to 650% enhancement in thermal conductivity and 20% reduction in the thermal expansion coefficient.

Some unconventional approaches have also been proposed to realize the hydroxylation of BNNTs. A solvothermal treatment of BNNTs in DMSO can peel B–N bonds and introduce OH on B sites (Fig. 16d) [119]. Guan et al. [120] designed a liquid bromine treatment to yield both hydroxyl- and amino-functionalized BNNTs (see Fig. 16e). OH and NH₂ groups provide BNNTs with pH-switchable solubility in water. In fact, the bi-functionalized BNNTs are stable in an aqueous solution with pH in the range of 4–8, but precipitate when pH is out of range.

**Other hydroxylated BN materials**

Apart from BNNSs and BNNTs, there are some other hydroxylated BN-based materials that are progressively attracting the attention, such as porous BNs, nanospheres, nanoparticles, nanowires, hollow shells, and nanoscrolls.

**Hydroxylated porous BN materials**

Porous h-BN materials offer the opportunity to modulate the properties and expand the fields of application. A high-temperature treatment of H₃BO₃ to substitute boron atoms with carbon in g-C₃N₄ [121] is a feasible method to prepare porous HO-BNs (Fig. 17a). Due to ~ 30 at% hydroxylation degree, the aqueous solutions result stable at high HO-BNs concentration (2 mg mL⁻¹) and represent good systems for potential applications in anticancer drug loading and delivery. A non-template strategy to fabricate a 3D h-BN foam-like porous monolith has been developed; it provides samples with a high surface area, up to 1406 m² g⁻¹ (Fig. 17b) [122]. It has also been developed a template-free synthesis to obtain porous HO-BNNSs from B₂O₃ and ethylene diamine for water cleaning [123].

Under hydroxyl functionalization of exposed (002) plane, porous BNNSs display an enhancement of visible optical absorption [124]. BNNSs incorporation into TiO₂ promotes a strong visible-light photocatalytic oxidation of organics in solution due to the synergic effect of h-BNs visible absorption and charge transfer to TiO₂ [124]. In addition, it was found that the electrical resistivity of porous HO-BNs is lower and the photocurrent is higher in comparison with commercial h-BNs, extending the application field to photocatalysis and photovoltaic engineering [125].

**Hydroxylated BN nanospheres**

Compared with other nanostructures, h-BN nanospheres possess several advantages in pollution cleaning and drug delivery, especially when they have a hollow structure. Liu et al. [126] prepared h-BN spheres with nanosheet-structure (see Fig. 18a) by thermally evaporating boron powders in NH₃ atmosphere under catalyzation of CoFe₂O₄ and CoO. The corresponding adsorption capacities for dyes, tested with malachite green and methylene blue, were up to 324 and 233 mg g⁻¹ (wt(dye)/wt(material)). The adsorption of Cu²⁺, Pb²⁺, and Cd²⁺ heavy metal ions could reach 678.7, 536.7, and 107.0 mg g⁻¹, respectively, which are much higher than other reported materials (e.g., activated BN and carbon, soy protein hollow microspheres) [126].

As shown in Fig. 18b, another 3D h-BN sphere with nanosheet-structure has been obtained by growing BNNSs in a crystalline metal–organic framework (MOF) [127]. In this work, a novel MOF-sphere containing boron was firstly designed and function as template. In the first step, the zeolitic nanoparticles react with boric acid by a solvothermal treatment and form a new spherical boron-containing MOF. The subsequent annealing at 1000 °C in ammonia environment produces h-BN spheres. Beside the intriguing morphological characteristics, the obtained material has shown interesting properties in both catalysis and energy storage.
A sol–gel route has also been developed to prepare h-BN nanospheres from H$_3$BO$_3$ and melamine [128]. In brief, a carbon spheres template has been impregnated by H$_3$BO$_3$/C$_1$melamine gel, and then h-BN spheres have been formed through a thermal process (500 and 1000 °C, in N$_2$ flow). The nanospheres present a high negative charge, which has been attributed to the adsorption of OH groups. This results in an excellent long-term dispersion stability in water without the need of further surface modification.

h-BN nanospheres have been loaded for delivering anticancer drugs [129, 130]. The h-BN spheres can be hydroxylated by HNO$_3$ and modified by APTES and poly(allylamine hydrochloride)-citraconic anhydride (Fig. 19). The functionalized h-BN system can deliver drugs, such as doxorubicin hydrochloride (DOX), to be released into cancer cells for therapy [130]. It has also been observed that hollow h-BN nanospheres are effective systems against tumor growth and may work for cancer treatment [131]. In this report, the authors find that controlled boron release from hollow h-BN spheres can inhibit the growth of both androgen-sensitive (LNCap) and androgen-independent (DU145) prostate cancer cells. The preventive properties, along with the high hydrophilicity, make hollow h-BN spheres a potential candidate for future anticancer treatments.

Figure 17  a Scheme of the thermal substitution reaction between g-C$_3$N$_4$ and H$_3$BO$_3$ to prepare porous HO-BNs, and the relevant SEM and TEM images. b Picture of 3D h-BN foam-like porous monolith, and the relevant SEM and TEM image. Reproduced from Refs. [121, 122] with permissions of Copyright (2014) American Chemical Society, and Copyright (2016) The Royal Society of Chemistry.
Hydroxylated BN nanoparticles, nanowires and hollow shells

$h$-BN nanoparticles can also be produced via CVD using B(OMe)$_3$ as a precursor in NH$_3$ atmosphere. The CVD temperature is a critical parameter for the morphologies of the products [132]; the $h$-BN nanoparticles aggregate and have an irregular shape if the CVD working temperature is lower than 900 °C, while they result well-dispersed and isotropic if the temperatures are higher. Additionally, the intensity of OH groups at $h$-BN surfaces, studied by monitoring the infrared band OH stretching vibration at $\sim 3600$ cm$^{-1}$, increases with the rise of temperature. Although the authors enlightened the role of oxygen defects in the morphology of $h$-BN nanoparticles, the mechanism related to the enhanced humidity sensitivity of $h$-BN is not fully comprehended even though the crystalline state prevails on the amorphous state.

Mesoporous $h$-BN nanowires can be synthesized from H$_3$BO$_3$ and melamine using Pluronic P123 as templating agent for porosity [133]. A thermal treatment at 1000 °C for 3 h in N$_2$ atmosphere has been employed for the synthesis. The nanowires porosity and oxygen-doping (B–O, B–OH) significantly improve the adsorptive desulfurization activity for dibenzothiophene, and adsorption ability for 4,6-dimethyldibenzothiophene.
$h$-BNNSs and their hollow shells can be prepared via a bottom-up route with $B_2O_3$ and melamine as precursors by adding Mg powders [134]. In particular, Mg powders reduces $B_2O_3$ to B element via a combustive reaction with heat release; this thermal energy further causes the decomposition of melamine and provides the condition for the formation of BN structure.

**Summary and outlook**

The recent interest caused by BN has required the development of new synthesis techniques to produce materials of various shapes and sizes and achieve effective control of surface properties. The current methods for obtaining hydroxylated BN-based materials are summarized in Table 2, coupled with an evaluation on their merits. Hydroxyl functionalization can endow BN-based materials with wide-ranging interesting properties. The advantages of HO-BNs include:

(i) The improvement of solvent-soluble and dispersion in the supporting matrices, as well as biocompatibility, non-toxicity;
(ii) The OH-tailored modification by other molecules, such as APTES, APS, glutaraldehyde, and carboxylic acid compound;
(iii) The empowered abilities in drugs delivery, therapeutics, and pharmaceuticals, as well as adsorption and filtration of pollutants (i.e., dyes, oils, heavy metal ions);

| Table 2 | Evaluation of different methods to prepare hydroxylated BN materials |
|---|---|
| **BN materials** | **Methods** | **Evaluation** | **Refs** |
| BNS in micron-scale | Heating in air | It is the simplest and most direct way to oxidize and hydroxylate bulk $h$-BNs. The thermal treatment is performed at high temperature (over 1000 °C). The hydroxyl degree depends on the temperature and time | [30, 34, 35] |
| Treatment in acidic or alkaline solution | It is efficient and facile, but needs the assist of sonication, heating (< 100 °C), and/or stirring | [32, 36–43] |
| Treatment with strong oxidants | The use of strong oxidants (i.e., $H_2O_2$, $KMnO_4$) requires special attention. Acids, e.g., $H_2SO_4$, $H_3PO_4$, are usually added to enhance the process. Sometimes the strong oxidization process can lead to the presence of a few by-products of nanosheets | [44, 45] |
| Bottom-up routes | Hydroxylated $h$-BNs can be directly obtained by temperature lower than 1100 °C. The treatment can generate some other defects in HO-BNs. If the temperature reaches 1500 °C, the products have to be hydroxylated by the above methods | [47–53] |
| $h$-BNNSs | Sonication-assisted exfoliation | It is a facile method to directly exfoliate BN bulk into HO-BNNSs. The hydroxyl degree is relatively low. For a high hydroxyl degree, some additives have to be introduced, such as, acid, base, salt, etc | [31, 55–63] |
| Ball-milling-assisted exfoliation | Situations similar to “sonication-assisted exfoliation” | [64–72] |
| Hummer’s-like routes | The use of strong oxidants (i.e., $H_2SO_4$, $H_2O_2$, $KMnO_4$, and their mixtures) requires an extra attention. Sometimes the oxidization process can lead to the presence of a few by-products of BN dots | [73–76] |
| Multi-step functionalization method | It can be divided into two stages: one is from BN to BNNS and then to HO-BNNS, the other is from BN to HO-BN and then to HO-BNNS | [82–92] |
| Bottom-up routes | In situ hydroxylation | It generally requires high temperatures and pressures | [93–98] |
| BNNTs | Treatment in $H_2O_2$ | Hydroxylated BNNTs can be directly obtained via a bottom-up approach. It is generally carried out by hydrothermal route and it does not require further purification | [111, 112] |
(iv) The optical properties modification and tuning, e.g., UV–Vis absorption, fluorescence, and NLO;
(v) The possibility to support catalysts, electrodes, etc.

Despite the great efforts that have been made, there are still many challenging phenomena to address and basic studies to perform. The hydroxylation process of BNs is still far from being well understood. Although it is predicted that OH groups can attach to both B and N sites, there are few experimental reports detecting the presence of N–OH in BN systems. Thus, it remains an unanswered question whether N–OH can depict a scenario other than B–OH. On the other hand, OH can be introduced on both surface and edges (armchair and zigzag) of BNNSs, but it is still difficult to appreciate or measure the differences in terms of properties. Furthermore, it is not yet clear which of the OH sites, whether on the surface or on edge, has the greatest influence on the functional properties.

As we know, hydroxyl groups usually are unstable under a thermal process. The thermal stability of nanomaterials and nanocomposites based on hydroxyl functionalized BN materials has, therefore, to be critically evaluated. For instance, the hydroxylation can enhance the filter capacity of BN-membrane. How about its filtering capability after going through a high-temperature process? The same question should also be considered about hydroxyl BN-based thermal conductors, energy storages, activators, mechanical materials, etc.

Although some works show that hydroxyl BN nanomaterials display low or non-toxicity in in vitro experimental results, it should be noted that their toxicity largely depends on the cellular systems, the concentrations, and sizes of BNs. Therefore, it is still important to verify the toxicity for more cell types and carry out more in vivo investigations in the future.

Significant results on HO-BNs systems have been accomplished. However, more theoretical and experimental efforts should be devoted to understand the mechanism through which the OH groups contribute to affect the functional properties. Another important direction of research is to develop advanced technologies using a property-tailored design of hydroxylated BN-based systems. In all mentioned fields, a feasible route is necessary to translate the laboratory findings into commercial engineering, supporting large-scale synthesis of high-quality hydroxylated BN materials.

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Compliance with ethical standards
Conflict of interest The authors declare that they have no conflict of interest.

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